

Lecture notes for
Advanced Data Analysis 1 (ADA1)
Stat 427/527
University of New Mexico

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Chapter 0

Introduction to R, Rstudio, and ggplot

Learning objectives

- identify** a function or operation and describe its use
- apply** functions and operations to achieve a specific result
- predict** answers of calculations written in R
- use** R's functions to get help and numerically summarize data
- apply** ggplot() to organize and reveal patterns visually
- explain** what each plotting option does

Achieving these goals contributes to mastery in these course learning outcomes:

- 1. organize knowledge
- 6. summarize data visually, numerically, and descriptively
- 8. use statistical software

0.1 Syllabus

Tools for this course

Computer: Windows/Mac/Linux

Software: R, text editor (Rstudio)

Brain: scepticism, curiosity, organization
planning, execution, clarity

Syllabus Website: <http://statacumen.com/teaching/ada1>

Let's see what's there:

- Step 0
- Tentative timetable
- Textbook
- i>Clickers
- Learning outcomes – HW00
- Rubrics
- Grading: Homework, team projects, midterm exam, clickers

To do by next class:

- HW 00 due Thursday (or at end of class today).
- Set up R and Rstudio.
- Obtain and register an iClicker.
- Read Hadley Wickham's R style guide stat405.had.co.nz/r-style.html
- Read the rubric <http://statacumen.com/teach/rubrics.pdf> for goal setting and definitions of quality work.
- Obtain (optional) text book for an R programming reference.
- If you need a disability requiring accommodation, please see me and register with the UNM Accessibility Resource Center.

Learning outcomes, syllabus

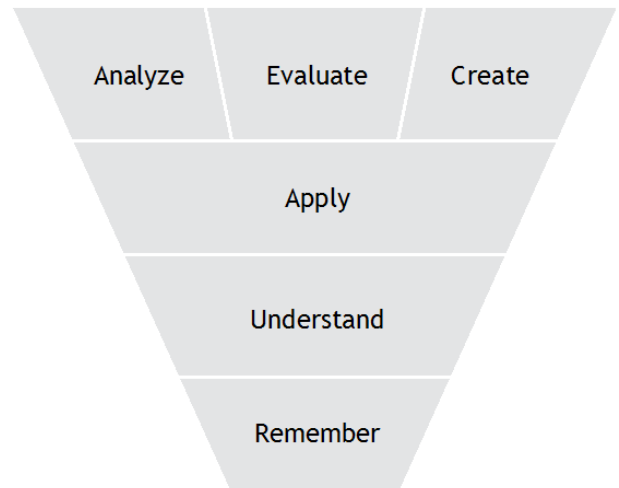
- Which of the learning outcomes on the syllabus are important for you to achieve?

- **Circle** these on your HW00 sheet.
- Put a **check mark** by the one that is *most* important to you.

■ CLICKER Qs — Ch 0, Learning outcomes, General ■

Statistics can be challenging because we operate at the higher levels of Bloom's Taxonomy¹.

- 6 * Create/synthesize
- 5 * Evaluate
- 4 * Analyze
- 3 Apply
- 2 Understand
- 1 Remember

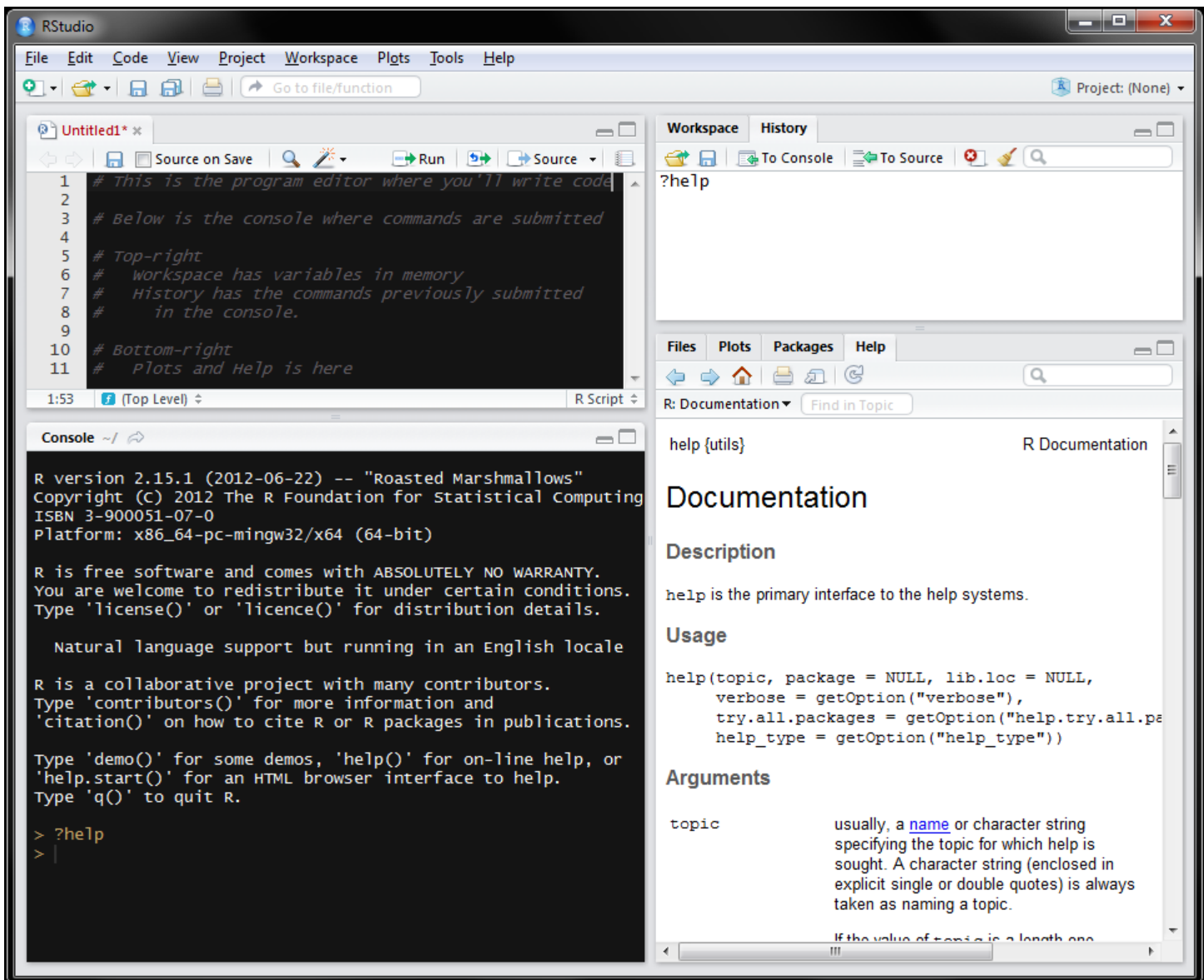


0.2 Rstudio

Setup Install R and Rstudio on your computer, as outlined in Step 0 on the course webpage.

Quick tour Note that I changed my background to black for stealth coding at night...

¹en.wikipedia.org/wiki/Bloom%27s_Taxonomy



1. Upper-left: Program editor window, where you'll write code.
2. Lower-left: Console, where you'll submit commands to be executed.
3. Upper-right: Workspace lists variables in memory, History lists the commands you've previously submitted in the console.
4. Lower-right: Files in current working directory, Plots you've produced, Packages installed and whether they're loaded, Help on commands.

Learning the keyboard shortcuts will make your life more wonderful. (Under Help menu)

Rstudio

Keyboard Shortcuts

Console

Description	Windows & Linux	Mac
Move cursor to Console	Ctrl+2	Ctrl+2
Clear console	Ctrl+L	Command+L
Move cursor to beginning of line	Home	Command+Left
Move cursor to end of line	End	Command+Right
Navigate command history	Up/Down	Up/Down
Popup command history	Ctrl+Up	Command+Up
Interrupt currently executing command	Esc	Esc
Change working directory	Ctrl+Shift+K	Ctrl+Shift+K

Source

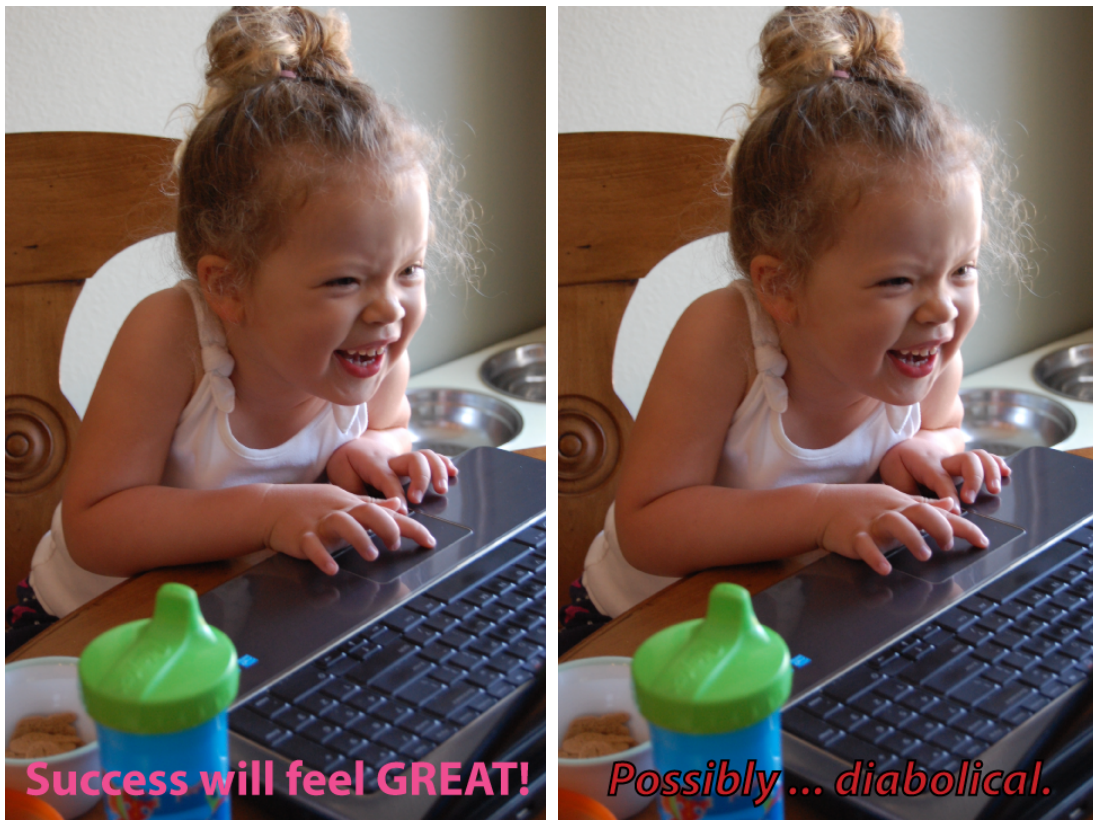
In particular:

- Ctrl-Enter (submit current line or selection in editor window to console)
- Ctrl-1 (move cursor to editor window)
- Ctrl-2 (move cursor to console)
- Ctrl-6 (move cursor to plots)
- Esc (interrupt currently executing command)

Workflow is how you get things done. Here's my recommendation:

1. Work out commands in the editor, then copy/paste into console (or use Ctrl-Enter to submit current line or selection).
2. Your editor will keep a history of what you've done.
3. Make comments to help your future-self recall *why* you did what you did.

0.3 R building blocks





R as calculator In the following sections, look at the commands and output and understand how the command was interpreted to give the output.

```
#### Calculator
# Arithmetic
2 * 10
## [1] 20
1 + 2
## [1] 3
# Order of operations is preserved
1 + 5 * 10
## [1] 51
(1 + 5) * 10
## [1] 60
# Exponents use the ^ symbol
2^5
## [1] 32
9^(1/2)
## [1] 3
```

Vectors A vector is a set of numbers like a column of a spreadsheet. Below these sets of numbers are not technically “vectors”, since they are not in a

row/column structure, though they are ordered and can be referred to by index.

```
#### Vectors
# Create a vector with the c (short for combine) function
c(1, 4, 6, 7)
## [1] 1 4 6 7
c(1:5, 10)
## [1] 1 2 3 4 5 10

# or use a function
# (seq is short for sequence)
seq(1, 10, by = 2)
## [1] 1 3 5 7 9
seq(0, 50, length = 11)
## [1] 0 5 10 15 20 25 30 35 40 45 50
seq(1, 50, length = 11)
## [1] 1.0 5.9 10.8 15.7 20.6 25.5 30.4 35.3 40.2 45.1 50.0
1:10 # short for seq(1, 10, by = 1), or just
## [1] 1 2 3 4 5 6 7 8 9 10
seq(1, 10)
## [1] 1 2 3 4 5 6 7 8 9 10
5:1
## [1] 5 4 3 2 1

# non-integer sequences
# (Note: the [1] at the beginning of lines indicates
#       the index of the first value in that row)
seq(0, 100*pi, by = pi)
## [1] 0.000 3.142 6.283 9.425 12.566 15.708 18.850 21.991
## [9] 25.133 28.274 31.416 34.558 37.699 40.841 43.982 47.124
## [17] 50.265 53.407 56.549 59.690 62.832 65.973 69.115 72.257
## [25] 75.398 78.540 81.681 84.823 87.965 91.106 94.248 97.389
## [33] 100.531 103.673 106.814 109.956 113.097 116.239 119.381 122.522
## [41] 125.664 128.805 131.947 135.088 138.230 141.372 144.513 147.655
## [49] 150.796 153.938 157.080 160.221 163.363 166.504 169.646 172.788
## [57] 175.929 179.071 182.212 185.354 188.496 191.637 194.779 197.920
## [65] 201.062 204.204 207.345 210.487 213.628 216.770 219.911 223.053
## [73] 226.195 229.336 232.478 235.619 238.761 241.903 245.044 248.186
## [81] 251.327 254.469 257.611 260.752 263.894 267.035 270.177 273.319
## [89] 276.460 279.602 282.743 285.885 289.027 292.168 295.310 298.451
## [97] 301.593 304.734 307.876 311.018 314.159
```

Assign variables Assigning objects to variables.

```
##### Assign variables
# Assign a vector to a variable with <-
a <- 1:5
a

## [1] 1 2 3 4 5

b <- seq(15, 3, length = 5)
b

## [1] 15 12 9 6 3

c <- a * b
c

## [1] 15 24 27 24 15
```

Basic functions There are lots of functions available in the base package.

Type `?base` and click on `Index` at the bottom of the help page for a complete list of functions. Other functions to look at are in the `?stats` and `?datasets` packages.

```
##### Basic functions
# Lots of familiar functions work
a

## [1] 1 2 3 4 5

sum(a)
## [1] 15

prod(a)
## [1] 120

mean(a)
## [1] 3

sd(a)
## [1] 1.581

var(a)
## [1] 2.5

min(a)
## [1] 1

median(a)
## [1] 3

max(a)
## [1] 5

range(a)
## [1] 1 5
```

Extracting subsets It will be an extremely important skill to understand the structure of your variables and pull out the information you need from them.

```
#### Extracting subsets
# Specify the indices you want in the square brackets []
a <- seq(0, 100, by = 10)
# blank = include all
a

## [1] 0 10 20 30 40 50 60 70 80 90 100
a[]

## [1] 0 10 20 30 40 50 60 70 80 90 100
# integer +=include, 0=include none, -=exclude
a[5]

## [1] 40
a[c(2, 4, 6, 8)]
## [1] 10 30 50 70
a[0]

## numeric(0)
a[-c(2, 4, 6, 8)]
## [1] 0 20 40 60 80 90 100
a[c(1, 1, 1, 6, 6, 9)]    # subsets can be bigger than the original set
## [1] 0 0 0 50 50 80
a[c(1,2)] <- c(333, 555) # update a subset
a

## [1] 333 555 20 30 40 50 60 70 80 90 100
```

Boolean: True/False Subsets can be selected based on which elements meet specific conditions.

```
#### Boolean
a

## [1] 333 555 20 30 40 50 60 70 80 90 100
(a > 50)           # TRUE/FALSE for each element
## [1] TRUE FALSE FALSE FALSE FALSE FALSE TRUE TRUE TRUE TRUE TRUE
which(a > 50)      # which indices are TRUE
## [1] 1 2 7 8 9 10 11
a[(a > 50)]

## [1] 333 555 60 70 80 90 100
!(a > 50)          # ! negates (flips) TRUE/FALSE values
## [1] FALSE FALSE TRUE TRUE TRUE TRUE FALSE FALSE FALSE FALSE
a[!(a > 50)]

## [1] 20 30 40 50
```

Comparison functions

All your favorite comparisons are available.

```
#### Comparison
# Here they are: < > <= >= != == %in%
a
## [1] 333 555 20 30 40 50 60 70 80 90 100
# equal to
a[(a == 50)]
## [1] 50
# equal to
a[(a == 55)]
## numeric(0)
# not equal to
a[(a != 50)]
## [1] 333 555 20 30 40 60 70 80 90 100
# greater than
a[(a > 50)]
## [1] 333 555 60 70 80 90 100
# less than
a[(a < 50)]
## [1] 20 30 40
# less than or equal to
a[(a <= 50)]
## [1] 20 30 40 50

# which values on left are in the vector on right
(c(10, 14, 40, 60, 99) %in% a)
## [1] FALSE FALSE TRUE TRUE FALSE
```

Boolean operators compare TRUE/FALSE values and return TRUE/-FALSE values.

```
#### Boolean
# & and, | or, ! not
a
## [1] 333 555 20 30 40 50 60 70 80 90 100
a[(a >= 50) & (a <= 90)]
## [1] 50 60 70 80 90
a[(a < 50) | (a > 100)]
## [1] 333 555 20 30 40
a[(a < 50) | !(a > 100)]
## [1] 20 30 40 50 60 70 80 90 100
a[(a >= 50) & !(a <= 90)]
## [1] 333 555 100
```

Missing values The value `NA` (not available) means the value is missing. Any calculation involving `NA` will return an `NA` by default.

```
#### Missing values
NA + 8
## [1] NA
3 * NA
## [1] NA
mean(c(1, 2, NA))
## [1] NA

# Many functions have an na.rm argument (NA remove)
mean(c(NA, 1, 2), na.rm = TRUE)
## [1] 1.5
sum(c(NA, 1, 2))
## [1] NA
sum(c(NA, 1, 2), na.rm = TRUE)
## [1] 3

# Or you can remove them yourself
a <- c(NA, 1:5, NA)
a
## [1] NA 1 2 3 4 5 NA
is.na(a)      # which values are missing?
## [1] TRUE FALSE FALSE FALSE FALSE FALSE TRUE
!is.na(a)      # which values are NOT missing?
## [1] FALSE TRUE TRUE TRUE TRUE TRUE FALSE
a[!is.na(a)]  # return those which are NOT missing
## [1] 1 2 3 4 5
a              # note, this did not change the variable a
## [1] NA 1 2 3 4 5 NA

# To save the results of removing the NAs,
# assign to another variable or reassign to the original variable
# Warning: if you write over variable a then the original version is gone forever!
a <- a[!is.na(a)]
a
## [1] 1 2 3 4 5
```

Your turn!

■ CLICKER Qs — Ch 0, R building blocks, T/F selection 1 ■

■ CLICKER Qs — Ch 0, R building blocks, T/F selection 2 ■

How'd you do?

Outstanding Understanding the operations and how to put them together, without skipping steps.

Good Understanding most of the small steps, missed a couple details.

Hang in there Understanding some of the concepts but all the symbols make my eyes spin.

Reading and writing a new language takes work.

You'll get better as you practice, practice, practice².

Having a buddy to work with will help.

R command review This is a summary of the commands we've seen so far.

```
##### Review
<-
+ - * / ^
c()
seq() # by=, length=
sum(), prod(), mean(), sd(), var(),
min(), median(), max(), range()
a[]
(a > 1), ==, !=, >, <, >=, <=, %in%
&, |, !
NA, mean(a, na.rm = TRUE), !is.na()
```

0.4 Plotting with ggplot2

There are three primary strategies for plotting in R. The first is base graphics, using functions such as `plot()`, `points()`, and `par()`. A second is use of the package `lattice`, which creates very nice graphics quickly, though can be more

²What's your Carnegie Hall that you're working towards?

difficult to create high-dimensional data displays. A third, and the one I will use throughout this course, is using package `ggplot2`, which is an implementation of the “Grammar of Graphics”. While creating a very simple plot requires an extra step or two, creating very complex displays are relatively straightforward as you become comfortable with the syntax.

This section is intended as an introduction to `ggplot()` and some of its capabilities (we will cover these plotting functions and others in detail in later chapters). As a basic introduction, it requires a `data.frame` object as input (a table where each row represents an observation or data point, and each column can have a different data type), and then you define plot layers that stack on top of each other, and each layer has visual/text elements that are mapped to aesthetics (colors, size, opacity). In this way, a simple set of commands can be combined to produce extremely informative displays.

In the example that follows, we consider a dataset `mpg` consisting of fuel economy data from 1999 and 2008 for 38 popular models of car.

```
#### Installing
# only needed once after installing or upgrading R
install.packages("ggplot2")
```

```
#### Library
# each time you start R
# load package ggplot2 for its functions and datasets
library(ggplot2)

# ggplot2 includes a dataset "mpg"

# ? gives help on a function or dataset
?mpg
```

Let's get a look at the dataset we're using.

```
#### mpg dataset
# head() lists the first several rows of a data.frame
head(mpg)

##   manufacturer model displ year cyl      trans drv cty hwy fl class
## 1          audi    a4  1.8 1999    4  auto(15)   f 18 29  p compact
## 2          audi    a4  1.8 1999    4 manual(m5)   f 21 29  p compact
## 3          audi    a4  2.0 2008    4 manual(m6)   f 20 31  p compact
## 4          audi    a4  2.0 2008    4  auto(av)    f 21 30  p compact
## 5          audi    a4  2.8 1999    6  auto(15)    f 16 26  p compact
## 6          audi    a4  2.8 1999    6 manual(m5)   f 18 26  p compact

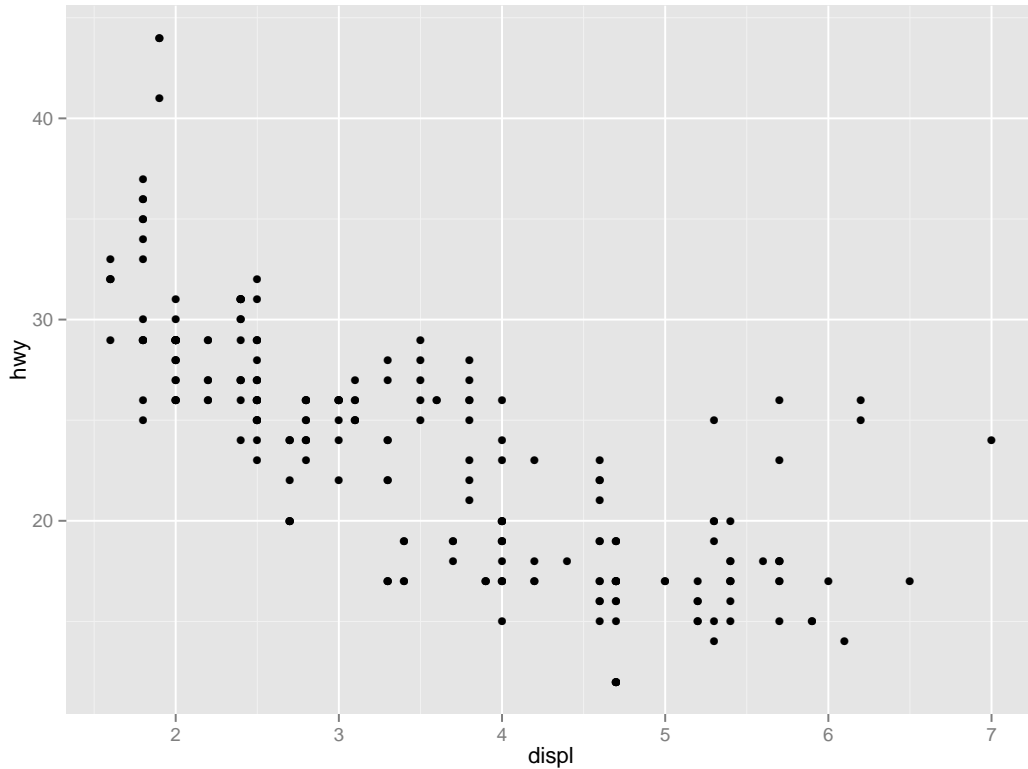
# str() gives the structure of the object
str(mpg)
```

```
## 'data.frame': 234 obs. of 11 variables:
##   $ manufacturer: Factor w/ 15 levels "audi","chevrolet",...: 1 1 1 1 1 1 1 1 1 ...
##   $ model       : Factor w/ 38 levels "4runner 4wd",...: 2 2 2 2 2 2 2 3 3 3 ...
##   $ displ       : num  1.8 1.8 2 2 2.8 2.8 3.1 1.8 1.8 2 ...
##   $ year        : int  1999 1999 2008 2008 1999 1999 2008 1999 1999 2008 ...
##   $ cyl         : int  4 4 4 4 6 6 6 4 4 4 ...
##   $ trans       : Factor w/ 10 levels "auto(av)","auto(13)",...: 4 9 10 1 4 9 1 9 4 10 ...
##   $ drv         : Factor w/ 3 levels "4","f","r": 2 2 2 2 2 2 2 1 1 1 ...
##   $ cty         : int  18 21 20 21 16 18 18 18 16 20 ...
##   $ hwy         : int  29 29 31 30 26 26 27 26 25 28 ...
##   $ fl          : Factor w/ 5 levels "c","d","e","p",...: 4 4 4 4 4 4 4 4 4 4 ...
##   $ class       : Factor w/ 7 levels "2seater","compact",...: 2 2 2 2 2 2 2 2 2 2 ...

# summary() gives frequency tables for categorical variables
# and mean and five-number summaries for continuous variables
summary(mpg)

##      manufacturer           model       displ
## dodge      :37    caravan 2wd      : 11   Min.   :1.60
## toyota     :34    ram 1500 pickup 4wd: 10   1st Qu.:2.40
## volkswagen:27   civic            :  9   Median :3.30
## ford       :25    dakota pickup 4wd :  9   Mean    :3.47
## chevrolet  :19    jetta           :  9   3rd Qu.:4.60
## audi       :18    mustang          :  9   Max.    :7.00
## (Other)    :74    (Other)          :177
##      year        cyl       trans      drv       cty
## Min.   :1999   Min.   :4.00   auto(14) :83   4:103   Min.   : 9.0
## 1st Qu.:1999  1st Qu.:4.00   manual(m5):58   f:106   1st Qu.:14.0
## Median :2004   Median :6.00   auto(15) :39   r: 25   Median :17.0
## Mean   :2004   Mean   :5.89   manual(m6):19   3:13   Mean   :16.9
## 3rd Qu.:2008  3rd Qu.:8.00   auto(s6) :16   5:12   3rd Qu.:19.0
## Max.   :2008   Max.   :8.00   auto(l4) : 6   6:10   Max.   :35.0
##                   (Other)   :13
##      hwy        fl       class
## Min.   :12.0   c:  1   2seater   : 5
## 1st Qu.:18.0   d:  5   compact    :47
## Median :24.0   e:  8   midsize   :41
## Mean   :23.4   p: 52  minivan   :11
## 3rd Qu.:27.0   r:168  pickup    :33
## Max.   :44.0                subcompact:35
##                   suv      :62

##### ggplot_mpg_displ_hwy
# specify the dataset and variables
p <- ggplot(mpg, aes(x = displ, y = hwy))
p <- p + geom_point() # add a plot layer with points
print(p)
```



Geoms, aesthetics, and facets are three concepts we'll see in this section.

Geom: is the “type” of plot

Aesthetics: shape, colour, size, alpha

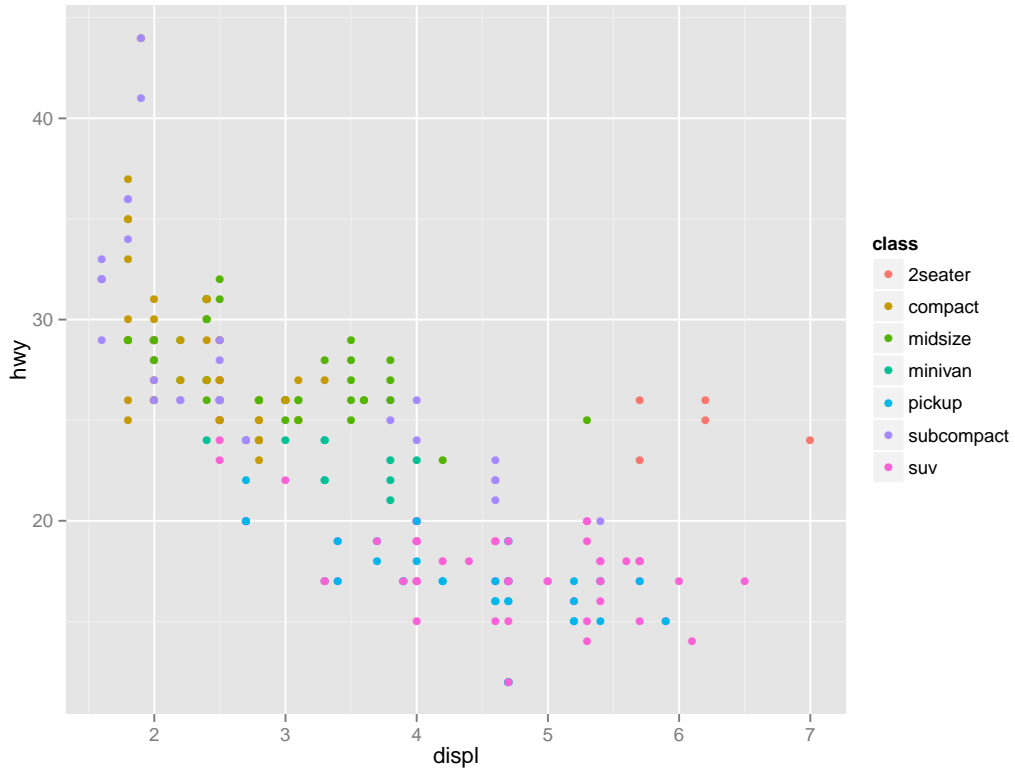
Faceting: “small multiples” displaying different subsets

Help is available. Try searching for examples, too.

- had.co.nz/ggplot2
- had.co.nz/ggplot2/geom_point.html

When certain aesthetics are defined, an appropriate legend is chosen and displayed automatically.

```
#### ggplot_mpg_displ_hwy_colour_class
p <- ggplot(mpg, aes(x = displ, y = hwy))
p <- p + geom_point(aes(colour = class))
print(p)
```



I encourage you to experiment with aesthetics!

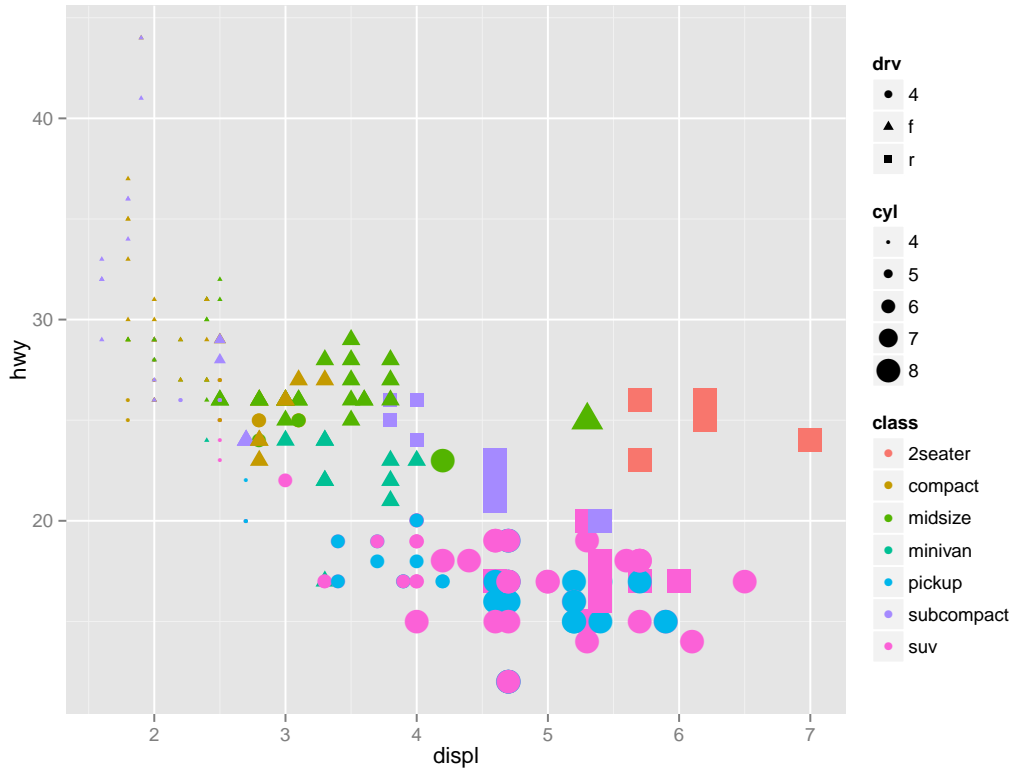
1. Assign variables to aesthetics colour, size, and shape.
2. What's the difference between discrete or continuous variables?
3. What happens when you combine multiple aesthetics?

The behavior of the aesthetics is predictable and customizable.

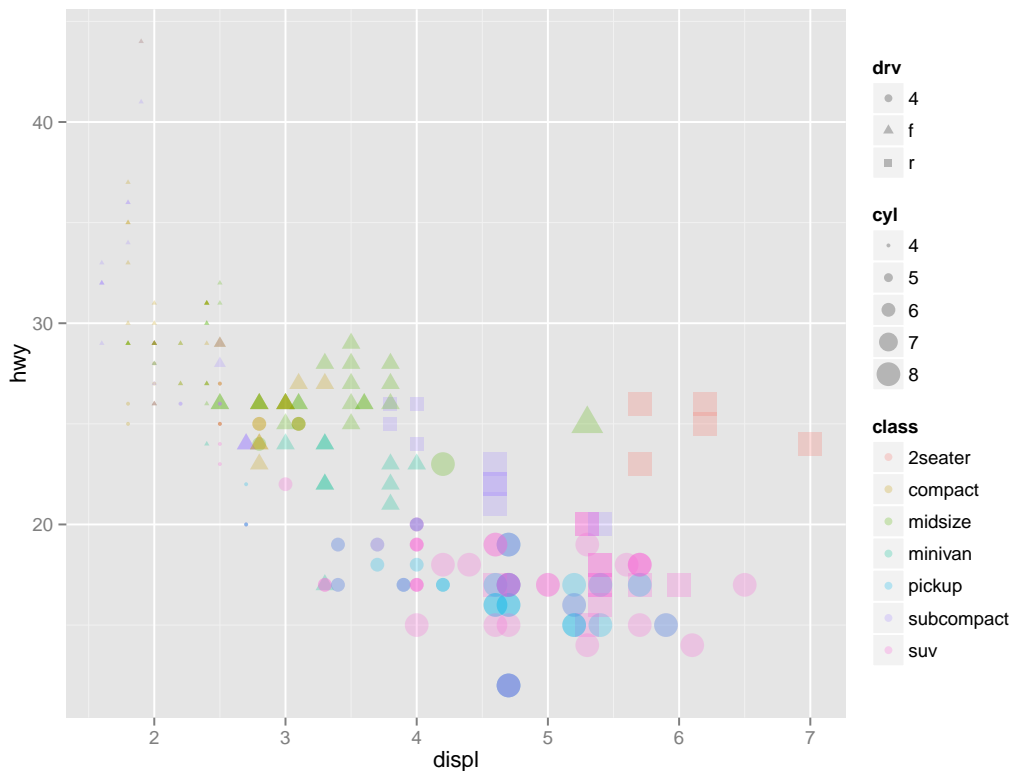
Aesthetic	Discrete	Continuous
colour	Rainbow of colors	Gradient from red to blue
size	Discrete size steps	Linear mapping between radius and value
shape	Different shape for each	Shouldn't work

Let's see a couple examples.

```
#### ggplot_mpg_displ_hwy_colour_class_size_cyl_shape_drv
p <- ggplot(mpg, aes(x = displ, y = hwy))
p <- p + geom_point(aes(colour = class, size = cyl, shape = drv))
print(p)
```



```
##### ggplot_mpg_displ_hwy_colour_class_size_cyl_shape_drv_alpha
p <- ggplot(mpg, aes(x = displ, y = hwy))
p <- p + geom_point(aes(colour = class, size = cyl, shape = drv),
                     , alpha = 1/4) # alpha is the opacity
print(p)
```



Faceting A small multiple³ (sometimes called faceting, trellis chart, lattice chart, grid chart, or panel chart) is a series or grid of small similar graphics or charts, allowing them to be easily compared.

- Typically, small multiples will display different subsets of the data.
- Useful strategy for exploring conditional relationships, especially for large data.

Experiment with faceting of different types. What relationships would you like to see?

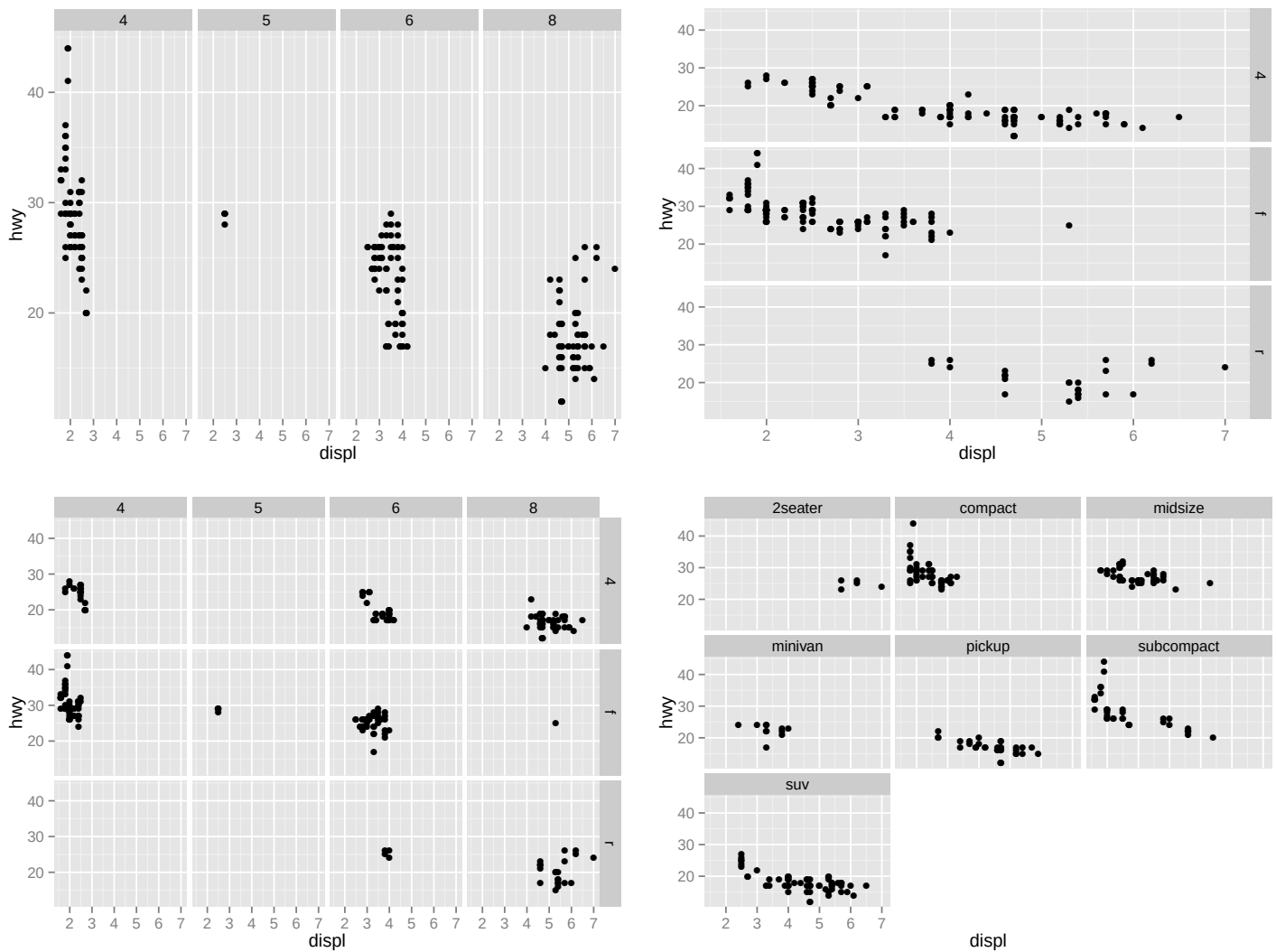
```
#### ggplot_mpg_displ_hwy_facet
# start by creating a basic scatterplot
p <- ggplot(mpg, aes(x = displ, y = hwy))
p <- p + geom_point()

## two methods
# facet_grid(rows ~ cols) for 2D grid, "." for no split.
# facet_wrap(~ var)      for 1D ribbon wrapped into 2D.

# examples of subsetting the scatterplot in facets
p1 <- p + facet_grid(. ~ cyl)      # columns are cyl categories
p2 <- p + facet_grid(drv ~ .)      # rows are drv categories
p3 <- p + facet_grid(drv ~ cyl)    # both
p4 <- p + facet_wrap(~ class)     # wrap plots by class category

# plot all four in one arrangement
library(gridExtra)
## Loading required package: grid
grid.arrange(p1, p2, p3, p4, ncol = 2)
```

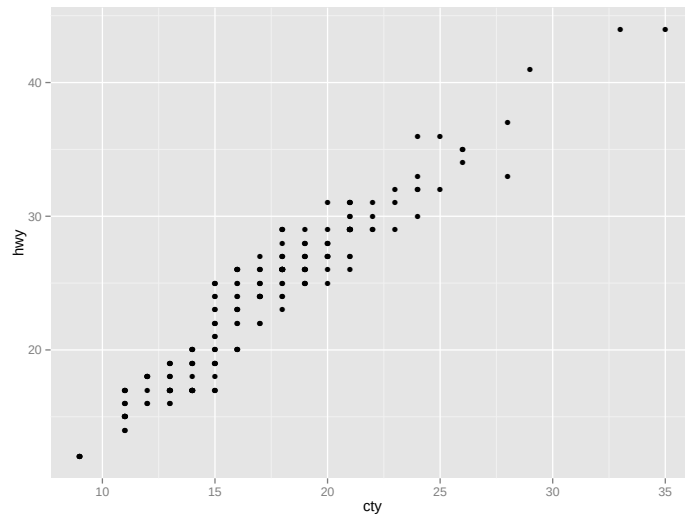
³According to Edward Tufte (*Envisioning Information*, p. 67): “At the heart of quantitative reasoning is a single question: Compared to what? Small multiple designs, multivariate and data bountiful, answer directly by visually enforcing comparisons of changes, of the differences among objects, of the scope of alternatives. For a wide range of problems in data presentation, small multiples are the best design solution.”



0.4.1 Improving plots

How can this plot be improved?

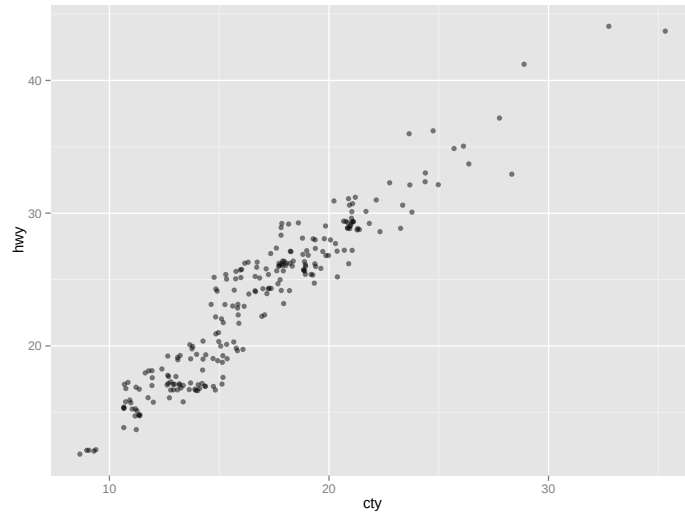
```
#### ggplot_mpg_cty_hwy
p <- ggplot(mpg, aes(x = cty, y = hwy))
p <- p + geom_point()
print(p)
```



Problem: points lie on top of each other, so it's impossible to tell how many observations each point represents.

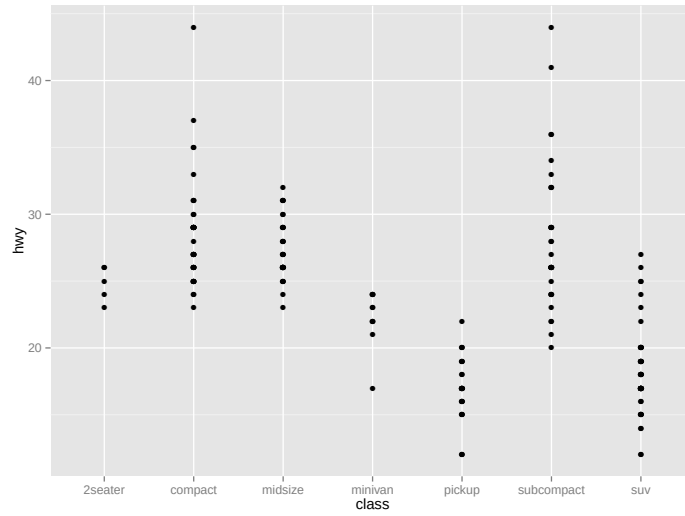
A solution: Jitter the points to reveal the individual points and reduce the opacity to 1/2 to indicate when points overlap.

```
#### ggplot_mpg_cty_hwy_jitter  
p <- ggplot(mpg, aes(x = cty, y = hwy))  
p <- p + geom_point(position = "jitter", alpha = 1/2)  
print(p)
```



How can this plot be improved?

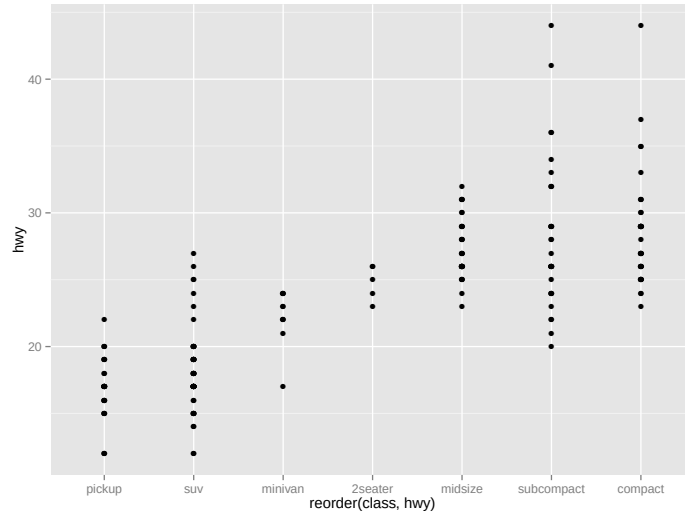
```
#### ggplot_mpg_class_hwy  
p <- ggplot(mpg, aes(x = class, y = hwy))  
p <- p + geom_point()  
print(p)
```



Problem: The classes are in alphabetical order, which is somewhat arbitrary.

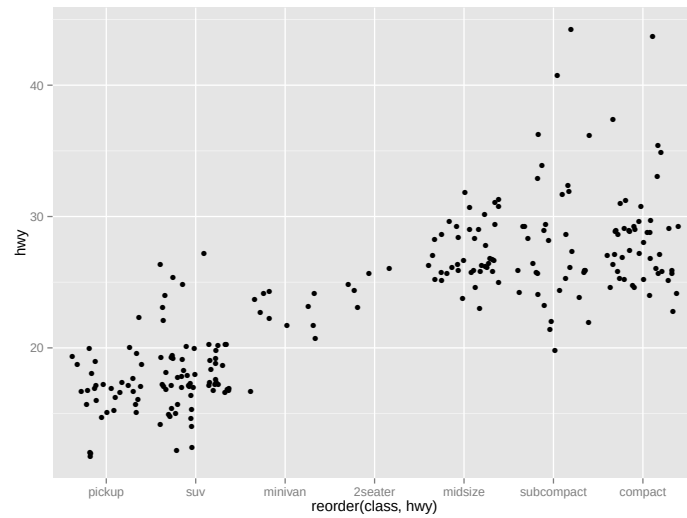
A solution: Reorder the class variable by the mean hwy for a meaningful ordering. Get help with `?reorder` to understand how this works.

```
#### ggplot_mpg_reorder_class_hwy
p <- ggplot(mpg, aes(x = reorder(class, hwy), y = hwy))
p <- p + geom_point()
print(p)
```



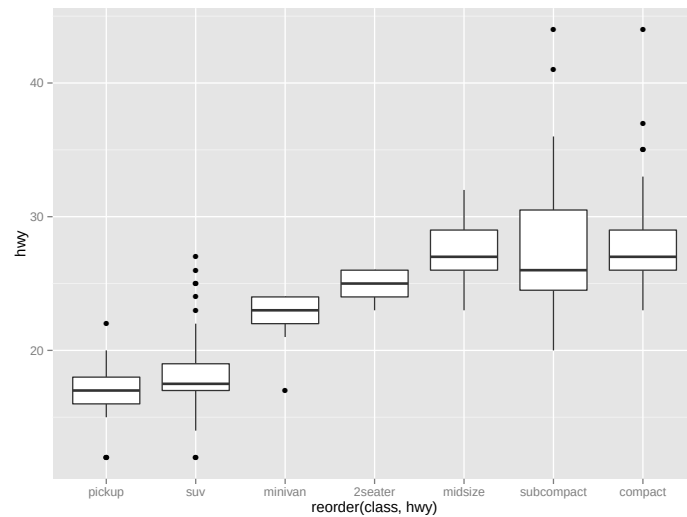
... add jitter

```
#### ggplot_mpg_reorder_class_hwy_jitter
p <- ggplot(mpg, aes(x = reorder(class, hwy), y = hwy))
p <- p + geom_point(position = "jitter")
print(p)
```



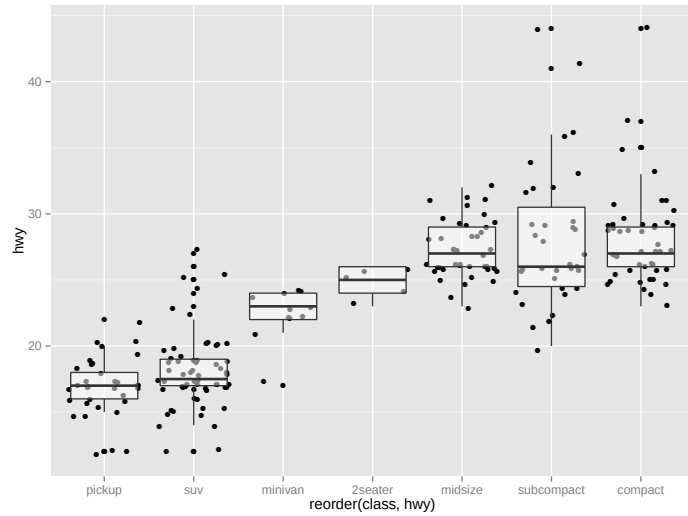
... or replace with boxplots

```
#### ggplot_mpg_reorder_class_hwy_boxplot
p <- ggplot(mpg, aes(x = reorder(class, hwy), y = hwy))
p <- p + geom_boxplot()
print(p)
```



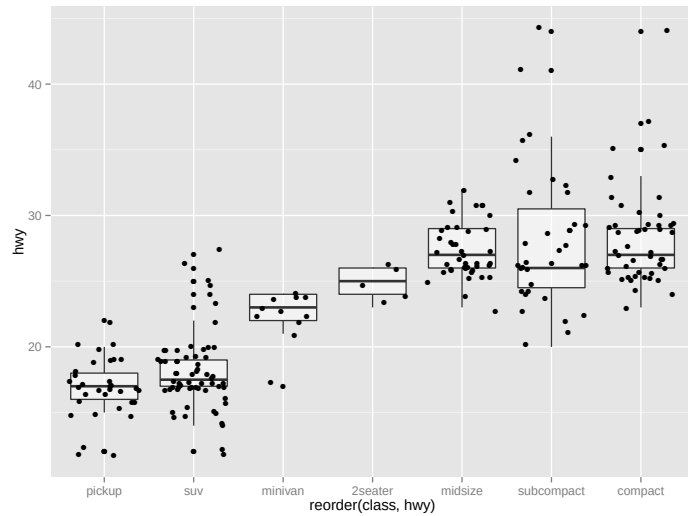
... or jitter those points with reduced-opacity boxplots on top

```
#### ggplot_mpg_reorder_class_hwy_jitter_boxplot
p <- ggplot(mpg, aes(x = reorder(class, hwy), y = hwy))
p <- p + geom_point(position = "jitter")
p <- p + geom_boxplot(alpha = 0.5)
print(p)
```



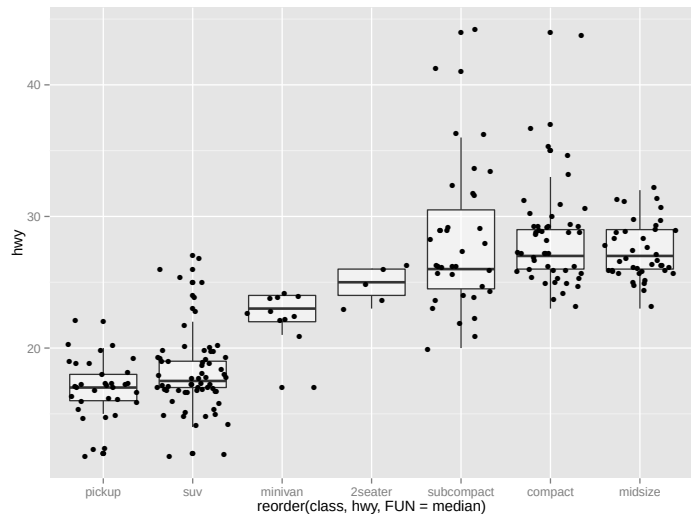
... even better with the boxplots with jittered points on top

```
#### ggplot_mpg_reorder_class_hwy_boxplot_jitter
p <- ggplot(mpg, aes(x = reorder(class, hwy), y = hwy))
p <- p + geom_boxplot(alpha = 0.5)
p <- p + geom_point(position = "jitter")
print(p)
```



... and can easily reorder by median() instead of mean() (mean is the default)

```
#### ggplot_mpg_reorder_class_hwy_boxplot_jitter_median
p <- ggplot(mpg, aes(x = reorder(class, hwy, FUN = median), y = hwy))
p <- p + geom_boxplot(alpha = 0.5)
p <- p + geom_point(position = "jitter")
print(p)
```



R command review This is a summary of the commands we saw in this plotting section.

```
#### Review
library(ggplot2)
?help
head()
str()
summary()
ggplot(df)
geom_point()
aes()
  colour, size, shape, alpha
  position = "jitter"
geom_boxplot()
facet_grid()
facet_wrap()
reorder()
median()
```

One-minute paper:

Muddy Any “muddy” points — anything that doesn’t make sense yet?

Thumbs up Anything you really enjoyed or feel excited about?

0.5 Course Overview

See ADA2 Chapter 1 notes for a brief overview of all we'll cover in this semester of ADA1.

Chapter 1

Summarizing and Displaying Data

Learning objectives

After completing this topic, you should be able to:

use R's functions to get help and numerically summarize data.

apply R's base graphics and ggplot to visually summarize data in several ways.

explain what each plotting option does.

describe the characteristics of a data distribution.

Achieving these goals contributes to mastery in these course learning outcomes:

1. organize knowledge.
6. summarize data visually, numerically, and descriptively.
8. use statistical software.

1.1 Random variables

A **random variable** is a variable whose value is subject to variations due to chance. Random variables fall into two broad categories: qualitative and quantitative.

Qualitative data includes categorical outcomes:

Nominal Outcome is one of several categories.

Ex: Blood group, hair color.

Ordinal Outcome is one of several *ordered* categories.

Ex: Likert data such as (strongly agree, agree, neutral, disagree, strongly disagree).

Quantitative data includes numeric outcomes:

Discrete Outcome is one of a fixed set of numerical values.

Ex: Number of children.

Continuous Outcome is any numerical value.

Ex: Birthweight.

It may not always be perfectly clear which type data belong to, and may sometimes be classified based on the question being asked of the data. Distinction between nominal and ordinal variables can be subjective. For example, for vertebral fracture types: (Wedge, Concavity, Biconcavity, Crush), one could argue that a crush is worse than a biconcavity which is worse than a concavity ..., but this is not self-evident. Distinction between ordinal and discrete variables can be subjective. For example, cancer staging (I, II, III, IV) sounds discrete, but better treated as ordinal because the “distance” between stages may be hard to define and unlikely to be equal. Continuous variables generally measured to a fixed level of precision, which makes them discrete. This “discreteness” of continuous variables is not a problem, providing there are enough levels.

■ CLICKER Qs — Random variables ■

1.2 Numerical summaries

Suppose we have a collection of n individuals, and we measure each individual’s response on one quantitative characteristic, say height, weight, or systolic blood pressure. For notational simplicity, the collected measurements are denoted by Y_1, Y_2, \dots, Y_n , where n is the **sample size**. The order in which the measurements are assigned to the place-holders (Y_1, Y_2, \dots, Y_n) is irrelevant.

Among the numerical summary measures we’re interested in are the **sample mean** \bar{Y} and the **sample standard deviation** s . The sample mean is a

measure of **central location**, or a measure of a “typical value” for the data set. The standard deviation is a measure of **spread** in the data set. These summary statistics should be familiar to you. Let us consider a simple example to refresh your memory on how to compute them.

Suppose we have a sample of $n = 8$ children with weights (in pounds): 5, 9, 12, 30, 14, 18, 32, 40. Then

$$\begin{aligned}\bar{Y} &= \frac{\sum_i Y_i}{n} = \frac{Y_1 + Y_2 + \cdots + Y_n}{n} \\ &= \frac{5 + 9 + 12 + 30 + 14 + 18 + 32 + 40}{8} = \frac{160}{8} = 20.\end{aligned}$$

```
##### Numerical summaries
##### mean
y <- c(5, 9, 12, 30, 14, 18, 32, 40)
mean(y)
## [1] 20
```

The sample standard deviation is the square root of the sample variance

$$\begin{aligned}s^2 &= \frac{\sum_i (Y_i - \bar{Y})^2}{n-1} = \frac{(Y_1 - \bar{Y})^2 + (Y_2 - \bar{Y})^2 + \cdots + (Y_k - \bar{Y})^2}{n-1} \\ &= \frac{(5 - 20)^2 + (9 - 20)^2 + \cdots + (40 - 20)^2}{7} = 156.3, \\ s &= \sqrt{s^2} = 12.5.\end{aligned}$$

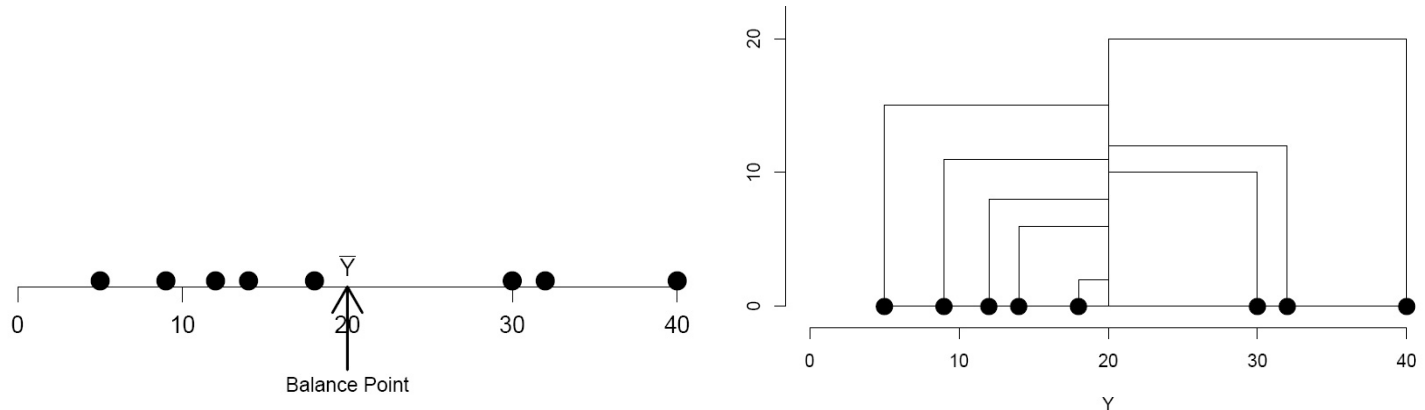
```
##### variance
var(y)
## [1] 156.3
sd(y)
## [1] 12.5
```

Summary statistics have well-defined units of measurement, for example, $\bar{Y} = 20$ lb, $s^2 = 156.3$ lb², and $s = 12.5$ lb. The standard deviation is often used instead of s^2 as a measure of spread because s is measured in the same units as the data.

Remark If the divisor for s^2 was n instead of $n - 1$, then the variance would be the average squared deviation observations are from the center of the data

as measured by the mean.

The following graphs should help you to see some physical meaning of the sample mean and variance. If the data values were placed on a “massless” ruler, the balance point would be the mean (20). The variance is basically the “average” (remember $n - 1$ instead of n) of the total areas of all the squares obtained when squares are formed by joining each value to the mean. In both cases think about the implication of unusual values (**outliers**). What happens to the balance point if the 40 were a 400 instead of a 40? What happens to the squares?



The **sample median** M is an alternative measure of central location. The measure of spread reported along with M is the **interquartile range**, $IQR = Q_3 - Q_1$, where Q_1 and Q_3 are the first and third quartiles of the data set, respectively. To calculate the median and interquartile range, order the data from lowest to highest values, all repeated values included. The ordered weights are

5 9 12 14 18 30 32 40.

```
#### sorting
sort(y)
## [1] 5 9 12 14 18 30 32 40
```

The median M is the value located at the half-way point of the ordered string. There is an even number of observations, so M is defined to be half-way between the two middle values, 14 and 18. That is, $M = 0.5(14 + 18) = 16$ lb. To get the quartiles, break the data into the lower half: 5 9 12 14, and the upper half: 18 30 32 and 40. Then

$Q_1 = \text{first quartile} = \text{median of lower half of data} = 0.5(9+12)=10.5$ lb,
and

Q_3 = third quartile = median of upper half of data = $0.5(30+32) = 31$ lb.

The interquartile range is

$$IQR = Q_3 - Q_1 = 31 - 10.5 = 20.5 \text{ lb.}$$

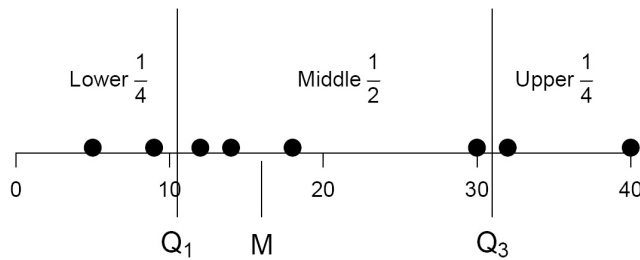
```
#### quartiles
median(y)
## [1] 16
fivenum(y)
## [1] 5.0 10.5 16.0 31.0 40.0

# The quantile() function can be useful, but doesn't calculate Q1 and Q3
# as defined above, regardless of the 9 types of calculations for them!
# summary() is a combination of mean() and quantile(y, c(0, 0.25, 0.5, 0.75, 1))
summary(y)

##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.
##      5.0    11.2   16.0    20.0    30.5    40.0

# IQR
fivenum(y)[c(2,4)]
## [1] 10.5 31.0
fivenum(y)[4] - fivenum(y)[2]
## [1] 20.5
diff(fivenum(y)[c(2,4)])
## [1] 20.5
```

The quartiles, with M being the second quartile, break the data set roughly into fourths. The first quartile is also called the 25^{th} percentile, whereas the median and third quartiles are the 50^{th} and 75^{th} percentiles, respectively. The IQR is the **range** for the middle half of the data.



Suppose we omit the largest observation from the weight data:

5 9 12 14 18 30 32.

How do M and IQR change? With an odd number of observations, there is a unique middle observation in the ordered string which is M . Here $M = 14$ lb. It is unclear which half the median should fall into, so M is placed into both

the lower and upper halves of the data. The lower half is 5 9 12 14, and the upper half is 14 18 30 32. With this convention, $Q_1 = 0.5(9 + 12) = 10.5$ and $Q_3 = 0.5(18 + 30) = 24$, giving $IQR = 24 - 10.5 = 13.5$ (lb).

```
#### remove largest
# remove the largest observation by removing the last of the sorted values
y2 <- sort(y)[-length(y)]
y2
## [1] 5 9 12 14 18 30 32
median(y2)
## [1] 14
fivenum(y2)
## [1] 5.0 10.5 14.0 24.0 32.0
diff(fivenum(y2)[c(2,4)])
## [1] 13.5
```

If you look at the data set with all eight observations, there actually are many numbers that split the data set in half, so the median is not uniquely defined¹, although “everybody” agrees to use the average of the two middle values. With quartiles there is the same ambiguity but no such universal agreement on what to do about it, however, so R will give slightly different values for Q_1 and Q_3 when using `summary()` and some other commands than we just calculated, and other packages will report even different values. This has no practical implication (all the values are “correct”) but it can appear confusing.

Example The data given below are the head breadths in mm for a sample of 18 modern Englishmen, with numerical summaries generated by R.

```
#### Englishmen
hb <- c(141, 148, 132, 138, 154, 142, 150, 146, 155
      , 158, 150, 140, 147, 148, 144, 150, 149, 145)

# see sorted values
sort(hb)
## [1] 132 138 140 141 142 144 145 146 147 148 148 148 149 150 150 150 154
## [17] 155 158

# number of observations is the length of the vector (when no missing values)
length(hb)
```

¹The technical definition of the median for an even set of values includes the entire range between the two center values. Thus, selecting any single value in this center range is convenient and the center of this center range is one sensible choice for the median, M .

```

## [1] 18

# default quartiles
summary(hb)
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.
##      132      142      148      146      150      158

# standard quartiles
fivenum(hb)
## [1] 132.0 142.0 147.5 150.0 158.0
# range() gives the min and max values
range(hb)
## [1] 132 158
# the range of the data is the (max - min), calculated using diff()
diff(range(hb))
## [1] 26

mean(hb)
## [1] 146.5
# standard deviation
sd(hb)
## [1] 6.382
# standard error of the mean
se <- sd(hb)/sqrt(length(hb))

```

Note that `se` is the standard error of the sample mean, $SE_{\bar{Y}} = s/\sqrt{n}$, and is a measure of the precision of the sample mean \bar{Y} .

■ CLICKER Qs — Numerical summaries ■

1.3 Graphical summaries for one quantitative sample

There are four graphical summaries of primary interest: the **dotplot**, the **histogram**, the **stem-and-leaf** display, and the **boxplot**. There are many more possible, but these will often be useful. The plots can be customized. Make liberal use of the help for learning how to customize them. Plots can also

be generated along with many statistical analyses, a point that we will return to repeatedly.

1.3.1 Dotplots

The **dotplot** breaks the range of data into many small-equal width intervals, and counts the number of observations in each interval. The interval count is superimposed on the number line at the interval midpoint as a series of dots, usually one for each observation. In the head breadth data, the intervals are centered at integer values, so the display gives the number of observations at each distinct observed head breadth.

A dotplot of the head breadth data is given below. Of the examples below, the R base graphics `stripchart()` with `method="stack"` resembles the traditional dotplot.

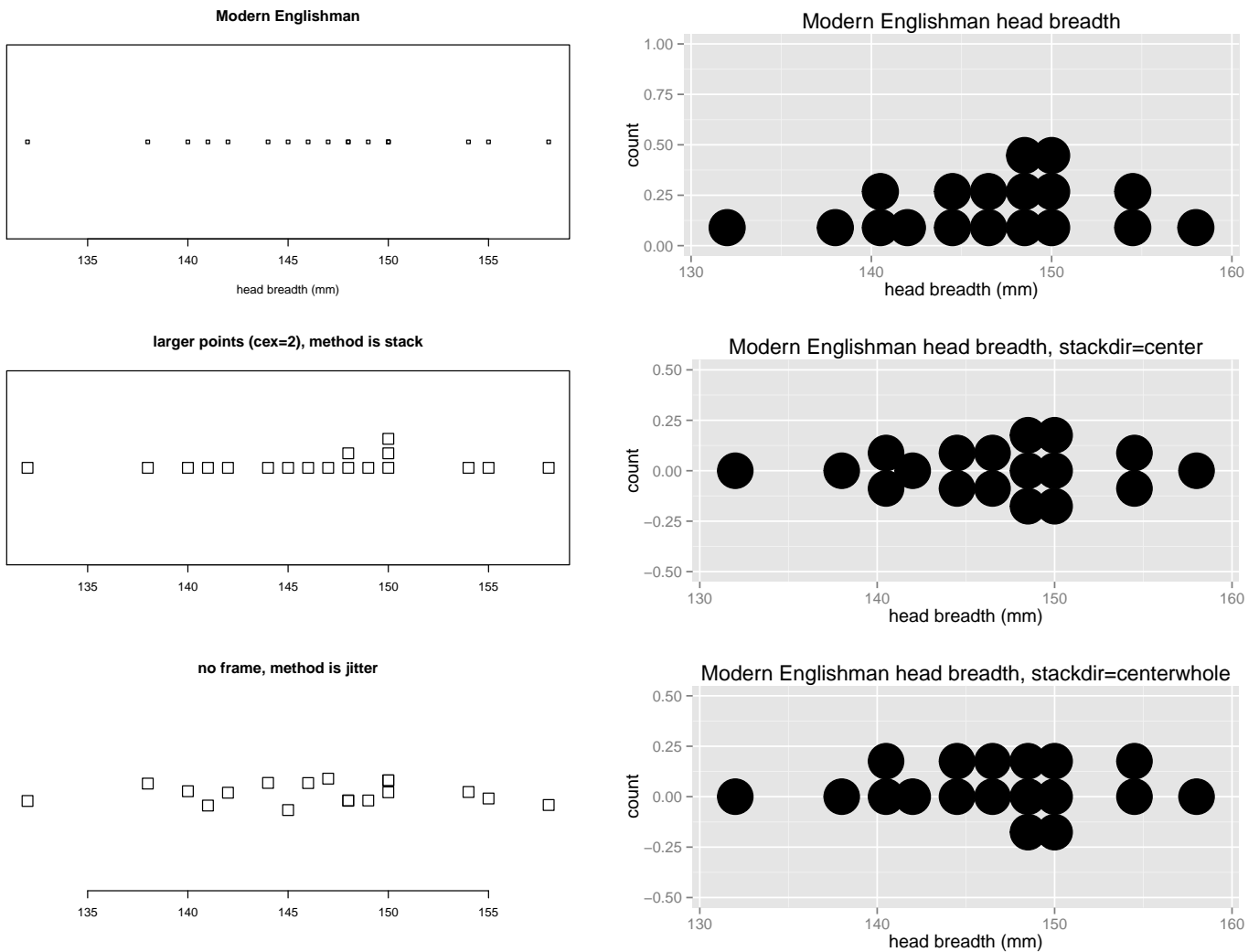
```
#### stripchart-ggplot
# stripchart (dotplot) using R base graphics
# 3 rows, 1 column
par(mfrow=c(3,1))
stripchart(hb, main="Modern Englishman", xlab="head breadth (mm)")
stripchart(hb, method="stack", cex=2
           , main="larger points (cex=2), method is stack")
stripchart(hb, method="jitter", cex=2, frame.plot=FALSE
           , main="no frame, method is jitter")

# dotplot using ggplot
library(ggplot2)
# first put hb vector into a data.frame
hb_df <- data.frame(hb)
p1 <- ggplot(hb_df, aes(x = hb))
p1 <- p1 + geom_dotplot(binwidth = 2)
p1 <- p1 + labs(title = "Modern Englishman head breadth")
p1 <- p1 + xlab("head breadth (mm)")

p2 <- ggplot(hb_df, aes(x = hb))
p2 <- p2 + geom_dotplot(binwidth = 2, stackdir = "center")
p2 <- p2 + labs(title = "Modern Englishman head breadth, stackdir=center")
p2 <- p2 + xlab("head breadth (mm)")

p3 <- ggplot(hb_df, aes(x = hb))
p3 <- p3 + geom_dotplot(binwidth = 2, stackdir = "centerwhole")
p3 <- p3 + labs(title = "Modern Englishman head breadth, stackdir=centerwhole")
p3 <- p3 + xlab("head breadth (mm)")

library(gridExtra)
grid.arrange(p1, p2, p3, ncol=1)
```



1.3.2 Histogram

The **histogram** and **stem-and-leaf** displays are similar, breaking the range of data into a smaller number of equal-width intervals. This produces graphical information about the observed distribution by highlighting where data values cluster. The histogram can use arbitrary intervals, whereas the intervals for the stem-and-leaf display use the base 10 number system. There is more arbitrariness to histograms than to stem-and-leaf displays, so histograms can sometimes be regarded a bit suspiciously.

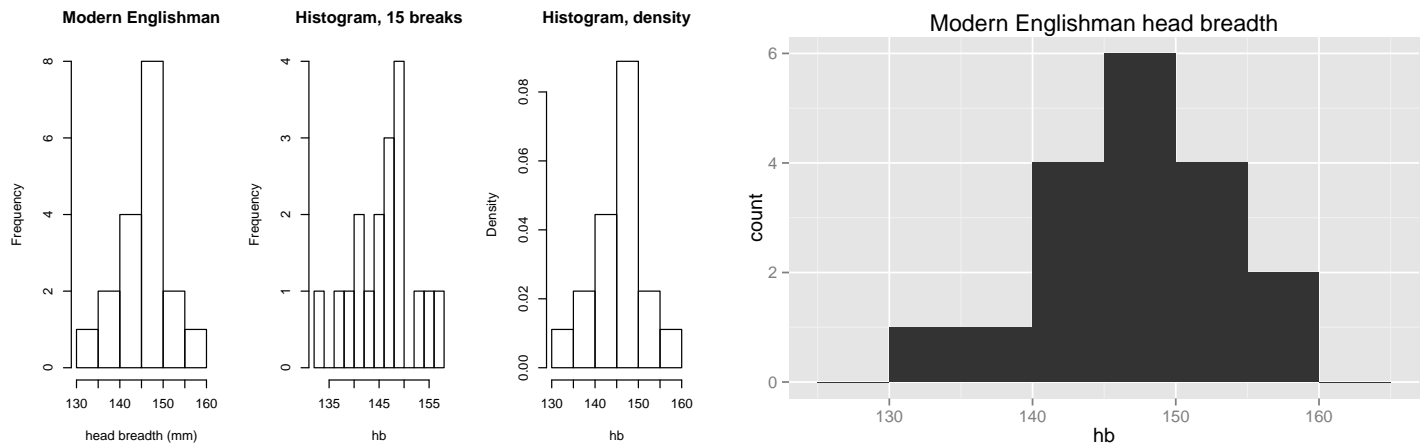
```
#### hist
# histogram using R base graphics
# par() gives graphical options
#   mfrow = "multifigure by row" with 1 row and 3 columns
par(mfrow=c(1,3))
# main is the title, xlab is x-axis label (ylab also available)
```

```

hist(hb, main="Modern Englishman", xlab="head breadth (mm)")
  # breaks are how many bins-1 to use
hist(hb, breaks = 15, main="Histogram, 15 breaks")
  # freq=FALSE changes the vertical axis to density,
  # so the total area of the bars is now equal to 1
hist(hb, breaks = 8, freq = FALSE, main="Histogram, density")

# histogram using ggplot
library(ggplot2)
# first put hb vector into a data.frame
hb_df <- data.frame(hb)
p <- ggplot(hb_df, aes(x = hb))
  # always specify a binwidth for the histogram (default is range/30)
  # try several binwidths
p <- p + geom_histogram(binwidth = 5)
p <- p + labs(title = "Modern Englishman head breadth")
print(p)

```



R allows you to modify the graphical display. For example, with the histogram you might wish to use different midpoints or interval widths. I will let you explore the possibilities.

1.3.3 Stem-and-leaf plot

A **stem-and-leaf** plot is a character display histogram defining intervals for a grouped frequency distribution using the base 10 number system. Intervals are generated by selecting an appropriate number of lead digits for the data values to be the stem. The remaining digits comprise the leaf. It is useful for small samples.

Character plots use typewriter characters to make graphs, and can be convenient for some simple displays, but require use of fixed fonts (like Courier) when copied to a word processing program or they get distorted.

The display almost looks upside down, since larger numbers are on the bottom rather than the top. It is done this way so that if rotated 90 degrees counter-clockwise it *is* a histogram.

The default stem-and-leaf display for the head breadth data is given below. The two columns give the stems and leaves. The data have three digits. The first two comprise the stem. The last digit is the leaf. Thus, a head breadth of 154 has a stem of 15 and leaf of 4. The possible stems are 13, 14, and 15, whereas the possible leaves are the integers from 0 to 9. In the first plot, each stem occurs once, while in the second each stem occurs twice. In the second instance, the first (top) occurrence of a stem value only holds leaves 0 through 4. The second occurrence holds leaves 5 through 9. The display is generated by placing the leaf value for each observation on the appropriate stem line. For example, the top 14 stem holds data values between 140 and 144.99. The stems on this line in the display tell us that four observations fall in this range: 140, 141, 142 and 144. Note that this stem-and-leaf display is an elaborate histogram with intervals of width 5. An advantage of the stem-and-leaf display over the histogram is that the original data values can essentially be recovered from the display.

```
#### stem-and-leaf
# stem-and-leaf plot
stem(hb)

##
## The decimal point is 1 digit(s) to the right of the |
##
## 13 | 28
## 14 | 0124567889
## 15 | 000458

# scale=2 makes plot roughly twice as wide
stem(hb, scale=2)

##
## The decimal point is 1 digit(s) to the right of the |
##
## 13 | 2
## 13 | 8
## 14 | 0124
## 14 | 567889
## 15 | 0004
## 15 | 58

# scale=5 makes plot roughly five times as wide
```

```
stem(hb, scale=5)

##
## The decimal point is at the |
##
## 132 | 0
## 134 |
## 136 |
## 138 | 0
## 140 | 00
## 142 | 0
## 144 | 00
## 146 | 00
## 148 | 000
## 150 | 000
## 152 |
## 154 | 00
## 156 |
## 158 | 0
```

The data values are always *truncated* so that a leaf has one digit. The leaf unit (location of the decimal point) tells us the degree of round-off. This will become clearer in the next example.

Of the three displays, which is the most informative? I think the middle option is best to see the clustering and shape of distributions of numbers.

■ CLICKER Qs — Stem-and-leaf plot ■

1.3.4 Boxplot or box-and-whiskers plot

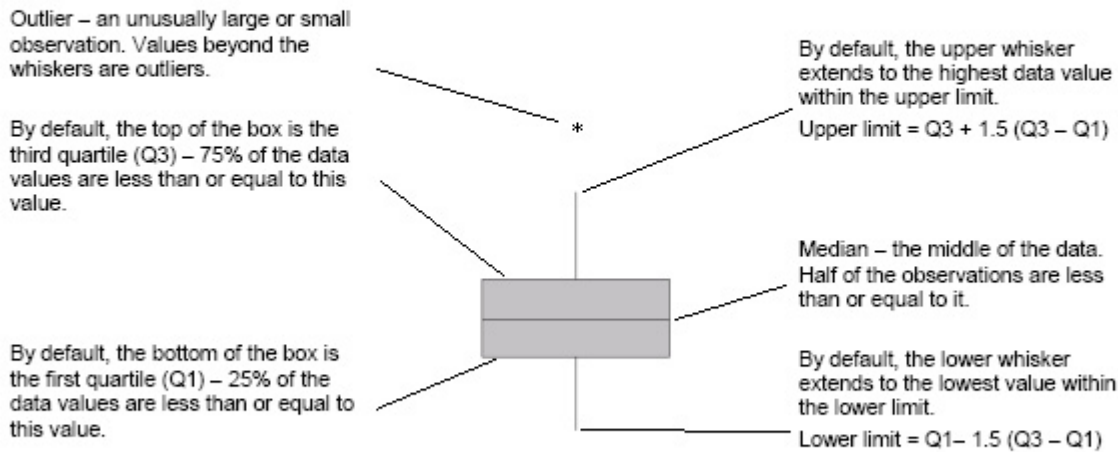
The **boxplot** breaks up the range of data values into regions about the center of the data, measured by the median. The boxplot highlights **outliers** and provides a visual means to assess “**normality**”. The following help entry outlines the construction of the boxplot, given the placement of data values on the axis.

Boxplots

Graph > Boxplot

Stat > EDA > Boxplot

Use boxplots (also called box-and-whisker plots) to assess and compare sample distributions. The figure below illustrates the components of a default boxplot.



Note By default, Minitab uses the quartile method for calculating box endpoints. To change the method for a specific graph to hinge or percentile, use Editor > Edit Interquartile Range Box > Options. To change the method for all future boxplots, use Tools > Options > Individual Graphs > Boxplots.

The endpoints of the box are placed at the locations of the first and third quartiles. The location of the median is identified by the line in the box. The whiskers extend to the data points closest to but not on or outside the outlier fences, which are $1.5IQR$ from the quartiles. Outliers are any values on or outside the outlier fences.

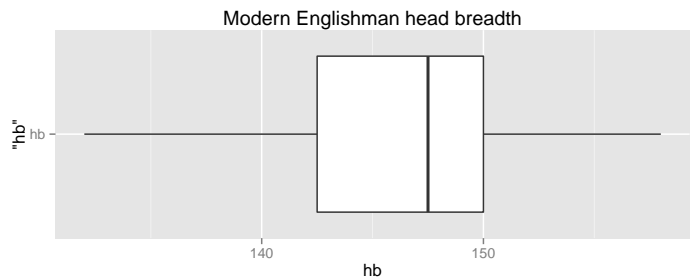
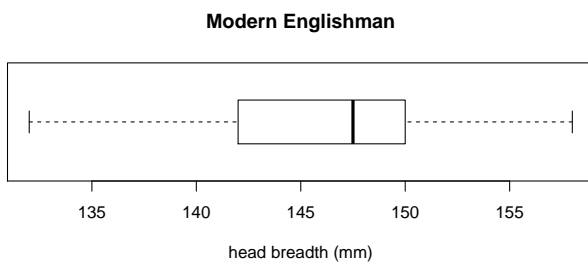
The boxplot for the head breadth data is given below. There are a lot of options that allow you to clutter the boxplot with additional information. Just use the default settings. We want to see the relative location of data (the median line), have an idea of the spread of data (IQR, the length of the box), see the shape of the data (relative distances of components from each other – to be covered later), and identify outliers (if present). The default boxplot has all these components.

Note that the boxplots below are horizontal to better fit on the page. The `horizontal=TRUE` and `coord_flip()` commands do this.

```
#### boxplot
fivenum(hb)

## [1] 132.0 142.0 147.5 150.0 158.0
# boxplot using R base graphics
par(mfrow=c(1,1))
boxplot(hb, horizontal=TRUE
        , main="Modern Englishman", xlab="head breadth (mm)")
```

```
# boxplot using ggplot
library(ggplot2)
# first put hb vector into a data.frame
hb_df <- data.frame(hb)
p <- ggplot(hb_df, aes(x = "hb", y = hb))
p <- p + geom_boxplot()
p <- p + coord_flip()
p <- p + labs(title = "Modern Englishman head breadth")
print(p)
```



■ CLICKER Qs — Boxplots ■

Improvements to the boxplot

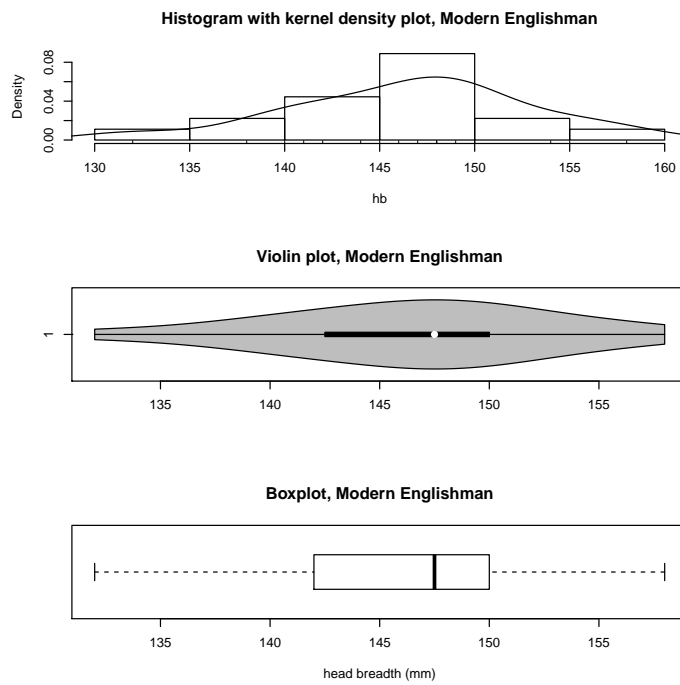
As a quick aside, a violin plot is a combination of a boxplot and a kernel density plot. They can be created using the **vioplot()** function from **vioplot** package.

```
#### vioplot
# vioplot using R base graphics
# 3 rows, 1 column
par(mfrow=c(3,1))

# histogram
hist(hb, freq = FALSE
      , main="Histogram with kernel density plot, Modern Englishman")
# Histogram overlaid with kernel density curve
points(density(hb), type = "l")
# rug of points under histogram
rug(hb)

# violin plot
library(vioplot)
vioplot(hb, horizontal=TRUE, col="gray")
title("Violin plot, Modern Englishman")

# boxplot
boxplot(hb, horizontal=TRUE
        , main="Boxplot, Modern Englishman", xlab="head breadth (mm)")
```



Example: income The data below are incomes in \$1000 units for a sample of 12 retired couples. Numerical and graphical summaries are given. There are two stem-and-leaf displays provided. The first is the default display.

```
#### Income examples
income <- c(7, 1110, 7, 5, 8, 12, 0, 5, 2, 2, 46, 7)
# sort in decreasing order
income <- sort(income, decreasing = TRUE)
income

## [1] 1110   46   12    8    7    7    7    5    5    2    2    0

summary(income)
##    Min. 1st Qu. Median      Mean 3rd Qu.      Max.
##    0.0     4.2     7.0    101.0     9.0   1110.0

# stem-and-leaf plot
stem(income)

##
##   The decimal point is 3 digit(s) to the right of the |
##
##   0 | 000000000000
##   0 |
##   1 | 1
```

Because the two large outliers, I trimmed them to get a sense of the shape of the distribution where most of the observations are.

```
#### remove largest
# remove two largest values (the first two)
income2 <- income[-c(1,2)]
```

```
income2
## [1] 12  8  7  7  7  5  5  2  2  0

summary(income2)
##    Min. 1st Qu. Median     Mean 3rd Qu.     Max.
##    0.00    2.75    6.00    5.50    7.00   12.00

# stem-and-leaf plot
stem(income2)

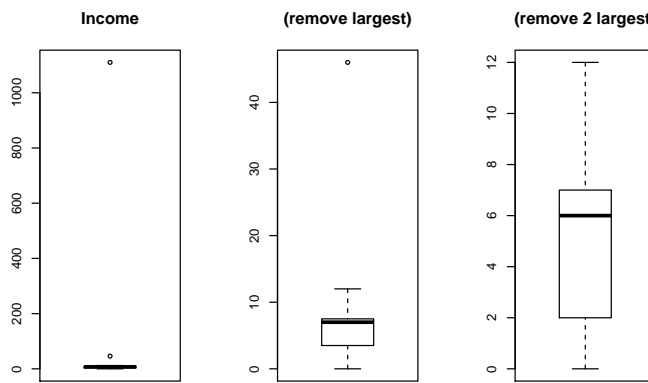
##
##    The decimal point is 1 digit(s) to the right of the |
##
##    0 | 022
##    0 | 557778
##    1 | 2

# scale=2 makes plot roughly twice as wide
stem(income2, scale=2)

##
##    The decimal point is at the |
##
##    0 | 0
##    2 | 00
##    4 | 00
##    6 | 000
##    8 | 0
##   10 |
##   12 | 0
```

Boxplots with full data, then incrementally removing the two largest outliers.

```
#### income-boxplot
# boxplot using R base graphics
# 1 row, 3 columns
par(mfrow=c(1,3))
boxplot(income, main="Income")
boxplot(income[-1], main="(remove largest)")
boxplot(income2, main="(remove 2 largest)")
```



1.4 Interpretation of Graphical Displays for Numerical Data

In many studies, the data are viewed as a subset or **sample** from a larger collection of observations or individuals under study, called the **population**. A primary goal of many statistical analyses is to generalize the information in the sample to **infer** something about the population. For this generalization to be possible, the sample must reflect the basic patterns of the population. There are several ways to collect data to ensure that the sample reflects the basic properties of the population, but the simplest approach, by far, is to take a random or “representative” sample from the population. A **random sample** has the property that every possible sample of a given size has the same chance of being the sample (eventually) selected (though we often do this only once). Random sampling eliminates any systematic biases associated with the selected observations, so the information in the sample should accurately reflect features of the population. The process of sampling introduces random variation or random errors associated with summaries. Statistical tools are used to calibrate the size of the errors.

Whether we are looking at a histogram (or stem-and-leaf, or dotplot) from a sample, or are conceptualizing the histogram generated by the population data, we can imagine approximating the “envelope” around the display with a smooth curve. The smooth curve that approximates the population histogram is called the **population frequency curve** or **population probability**

density function or population distribution². Statistical methods for inference about a population usually make assumptions about the shape of the population frequency curve. A common assumption is that the population has a normal frequency curve. In practice, the observed data are used to assess the reasonableness of this assumption. In particular, a sample display should resemble a population display, provided the collected data are a random or representative sample from the population. Several common shapes for frequency distributions are given below, along with the statistical terms used to describe them.

Unimodal, symmetric, bell-shaped, and no outliers The first display is **unimodal** (one peak), **symmetric**, and **bell-shaped** with no outliers. This is the prototypical normal curve. The boxplot (laid on its side for this display) shows strong evidence of symmetry: the median is about halfway between the first and third quartiles, and the tail lengths are roughly equal. The boxplot is calibrated in such a way that 7 of every 1000 observations are outliers (more than $1.5(Q_3 - Q_1)$ from the quartiles) in samples from a population with a normal frequency curve. Only 2 out of every 1 million observations are extreme outliers (more than $3(Q_3 - Q_1)$ from the quartiles). We do not have any outliers here out of 250 observations, but we certainly could have some without indicating nonnormality. If a sample of 30 observations contains 4 outliers, two of which are extreme, would it be reasonable to assume the population from which the data were collected has a normal frequency curve? Probably not.

```
#### Unimodal, symmetric, bell-shaped, and no outliers (Normal distribution)
## base graphics
# sample from normal distribution
x1 <- rnorm(250, mean = 100, sd = 15)

par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(x1, freq = FALSE, breaks = 20)
points(density(x1), type = "l")
rug(x1)

# violin plot
library(vioplot)
```

²“Distribution function” often refers to the “cumulative distribution function”, which is a different (but one-to-one related) function than what I mean here.

```
vioplot(x1, horizontal=TRUE, col="gray")

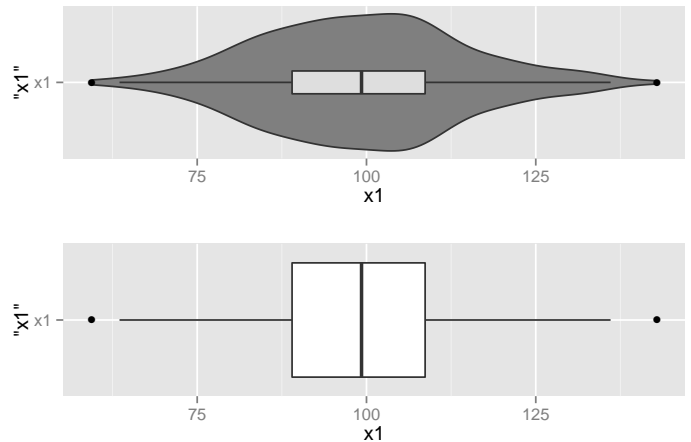
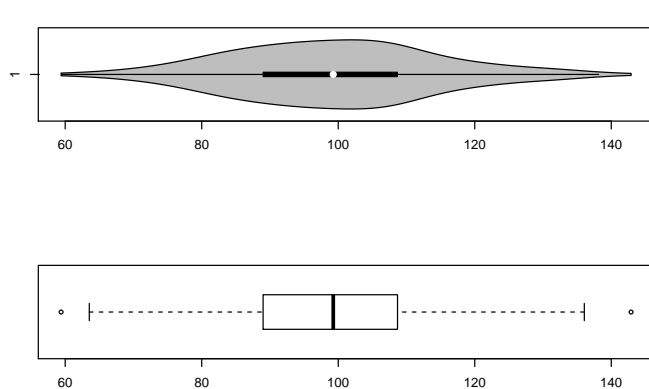
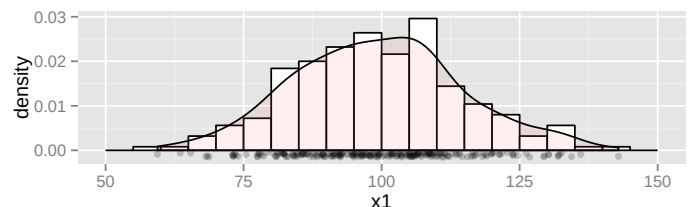
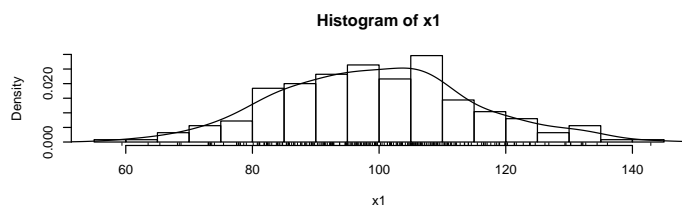
# boxplot
boxplot(x1, horizontal=TRUE)

## ggplot
# Histogram overlaid with kernel density curve
x1_df <- data.frame(x1)
p1 <- ggplot(x1_df, aes(x = x1))
  # Histogram with density instead of count on y-axis
p1 <- p1 + geom_histogram(aes(y=..density..)
                          , binwidth=5
                          , colour="black", fill="white")
  # Overlay with transparent density plot
p1 <- p1 + geom_density(alpha=0.1, fill="#FF6666")
p1 <- p1 + geom_point(aes(y = -0.001)
                      , position = position_jitter(height = 0.0005)
                      , alpha = 1/5)

# violin plot
p2 <- ggplot(x1_df, aes(x = "x1", y = x1))
p2 <- p2 + geom_violin(fill = "gray50")
p2 <- p2 + geom_boxplot(width = 0.2, alpha = 3/4)
p2 <- p2 + coord_flip()

# boxplot
p3 <- ggplot(x1_df, aes(x = "x1", y = x1))
p3 <- p3 + geom_boxplot()
p3 <- p3 + coord_flip()

library(gridExtra)
grid.arrange(p1, p2, p3, ncol=1)
```



```
#### Central statistical moments
```

```
# moments package for 3rd and 4th moments: skewness() and kurtosis()
```

```
library(moments)
```

```
summary(x1)
```

```
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.
##    59.4    89.0   99.3    99.3   109.0   143.0
```

```
sd(x1)
```

```
## [1] 15.1
```

```
skewness(x1)
```

```
## [1] 0.1433
```

```
kurtosis(x1)
```

```
## [1] 2.883
```

```
stem(x1)
```

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

```
##
##   5 | 9
##   6 | 4
##   6 | 5889
##   7 | 3333344
##   7 | 578888899
##   8 | 0111112222223344444
##   8 | 555556666677778888899999999
##   9 | 000111111222223333344
##   9 | 5555555566666667777888888999999999
##  10 | 000001112222223333344444
##  10 | 555555556666666777777888889999999999
##  11 | 000001111122233444
##  11 | 566677788999
```

```
## 12 | 00001123444
## 12 | 5679
## 13 | 00022234
## 13 | 6
## 14 | 3
```

Unimodal, symmetric, heavy-tailed The boxplot is better at highlighting outliers than are other displays. The histogram and stem-and-leaf displays below appear to have the same basic shape as a normal curve (unimodal, symmetric). However, the boxplot shows that we have a dozen outliers in a sample of 250 observations. We would only expect about two outliers in 250 observations when sampling from a population with a normal frequency curve. The frequency curve is best described as unimodal, symmetric, and **heavy-tailed**.

```
#### Unimodal, symmetric, heavy-tailed
# sample from normal distribution
x2.temp <- rnorm(250, mean = 0, sd = 1)
x2 <- sign(x2.temp)*x2.temp^2 * 15 + 100

par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(x2, freq = FALSE, breaks = 20)
points(density(x2), type = "l")
rug(x2)

# violin plot
library(vioplot)
vioplot(x2, horizontal=TRUE, col="gray")

# boxplot
boxplot(x2, horizontal=TRUE)

# Histogram overlaid with kernel density curve
x2_df <- data.frame(x2)
p1 <- ggplot(x2_df, aes(x = x2))
  # Histogram with density instead of count on y-axis
p1 <- p1 + geom_histogram(aes(y=..density..)
                          , binwidth=5
                          , colour="black", fill="white")
  # Overlay with transparent density plot
p1 <- p1 + geom_density(alpha=0.1, fill="#FF6666")
p1 <- p1 + geom_point(aes(y = -0.001)
                      , position = position_jitter(height = 0.0005)
                      , alpha = 1/5)

# violin plot
```

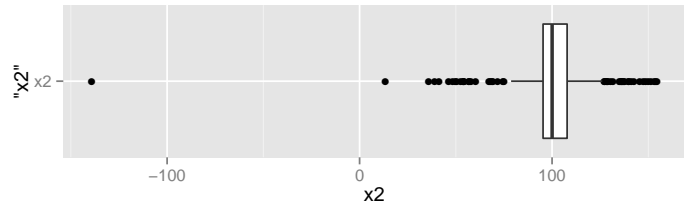
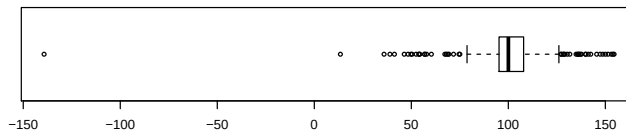
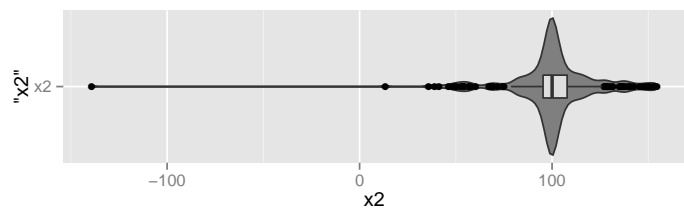
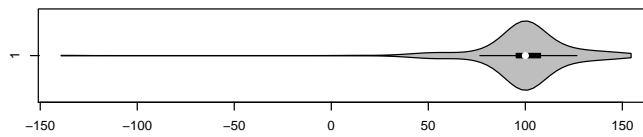
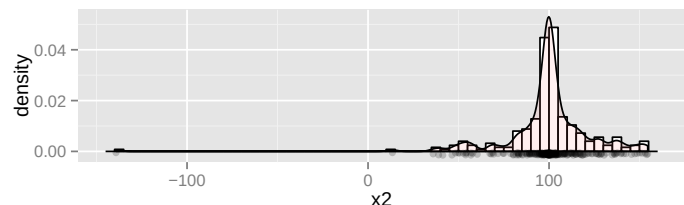
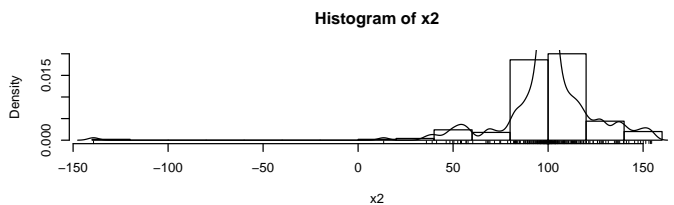
```

p2 <- ggplot(x2_df, aes(x = "x2", y = x2))
p2 <- p2 + geom_violin(fill = "gray50")
p2 <- p2 + geom_boxplot(width = 0.2, alpha = 3/4)
p2 <- p2 + coord_flip()

# boxplot
p3 <- ggplot(x2_df, aes(x = "x2", y = x2))
p3 <- p3 + geom_boxplot()
p3 <- p3 + coord_flip()

library(gridExtra)
grid.arrange(p1, p2, p3, ncol=1)

```



```

summary(x2)
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.
## -139.0    95.3   100.0    99.5   108.0   155.0

sd(x2)
## [1] 25.79

skewness(x2)
## [1] -3.365

kurtosis(x2)
## [1] 31.86

stem(x2)
##
##      The decimal point is 1 digit(s) to the right of the |
##      |
##      -12  | 9
##      -10  |
##      -8   |

```

```
##   -6 |
##   -4 |
##   -2 |
##   -0 |
##   0 | 3
##   2 | 69
##   4 | 168002444778
##   6 | 078992559
##   8 | 002233345557788999900011223334455556666666777777888888888899
##  10 | 0000000000000000000000000000000000000011111111122222222333334444+32
##  12 | 001244667789902566678
##  14 | 001267901345
```

Symmetric, (uniform,) short-tailed Not all symmetric distributions are mound-shaped, as the display below suggests. The boxplot shows symmetry, but the tails of the distribution are shorter (lighter) than in the normal distribution. Note that the distance between quartiles is roughly constant here.

```
#### Symmetric, (uniform,) short-tailed
# sample from uniform distribution
x3 <- runif(250, min = 50, max = 150)

par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(x3, freq = FALSE, breaks = 20)
points(density(x3), type = "l")
rug(x3)

# violin plot
library(vioplot)
vioplot(x3, horizontal=TRUE, col="gray")

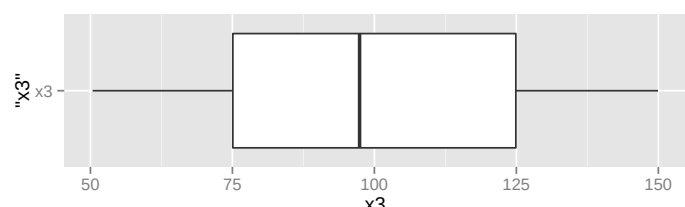
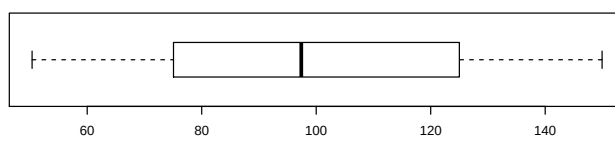
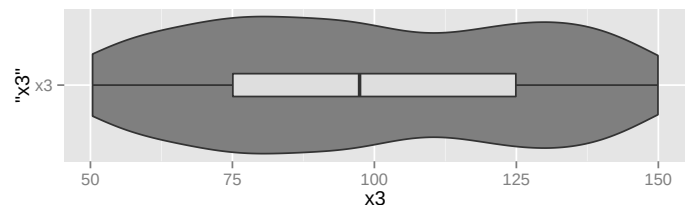
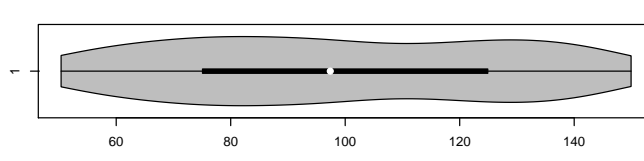
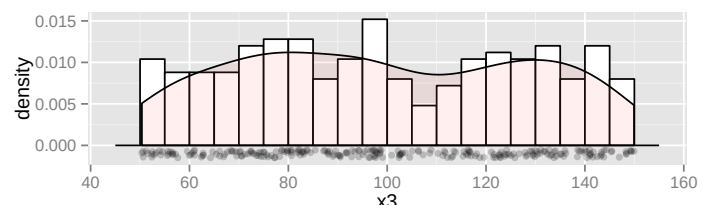
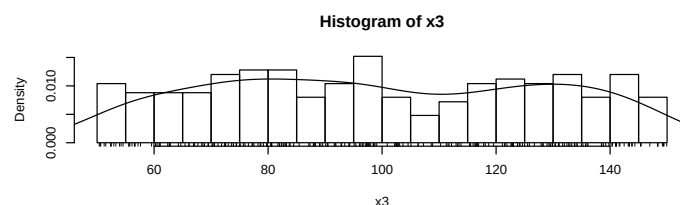
# boxplot
boxplot(x3, horizontal=TRUE)

# Histogram overlaid with kernel density curve
x3_df <- data.frame(x3)
p1 <- ggplot(x3_df, aes(x = x3))
  # Histogram with density instead of count on y-axis
p1 <- p1 + geom_histogram(aes(y=..density..)
                          , binwidth=5
                          , colour="black", fill="white")
  # Overlay with transparent density plot
p1 <- p1 + geom_density(alpha=0.1, fill="#FF6666")
p1 <- p1 + geom_point(aes(y = -0.001)
                      , position = position_jitter(height = 0.0005)
                      , alpha = 1/5)
```

```
# violin plot
p2 <- ggplot(x3_df, aes(x = "x3", y = x3))
p2 <- p2 + geom_violin(fill = "gray50")
p2 <- p2 + geom_boxplot(width = 0.2, alpha = 3/4)
p2 <- p2 + coord_flip()

# boxplot
p3 <- ggplot(x3_df, aes(x = "x3", y = x3))
p3 <- p3 + geom_boxplot()
p3 <- p3 + coord_flip()

library(gridExtra)
grid.arrange(p1, p2, p3, ncol=1)
```



```
summary(x3)
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.
##    50.4     75.1    97.4    99.5   125.0   150.0
sd(x3)
## [1] 28.87
skewness(x3)
## [1] 0.04839
kurtosis(x3)
## [1] 1.789

stem(x3)
##
##      The decimal point is 1 digit(s) to the right of the |
##      5 | 011122223344
##      5 | 5666677778
```

```

##   6 | 000111123334
##   6 | 55667788999
##   7 | 0000112233344444
##   7 | 5556677788999
##   8 | 0000111222223333444
##   8 | 577888899
##   9 | 00001222333444
##   9 | 556677777888888999
##  10 | 122333334
##  10 | 5566799
##  11 | 012223334
##  11 | 566667777889
##  12 | 0122233334444
##  12 | 5566788899
##  13 | 000000112223334444
##  13 | 556668999
##  14 | 00001111222333444
##  14 | 557888999
##  15 | 0

```

The mean and median are identical in a population with a (exact) symmetric frequency curve. The histogram and stem-and-leaf displays for a sample selected from a symmetric population will tend to be fairly symmetric. Further, the sample means and medians will likely be close.

Unimodal, skewed right The distribution below is unimodal, and asymmetric or **skewed**. The distribution is said to be **skewed to the right**, or upper end, because the right tail is much longer than the left tail. The boxplot also shows the skewness – the region between the minimum observation and the median contains half the data in less than 1/5 the range of values. In addition, the upper tail contains several outliers.

```

##### Unimodal, skewed right
# sample from exponential distribution
x4 <- rexp(250, rate = 1)

par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(x4, freq = FALSE, breaks = 20)
points(density(x4), type = "l")
rug(x4)

# violin plot
library(vioplot)
vioplot(x4, horizontal=TRUE, col="gray")

# boxplot

```

```

boxplot(x4, horizontal=TRUE)

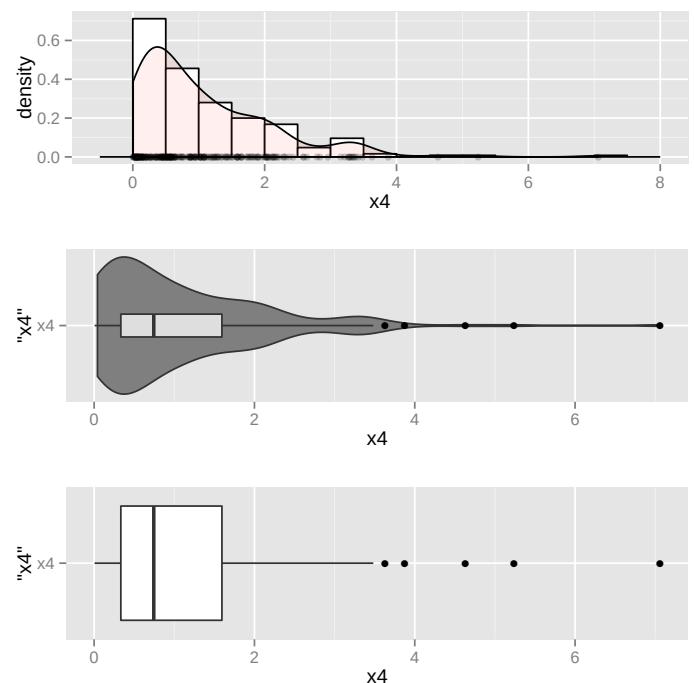
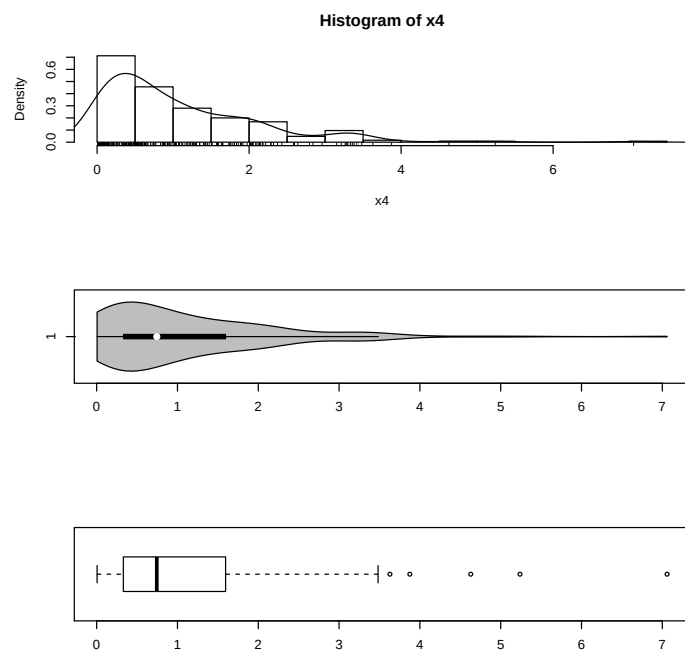
# Histogram overlaid with kernel density curve
x4_df <- data.frame(x4)
p1 <- ggplot(x4_df, aes(x = x4))
  # Histogram with density instead of count on y-axis
p1 <- p1 + geom_histogram(aes(y=..density..),
                           binwidth=0.5,
                           colour="black", fill="white")
  # Overlay with transparent density plot
p1 <- p1 + geom_density(alpha=0.1, fill="#FF6666")
p1 <- p1 + geom_point(aes(y = -0.001),
                      position = position_jitter(height = 0.0005),
                      alpha = 1/5)

# violin plot
p2 <- ggplot(x4_df, aes(x = "x4", y = x4))
p2 <- p2 + geom_violin(fill = "gray50")
p2 <- p2 + geom_boxplot(width = 0.2, alpha = 3/4)
p2 <- p2 + coord_flip()

# boxplot
p3 <- ggplot(x4_df, aes(x = "x4", y = x4))
p3 <- p3 + geom_boxplot()
p3 <- p3 + coord_flip()

library(gridExtra)
grid.arrange(p1, p2, p3, ncol=1)

```



```

summary(x4)
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.

```

```

##   0.005   0.333   0.744   1.080   1.590   7.060
sd(x4)
## [1] 1.044
skewness(x4)
## [1] 1.767
kurtosis(x4)
## [1] 7.77

stem(x4)
##
##      The decimal point is at the |
##
## 0 | 00000000000001111111111111111111111111111122222222222333333334444
## 0 | 55555555555555555555566666666666666667777778888999999999999
## 1 | 0000000000011111112222233333344444
## 1 | 555566666666777888899999
## 2 | 000000111111222233344
## 2 | 66688
## 3 | 022333333444
## 3 | 569
## 4 |
## 4 | 6
## 5 | 2
## 5 |
## 6 |
## 6 |
## 7 | 1

```

Unimodal, skewed left The distribution below is unimodal and **skewed to the left**. The two examples show that extremely skewed distributions often contain outliers in the longer tail of the distribution.

```

##### Unimodal, skewed left
# sample from uniform distribution
x5 <- 15 - rexp(250, rate = 0.5)

par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(x5, freq = FALSE, breaks = 20)
points(density(x5), type = "l")
rug(x5)

# violin plot
library(vioplot)
vioplot(x5, horizontal=TRUE, col="gray")

```

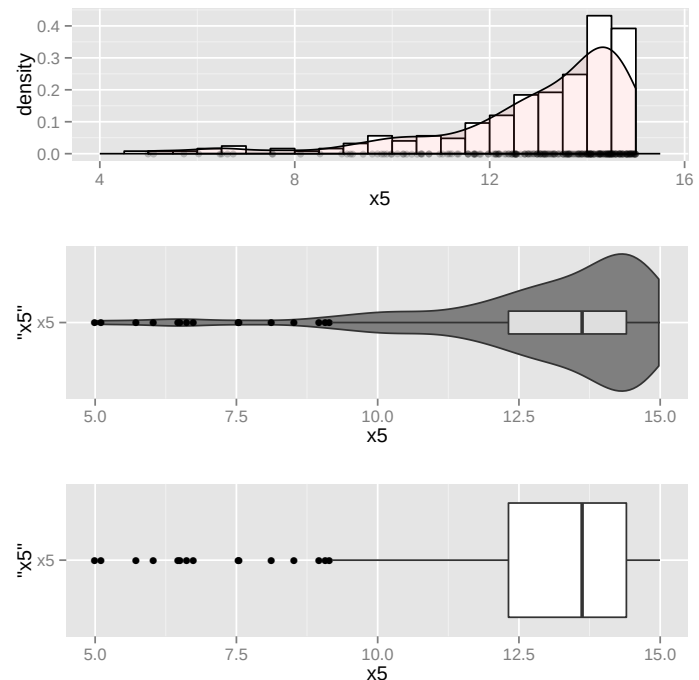
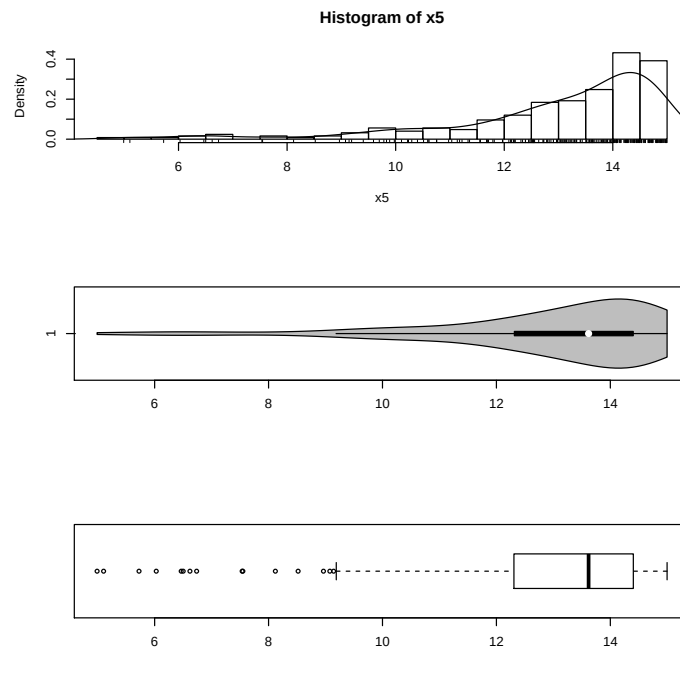
```
# boxplot
boxplot(x5, horizontal=TRUE)

# Histogram overlaid with kernel density curve
x5_df <- data.frame(x5)
p1 <- ggplot(x5_df, aes(x = x5))
  # Histogram with density instead of count on y-axis
p1 <- p1 + geom_histogram(aes(y=..density..),
                           binwidth=0.5,
                           colour="black", fill="white")
  # Overlay with transparent density plot
p1 <- p1 + geom_density(alpha=0.1, fill="#FF6666")
p1 <- p1 + geom_point(aes(y = -0.001),
                      position = position_jitter(height = 0.0005),
                      alpha = 1/5)

# violin plot
p2 <- ggplot(x5_df, aes(x = "x5", y = x5))
p2 <- p2 + geom_violin(fill = "gray50")
p2 <- p2 + geom_boxplot(width = 0.2, alpha = 3/4)
p2 <- p2 + coord_flip()

# boxplot
p3 <- ggplot(x5_df, aes(x = "x5", y = x5))
p3 <- p3 + geom_boxplot()
p3 <- p3 + coord_flip()

library(gridExtra)
grid.arrange(p1, p2, p3, ncol=1)
```



```

summary(x5)
##      Min. 1st Qu. Median     Mean 3rd Qu.    Max.
## 4.99    12.30   13.60    13.00   14.40   15.00

sd(x5)
## [1] 2.03

skewness(x5)
## [1] -1.724

kurtosis(x5)
## [1] 6.008

stem(x5)
##
##      The decimal point is at the |
##
##      4 |
##      5 | 017
##      6 | 05567
##      7 | 55
##      8 | 15
##      9 | 0112467889
##     10 | 0012234566688
##     11 | 00122346667777788
##     12 | 0011122223344455555556667778888999
##     13 | 0000000111111222233333444555566666677788888999999
##     14 | 0000000001111111112222233333334444444445555555555666666+19
##     15 | 0000000

```

Bimodal (multi-modal) Not all distributions are unimodal. The distribution below has two modes or peaks, and is said to be **bimodal**. Distributions with three or more peaks are called **multi-modal**.

```

##### Bimodal (multi-modal)
# sample from uniform distribution
x6 <- c(rnorm(150, mean = 100, sd = 15), rnorm(150, mean = 150, sd = 15))

```

```

par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(x6, freq = FALSE, breaks = 20)
points(density(x6), type = "l")
rug(x6)

```

```

# violin plot
library(vioplot)
vioplot(x6, horizontal=TRUE, col="gray")

# boxplot

```

```

boxplot(x6, horizontal=TRUE)

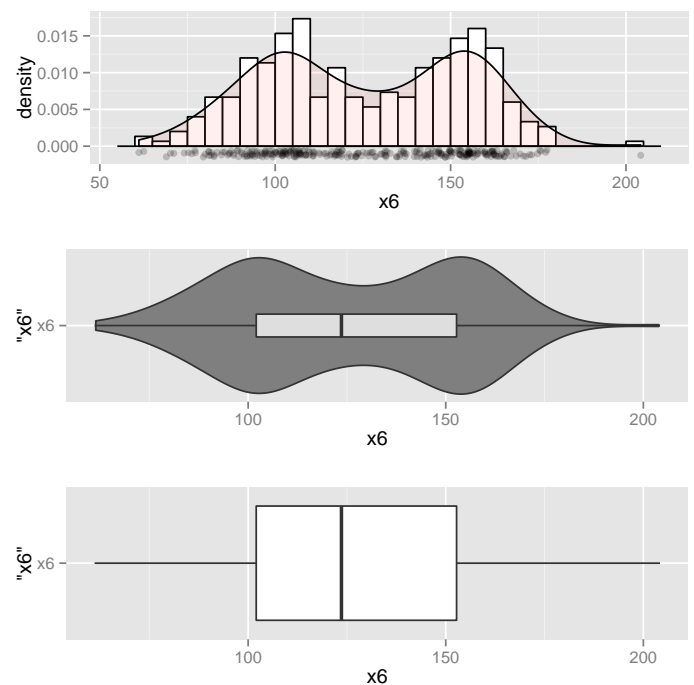
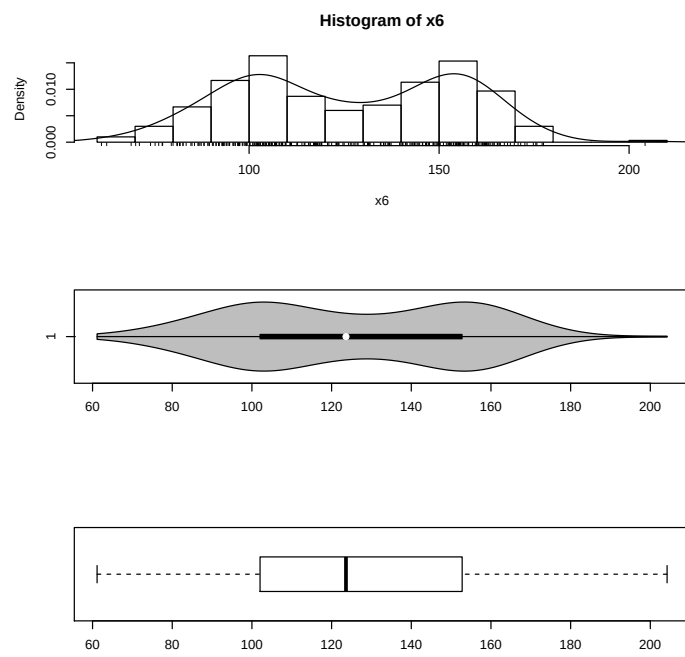
# Histogram overlaid with kernel density curve
x6_df <- data.frame(x6)
p1 <- ggplot(x6_df, aes(x = x6))
  # Histogram with density instead of count on y-axis
p1 <- p1 + geom_histogram(aes(y=..density..),
                           binwidth=5,
                           colour="black", fill="white")
  # Overlay with transparent density plot
p1 <- p1 + geom_density(alpha=0.1, fill="#FF6666")
p1 <- p1 + geom_point(aes(y = -0.001),
                      position = position_jitter(height = 0.0005),
                      alpha = 1/5)

# violin plot
p2 <- ggplot(x6_df, aes(x = "x6", y = x6))
p2 <- p2 + geom_violin(fill = "gray50")
p2 <- p2 + geom_boxplot(width = 0.2, alpha = 3/4)
p2 <- p2 + coord_flip()

# boxplot
p3 <- ggplot(x6_df, aes(x = "x6", y = x6))
p3 <- p3 + geom_boxplot()
p3 <- p3 + coord_flip()

library(gridExtra)
grid.arrange(p1, p2, p3, ncol=1)

```



```

summary(x6)
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.

```

```
##      61.1   102.0   124.0   126.0   153.0   204.0
sd(x6)
## [1] 28.84
skewness(x6)
## [1] -0.01526
kurtosis(x6)
## [1] 1.899

stem(x6)

##
##      The decimal point is 1 digit(s) to the right of the |
##
##      6 | 139
##      7 | 0145677
##      8 | 001112233355677778899
##      9 | 001112223333444555666677778889999
##     10 | 00011122222233334444555556666777777888999
##     11 | 0001111334556667777888899999
##     12 | 112333344556778889
##     13 | 0011122235555677777
##     14 | 00000112233334444566677777788999
##     15 | 000000122333333444455555555566666778889
##     16 | 0000001112222222333444555666789
##     17 | 0113355677
##     18 |
##     19 |
##     20 | 4
```

The boxplot and histogram or stem-and-leaf display (or dotplot) are used **together** to describe the distribution. The boxplot does not provide information about modality – it only tells you about skewness and the presence of outliers.

As noted earlier, many statistical methods assume the population frequency curve is normal. Small deviations from normality usually do not dramatically influence the operating characteristics of these methods. We worry most when the deviations from normality are severe, such as extreme skewness or heavy tails containing multiple outliers.



1.5 Interpretations for examples

The head breadth sample is slightly skewed to the left, unimodal, and has no outliers. The distribution does not deviate substantially from normality. The various measures of central location ($\bar{Y} = 146.5$, $M = 147.5$) are close, which is common with fairly symmetric distributions containing no outliers.

The income sample is extremely skewed to the right due to the presence of two extreme outliers at 46 and 1110. A normality assumption here is unrealistic.

It is important to recognize the influence that outliers can have on the values of \bar{Y} and s . The median and interquartile range are more robust (less sensitive) to the presence of outliers. For the income data $\bar{Y} = 100.9$ and $s = 318$, whereas $M = 7$ and $IQR = 8.3$. If we omit the two outliers, then $\bar{Y} = 5.5$ and $s = 3.8$, whereas $M = 6$ and $IQR = 5.25$.

The mean and median often have similar values in data sets without outliers, so it does not matter much which one is used as the “typical value”. This issue is important, however, in data sets with extreme outliers. In such instances, the median is often more reasonable. For example, is $\bar{Y} = 100.9$ a reasonable measure for a typical income in this sample, given that the second largest income is only 46?

R Discussion I have included basic pointers on how to use R in these notes. I find that copying an R code example from the internet, then modifying the code to apply to my data is a productive strategy. When you’re first learning, “pretty good” is often “good enough”, especially when it comes to plots (in other words, spending 20 minutes vs 2 hours on a plot is fine). I will demonstrate most of what you need and I will be happy to answer questions. You will learn a lot more by using and experimenting with R than by watching.

Chapter 2

Estimation in One-Sample Problems

Learning objectives

After completing this topic, you should be able to:

select graphical displays that meaningfully communicate properties of a sample.

assess the assumptions of the one-sample t-test visually.

decide whether the mean of a population is different from a hypothesized value.

recommend action based on a hypothesis test.

Achieving these goals contributes to mastery in these course learning outcomes:

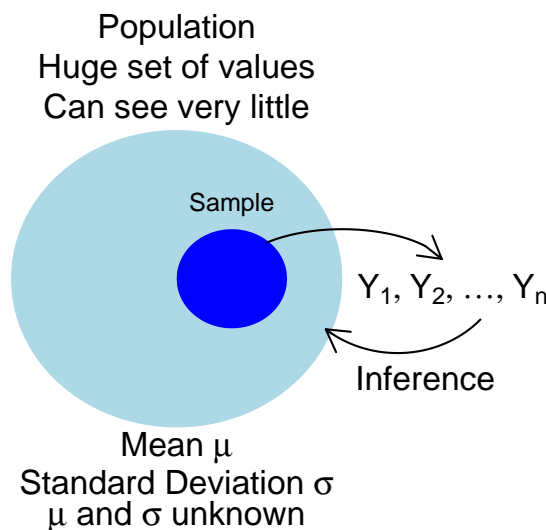
1. organize knowledge.
5. define parameters of interest and hypotheses in words and notation.
6. summarize data visually, numerically, and descriptively.
8. use statistical software.
12. make evidence-based decisions.

2.1 Inference for a population mean

Suppose that you have identified a population of interest where individuals are measured on a single quantitative characteristic, say, weight, height or IQ. You select a random or representative sample from the population with the goal

of estimating the (unknown) **population mean** value, identified by μ . You cannot see much of the population, but you would like to know what is typical in the population (μ). The only information you can see is that in the sample.

This is a standard problem in statistical inference, and the first inferential problem that we will tackle. For notational convenience, identify the measurements on the sample as Y_1, Y_2, \dots, Y_n , where n is the sample size. Given the data, our best guess, or estimate, of μ is the sample mean: $\bar{Y} = \frac{\sum_i Y_i}{n} = \frac{Y_1 + Y_2 + \dots + Y_n}{n}$.



There are two main methods that are used for inferences on μ : **confidence intervals** (CI) and **hypothesis tests**. The standard CI and test procedures are based on the sample mean and the sample standard deviation, denoted by s .



CLICKER Qs — Inference for a population mean, 2



2.1.1 Standard error, LLN, and CLT

The **standard error (SE)** is the standard deviation of the **sampling distribution** of a statistic.

The **sampling distribution** of a statistic is the distribution of that statistic, considered as a random variable, when derived from a random sample of size n .

The **standard error of the mean (SEM)** is the standard deviation of the sample-mean's estimate of a population mean. (It can also be viewed as the standard deviation of the error in the sample mean relative to the true mean, since the sample mean is an unbiased estimator.) SEM is usually estimated by the sample estimate of the population standard deviation (sample standard deviation) divided by the square root of the sample size (assuming statistical independence of the values in the sample):

$$SE_{\bar{Y}} = s/\sqrt{n}$$

where s is the sample standard deviation (i.e., the sample-based estimate of the standard deviation of the population), and n is the size (number of observations) of the sample.

In probability theory, the **law of large numbers (LLN)** is a theorem that describes the result of performing the same experiment a large number of times. According to the law, the average of the results obtained from a large number of trials (the sample mean, \bar{Y}) should be close to the expected value (the population mean, μ), and will tend to become closer as more trials are performed.

In probability theory, the **central limit theorem (CLT)** states that, given certain conditions, the mean of a sufficiently large number of independent random variables, each with finite mean and variance, will be approximately normally distributed¹.

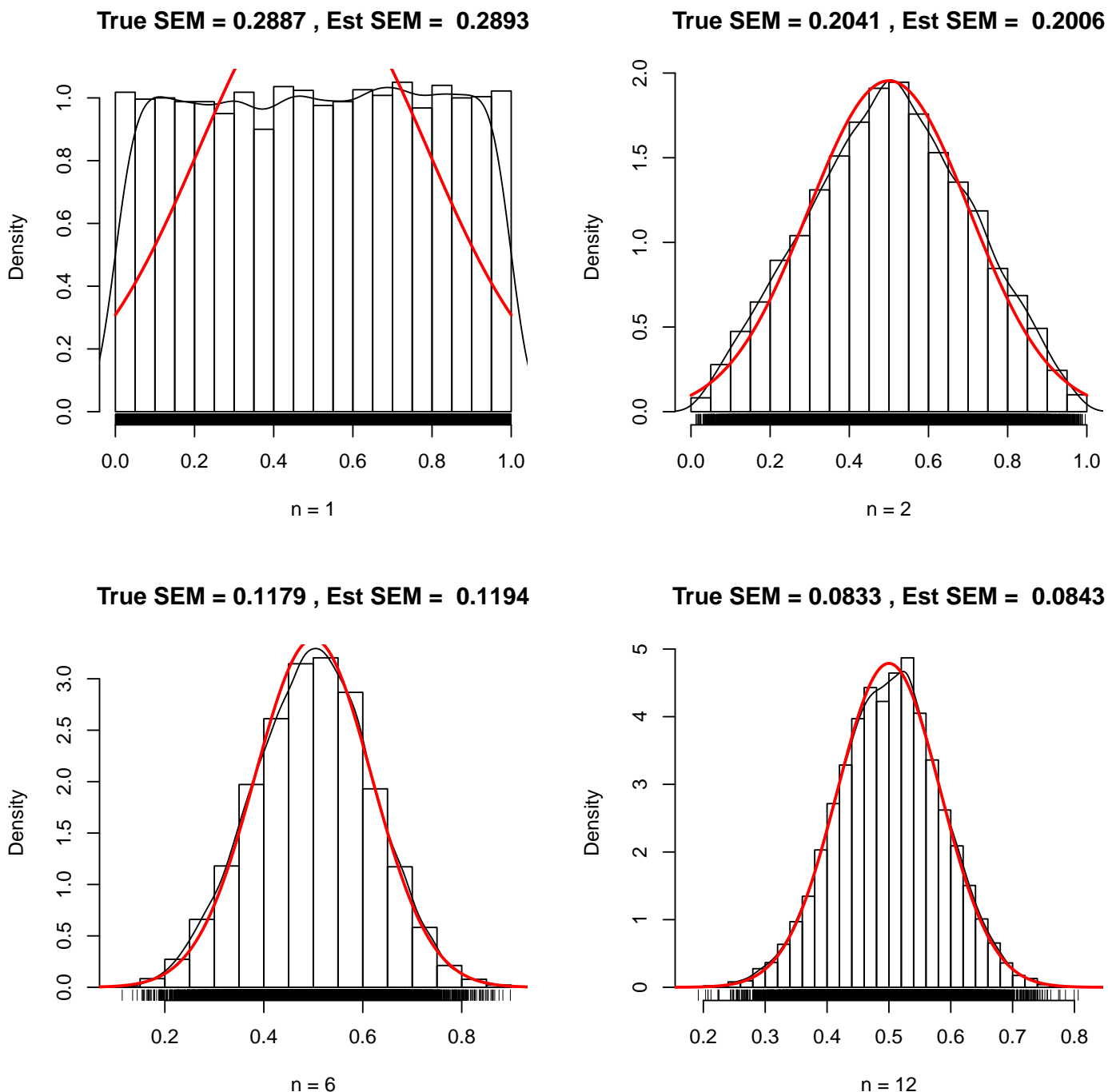
As a joint illustration of these concepts, consider drawing random variables following a Uniform(0,1) distribution, that is, any value in the interval $[0, 1]$ is equally likely. By definition, the mean of this distribution is $\mu = 1/2$ and the variance is $\sigma^2 = 1/12$ (so the standard deviation is $\sigma = \sqrt{1/12} = 0.289$). Therefore, if we draw a sample of size n , then the standard error of the mean will be σ/n , and as n gets larger the distribution of the mean will increasingly

¹The central limit theorem has a number of variants. In its common form, the random variables must be identically distributed. In variants, convergence of the mean to the normal distribution also occurs for non-identical distributions, given that they comply with certain conditions.

follow a normal distribution. We illustrate this by drawing $N = 10000$ samples of size n and plot those N means, computing the expected and observed SEM and how well the histogram of sampled means follows a normal distribution. Notice, indeed, that even with samples as small as 2 and 6 that the properties of the SEM and the distribution are as predicted.

```
#### Illustration of Central Limit Theorem, Uniform distribution
# demo.clt.unif(N, n)
#   draws N samples of size n from Uniform(0,1)
#   and plots the N means with a normal distribution overlay
demo.clt.unif <- function(N, n) {
  # draw sample in a matrix with N columns and n rows
  sam <- matrix(runif(N*n, 0, 1), ncol=N);
  # calculate the mean of each column
  sam.mean <- colMeans(sam)
  # the sd of the mean is the SEM
  sam.se <- sd(sam.mean)
  # calculate the true SEM given the sample size n
  true.se <- sqrt((1/12)/n)
  # draw a histogram of the means
  hist(sam.mean, freq = FALSE, breaks = 25
    , main = paste("True SEM =", round(true.se, 4)
      , ", Est SEM = ", round( sam.se, 4))
    , xlab = paste("n =", n))
  # overlay a density curve for the sample means
  points(density(sam.mean), type = "l")
  # overlay a normal distribution, bold and red
  x <- seq(0, 1, length = 1000)
  points(x, dnorm(x, mean = 0.5, sd = true.se), type = "l", lwd = 2, col = "red")
  # place a rug of points under the plot
  rug(sam.mean)
}

par(mfrow=c(2,2));
demo.clt.unif(10000, 1);
demo.clt.unif(10000, 2);
demo.clt.unif(10000, 6);
demo.clt.unif(10000, 12);
```

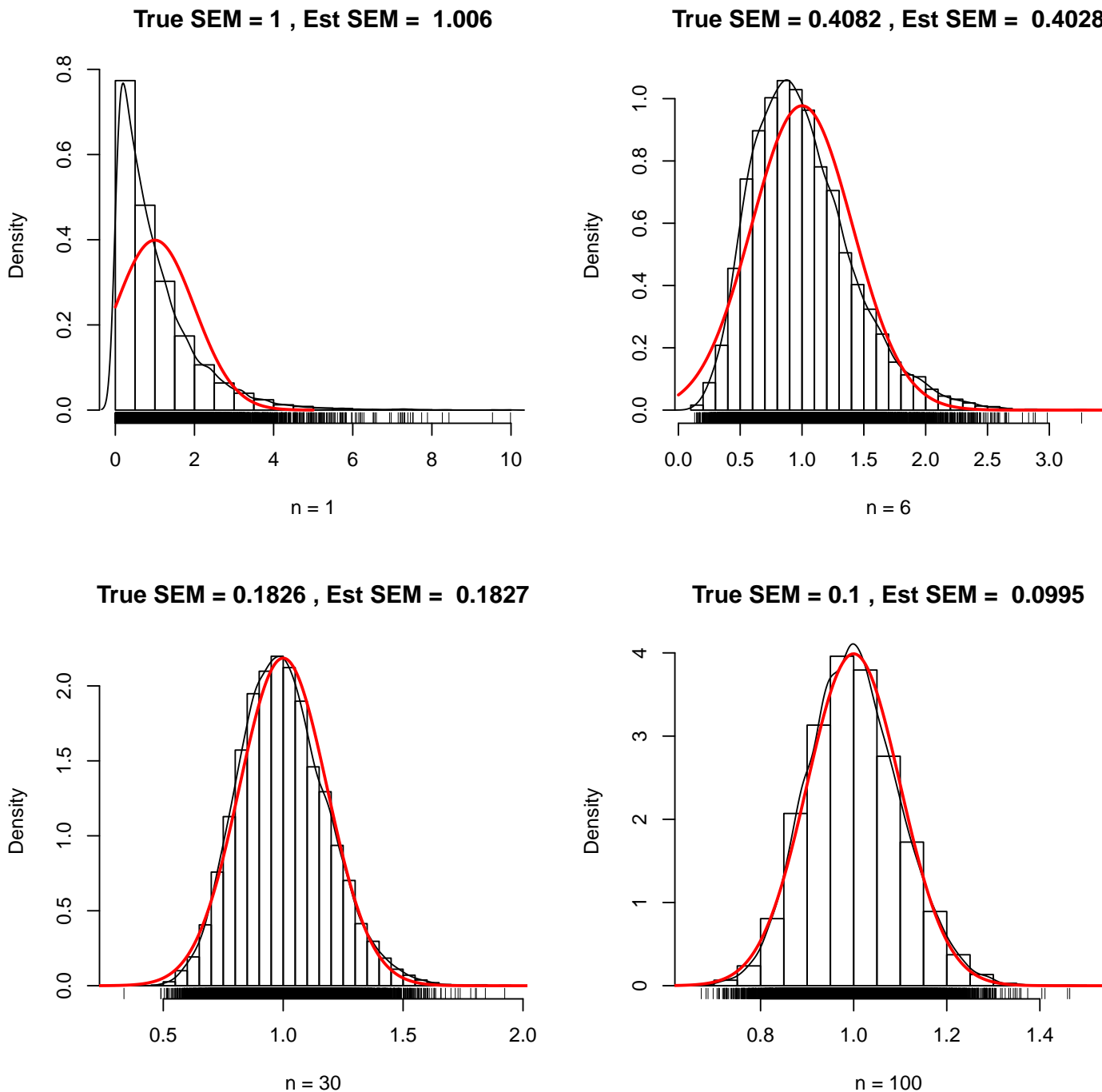


In a more extreme example, we draw samples from an $\text{Exponential}(1)$ distribution ($\mu = 1$ and $\sigma = 1$), which is strongly skewed to the right. Notice that the normality promised by the CLT requires larger samples sizes, about $n \geq 30$, than for the previous $\text{Uniform}(0,1)$ example, which required about $n \geq 6$.

```
#### Illustration of Central Limit Theorem, Exponential distribution
# demo.clt.exp(N, n) draws N samples of size n from Exponential(1)
# and plots the N means with a normal distribution overlay
demo.clt.exp <- function(N, n) {
  # draw sample in a matrix with N columns and n rows
  sam <- matrix(rexp(N*n, 1), ncol=N);
  # calculate the mean of each column
```

```
 sam.mean <- colMeans(sam)
# the sd of the mean is the SEM
sam.se <- sd(sam.mean)
# calculate the true SEM given the sample size n
true.se <- sqrt(1/n)
# draw a histogram of the means
hist(sam.mean, freq = FALSE, breaks = 25
    , main = paste("True SEM =", round(true.se, 4), ", Est SEM = ", round(sam.se, 4))
    , xlab = paste("n =", n))
# overlay a density curve for the sample means
points(density(sam.mean), type = "l")
# overlay a normal distribution, bold and red
x <- seq(0, 5, length = 1000)
points(x, dnorm(x, mean = 1, sd = true.se), type = "l", lwd = 2, col = "red")
# place a rug of points under the plot
rug(sam.mean)
}

par(mfrow=c(2,2));
demo.clt.exp(10000,    1);
demo.clt.exp(10000,    6);
demo.clt.exp(10000,   30);
demo.clt.exp(10000,  100);
```



Note well that the further the population distribution is from being normal, the larger the sample size is required to be for the sampling distribution of the sample mean to be normal. If the population distribution is normal, what's the minimum sample size for the sampling distribution of the mean to be normal?

For more examples, try:

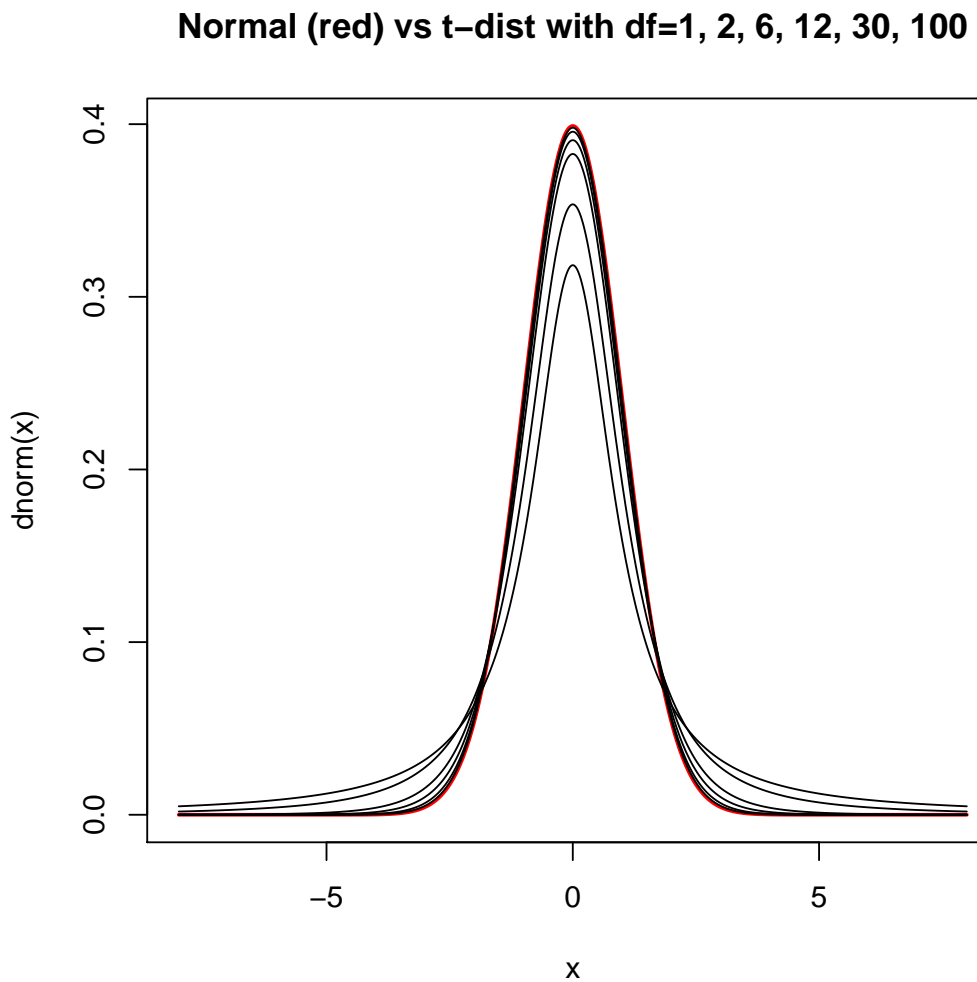
```
##### More examples for Central Limit Theorem can be illustrated with this code
# install.packages("TeachingDemos")
library(TeachingDemos)
# look at examples at bottom of the help page
```

```
?clt.examp
```

2.1.2 *t*-distribution

The Student's ***t*-distribution** is a family of continuous probability distributions that arises when estimating the mean of a normally distributed population in situations where the *sample size is small* and population *standard deviation is unknown*. The *t*-distribution is symmetric and bell-shaped, like the normal distribution, but has heavier tails, meaning that it is more prone to producing values that fall far from its mean. Effectively, the *t*-distribution is wider than the normal distribution because in addition to estimating the mean μ with \bar{Y} , we *also* have to estimate σ^2 with s^2 , so there's some additional uncertainty. The degrees-of-freedom (df) parameter of the *t*-distribution is the sample size n minus the number of variance parameters estimated. Thus, $df = n - 1$ when we have one sample and $df = n - 2$ when we have two samples. As n increases, the *t*-distribution becomes close to the normal distribution, and when $n = \infty$ the distributions are equivalent.

```
#### Normal vs t-distributions with a range of degrees-of-freedom
x <- seq(-8, 8, length = 1000)
par(mfrow=c(1,1))
plot(x, dnorm(x), type = "l", lwd = 2, col = "red"
      , main = "Normal (red) vs t-dist with df=1, 2, 6, 12, 30, 100")
points(x, dt(x, 1), type = "l")
points(x, dt(x, 2), type = "l")
points(x, dt(x, 6), type = "l")
points(x, dt(x, 12), type = "l")
points(x, dt(x, 30), type = "l")
points(x, dt(x,100), type = "l")
```



2.2 CI for μ

Statistical inference provides methods for drawing conclusions about a population from sample data. In this chapter, we want to make a claim about population mean μ given sample statistics \bar{Y} and s .

A CI for μ is a range of plausible values for the unknown population mean μ , based on the observed data, of the form “Best Guess \pm Reasonable Error of the Guess”. To compute a CI for μ :

1. Define the **population parameter**, “Let $\mu = \text{mean [characteristic]}$ for population of interest”.
2. Specify the **confidence coefficient**, which is a number between 0 and 100%, in the form $100(1 - \alpha)\%$. Solve for α . (For example, 95% has $\alpha = 0.05$.)

3. Compute the t -critical value: $t_{\text{crit}} = t_{0.5\alpha}$ such that the area under the t -curve ($df = n - 1$) to the right of t_{crit} is 0.5α . See appendix or internet for a t -table.
4. Report the CI in the form $\bar{Y} \pm t_{\text{crit}}SE_{\bar{Y}}$ or as an interval (L, U) . The desired CI has lower and upper endpoints given by $L = \bar{Y} - t_{\text{crit}}SE_{\bar{Y}}$ and $U = \bar{Y} + t_{\text{crit}}SE_{\bar{Y}}$, respectively, where $SE_{\bar{Y}} = s/\sqrt{n}$ is the standard error of the sample mean.
5. Assess method assumptions (see below).

In practice, the confidence coefficient is large, say 95% or 99%, which correspond to $\alpha = 0.05$ and 0.01, respectively. The value of α expressed as a percent is known as the **error rate** of the CI.

The CI is determined once the confidence coefficient is specified and the data are collected. Prior to collecting the data, the interval is unknown and is viewed as random because it will depend on the actual sample selected. Different samples give different CIs. The “**confidence**” in, say, the 95% CI (which has a 5% error rate) can be interpreted as follows. *If you repeatedly sample the population and construct 95% CIs for μ , then 95% of the intervals will contain μ , whereas 5% will not.* The interval you construct from your data will either cover μ , or it will not.

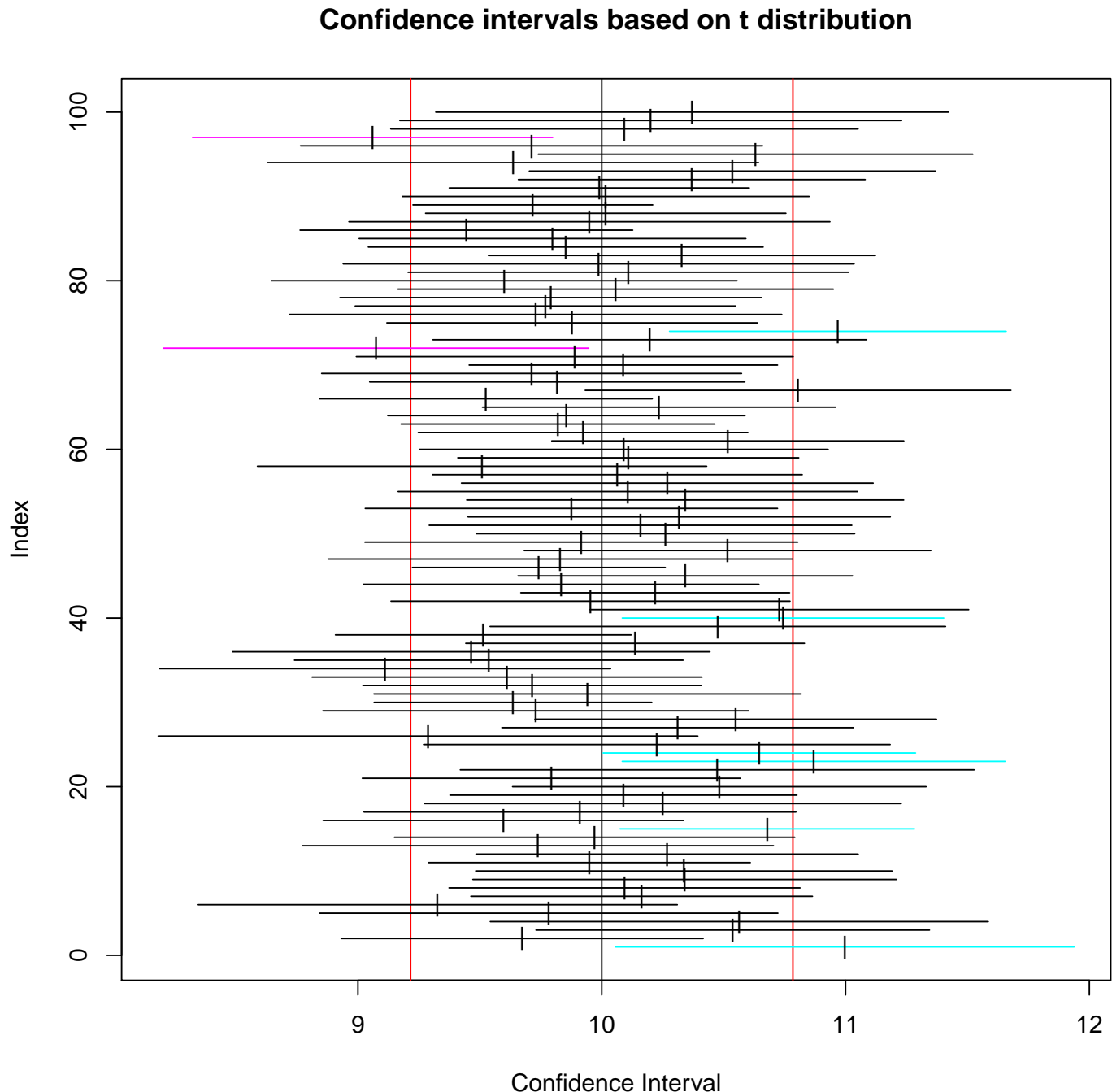
The length of the CI

$$U - L = 2t_{\text{crit}}SE_{\bar{Y}}$$

depends on the accuracy of our estimate \bar{Y} of μ , as measured by the standard error of \bar{Y} , $SE_{\bar{Y}} = s/\sqrt{n}$. Less precise estimates of μ lead to wider intervals for a given level of confidence.

An example with 100 CIs Consider drawing a sample of 25 observations from a normally distributed population with mean 10 and sd 2. Calculate the 95% t -CI. Now do that 100 times. The plot below reflects the variability of that process. We expect 95 of the 100 CIs to contain the true population mean of 10, that is, on average 5 times out of 100 we draw the incorrect inference that the population mean is in an interval when it does not contain the true value of 10.

```
#### Illustration of Confidence Intervals (consistent with their interpretation)
library(TeachingDemos)
ci.examp(mean.sim = 10, sd = 2, n = 25
         , reps = 100, conf.level = 0.95, method = "t")
```



2.2.1 Assumptions for procedures

I described the classical CI. The procedure is based on the assumptions that the data are a **random sample** from the population of interest, and that the **population frequency curve is normal**. The population frequency curve can be viewed as a “smoothed histogram” created from the population data.

The normality assumption can never be completely verified without having the entire population data. You can assess the reasonableness of this assumption using a stem-and-leaf display or a boxplot of the sample data. The stem-and-leaf display from the data should resemble a normal curve.

In fact, the assumptions are slightly looser than this, the population frequency curve can be anything provided the sample size is large enough that it's reasonable to assume that the **sampling distribution of the mean is normal**.

Assessing assumptions using the bootstrap

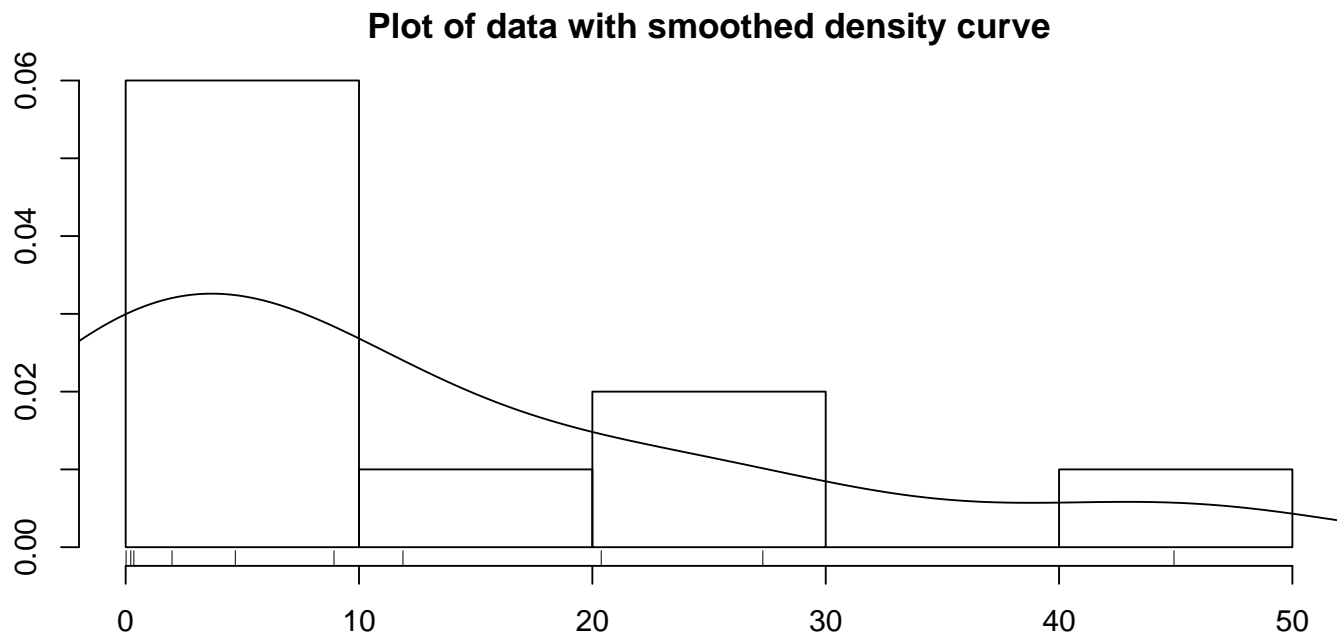
We will cover the bootstrap at the end of this course, but a brief introduction here can help us check model assumptions. Recall in Section 2.1.1 the sampling distribution examples. Assume the sample is representative of the population. Let's use our sample as a proxy for the population and repeatedly draw samples (with replacement) of size n and calculate the mean, then plot the bootstrap sampling distribution of means. If this bootstrap sampling distribution strongly deviates from normal, then that's evidence from the data that inference using the t -distribution is not appropriate. Otherwise, if roughly normal, then the t -distribution may be sensible.

```
#### Visual comparison of whether sampling distribution is close to Normal via Bootstrap
# a function to compare the bootstrap sampling distribution with
#   a normal distribution with mean and SEM estimated from the data
bs.one.samp.dist <- function(dat, N = 1e4) {
  n <- length(dat);
  # resample from data
  sam <- matrix(sample(dat, size = N * n, replace = TRUE), ncol=N);
  # draw a histogram of the means
  sam.mean <- colMeans(sam);
  # save par() settings
  old.par <- par(no.readonly = TRUE)
  # make smaller margins
  par(mfrow=c(2,1), mar=c(3,2,2,1), oma=c(1,1,1,1))
```

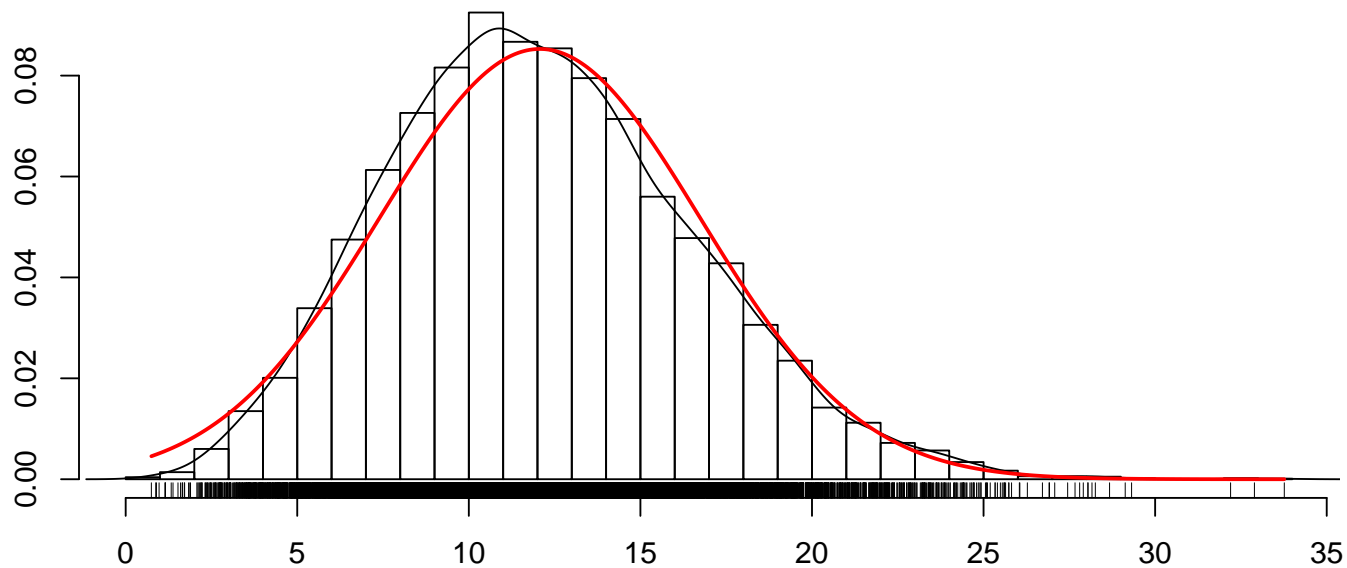
```
# Histogram overlaid with kernel density curve
hist(dat, freq = FALSE, breaks = 6
      , main = "Plot of data with smoothed density curve")
points(density(dat), type = "l")
rug(dat)

hist(sam.mean, freq = FALSE, breaks = 25
      , main = "Bootstrap sampling distribution of the mean"
      , xlab = paste("Data: n =", n
                     , ", mean =", signif(mean(dat), digits = 5)
                     , ", se =", signif(sd(dat)/sqrt(n)), digits = 5))
# overlay a density curve for the sample means
points(density(sam.mean), type = "l")
# overlay a normal distribution, bold and red
x <- seq(min(sam.mean), max(sam.mean), length = 1000)
points(x, dnorm(x, mean = mean(dat), sd = sd(dat)/sqrt(n))
       , type = "l", lwd = 2, col = "red")
# place a rug of points under the plot
rug(sam.mean)
# restore par() settings
par(old.par)
}

# example data, skewed --- try others datasets to develop your intuition
x <- rgamma(10, shape = .5, scale = 20)
bs.one.samp.dist(x)
```



Bootstrap sampling distribution of the mean



Example: Age at First Heart Transplant Let us go through a hand-calculation of a CI, using R to generate summary data. I'll show you later how to generate the CI in R.

We are interested in the mean age at first heart transplant for a population of patients.

1. Define the population parameter

Let μ = mean age at the time of first heart transplant for population of

patients.

2. Calculate summary statistics from sample

The ages (in years) at first transplant for a sample of 11 heart transplant patients are as follows:

$$54, 42, 51, 54, 49, 56, 33, 58, 54, 64, 49.$$

Summaries for the data are: $n = 11$, $\bar{Y} = 51.27$, and $s = 8.26$ so that $SE_{\bar{Y}} = 8.26/\sqrt{11} = 2.4904$. The degrees of freedom are $df = 11 - 1 = 10$.

3. Specify confidence level, find critical value, calculate limits

Let us calculate a 95% CI for μ . For a 95% CI $\alpha = 0.05$, so we need to find $t_{\text{crit}} = t_{0.025}$, which is 2.228. Now $t_{\text{crit}}SE_{\bar{Y}} = 2.228 \times 2.4904 = 5.55$. The lower limit on the CI is $L = 51.27 - 5.55 = 45.72$. The upper limit is $U = 51.27 + 5.55 = 56.82$.

4. Summarize in words For example, I am 95% confident that the population mean age at first transplant is 51.3 ± 5.55 , that is, between 45.7 and 56.8 years (rounding off to 1 decimal place).

5. Check assumptions We will see this in several pages, sampling distribution is reasonably normal.

2.2.2 The effect of α on a two-sided CI

A two-sided $100(1 - \alpha)\%$ CI for μ is given by $\bar{Y} \pm t_{\text{crit}}SE_{\bar{Y}}$. The CI is centered at \bar{Y} and has length $2t_{\text{crit}}SE_{\bar{Y}}$. The confidence coefficient $100(1 - \alpha)\%$ is **increased** by **decreasing** α , which increases t_{crit} . That is, increasing the confidence coefficient makes the CI wider. This is sensible: to increase your confidence that the interval captures μ you must pinpoint μ with less precision by making the CI wider. For example, a 95% CI is wider than a 90% CI.



CLICKER Qs — CI, quickie



2.3 Hypothesis Testing for μ

A **hypothesis test** is used to make a **decision** about a population parameter.

Suppose you are interested in checking whether the population mean μ is equal to some prespecified value, say μ_0 . This question can be formulated as a two-sided hypothesis test, where you are trying to decide which of two contradictory claims or hypotheses about μ is more reasonable given the observed data. The **null hypothesis**, or the hypothesis under test, is $H_0 : \mu = \mu_0$, whereas the **alternative hypothesis** is $H_A : \mu \neq \mu_0$.

I will explore the ideas behind hypothesis testing later. At this point, I focus on the mechanics behind the test. The steps in carrying out the test are:

1. Set up the **null and alternative hypotheses** in words and notation.

In words: “The population mean for [what is being studied] is different from [value of μ_0].” (Note that the statement in words is in terms of the alternative hypothesis.)

In notation: $H_0 : \mu = \mu_0$ versus $H_A : \mu \neq \mu_0$ (where μ_0 is specified by the context of the problem).

2. Choose the **size** or **significance level** of the test, denoted by α . In practice, α is set to a small value, say, 0.01 or 0.05, but theoretically can be any value between 0 and 1.
3. Compute the **test statistic**

$$t_s = \frac{\bar{Y} - \mu_0}{SE_{\bar{Y}}},$$

where $SE_{\bar{Y}} = s/\sqrt{n}$ is the standard error.

Note: I sometimes call the test statistic t_{obs} to emphasize that the computed value depends on the observed data.

4. Compute the **critical value** $t_{\text{crit}} = t_{0.5\alpha}$ (or p -value from the test statistic) in the direction of the alternative hypothesis from the t -distribution table with degrees of freedom $df = n - 1$.
5. State the **conclusion** in terms of the problem.

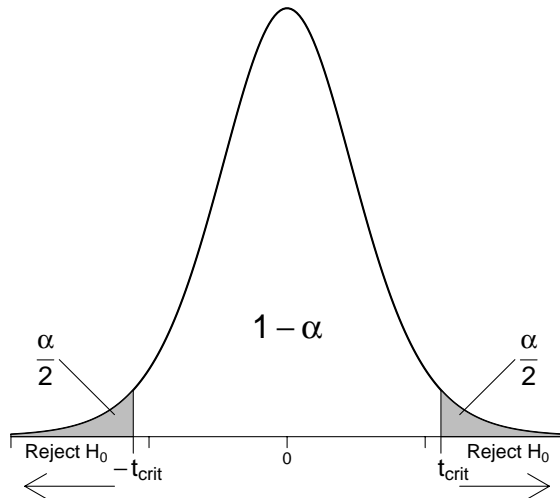
Reject H_0 in favor of H_A (i.e., decide that H_0 is false, based on the data) if $|t_s| > t_{\text{crit}}$ or p -value $< \alpha$, that is, reject if $t_s < -t_{\text{crit}}$ or if $t_s > t_{\text{crit}}$.

Otherwise, **Fail to reject** H_0 .

(Note: We DO NOT *accept* H_0 — more on this later.)

6. **Check assumptions** of the test, when possible (could do earlier to save yourself some effort if they are not met).

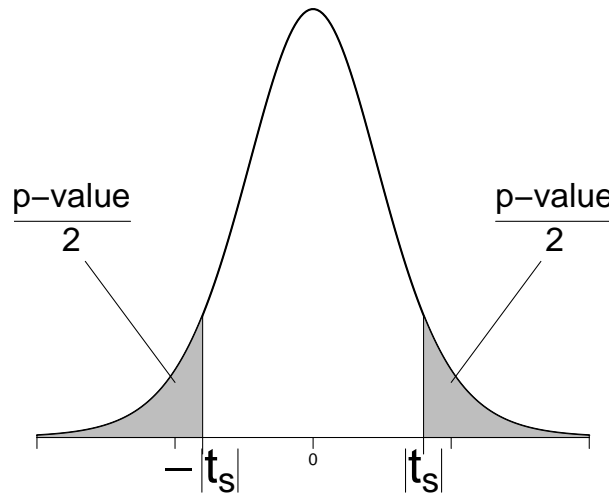
The process is represented graphically below. The area under the t -probability curve outside $\pm t_{\text{crit}}$ is the size of the test, α . One-half α is the area in each tail. You reject H_0 in favor of H_A only if the test statistic is outside $\pm t_{\text{crit}}$.



2.3.1 P-values

The **p-value**, or **observed significance level** for the test, provides a measure of plausibility for H_0 . Smaller values of the p-value imply that H_0 is less plausible. To compute the p-value for a two-sided test, you

1. Compute the test statistic t_s as above.
2. Evaluate the area under the t -probability curve (with $df = n - 1$) outside $\pm |t_s|$.



The p-value is the total shaded area, or twice the area in either tail. A useful **interpretation** of the p-value is that *it is the chance of obtaining data favoring H_A by this much or more if H_0 actually is true*. Another interpretation is that

the **p-value** is *the probability of observing a sample mean at least as extreme as the one observed assuming μ_0 from H_0 is the true population mean.*

If the p-value is small then the sample we obtained is pretty unusual to have obtained if H_0 is true — but we actually got the sample, so probably it is not very unusual, so we would conclude H_0 is false (it would not be unusual if H_A is true).

Most, if not all, statistical packages summarize hypothesis tests with a p-value, rather than a decision (i.e., reject or not reject at a given α level). You can make a decision to reject or not reject H_0 for a size α test based on the p-value as follows — *reject H_0 if the p-value is less than α* . This decision is identical to that obtained following the formal rejection procedure given earlier. The reason for this is that the p-value can be interpreted as the smallest value you can set the size of the test and still reject H_0 given the observed data.

There are a lot of terms to keep straight here. α and t_{crit} are constants we choose (actually, one determines the other so we really only choose one,

usually α) to set how rigorous evidence against H_0 needs to be. t_s and the p-value (again, one determines the other) are random variables because they are calculated from the random sample. They are the evidence against H_0 .

2.3.2 Assumptions for procedures

I described the classical t -test, which assumes that the data are a random sample from the population and that the population frequency curve is normal. These are the same assumptions as for the CI.

Example: Age at First Transplant (Revisited) The ages (in years) at first transplant for a sample of 11 heart transplant patients are as follows: 54, 42, 51, 54, 49, 56, 33, 58, 54, 64, 49. Summaries for these data are: $n = 11$, $\bar{Y} = 51.27$, $s = 8.26$ and $SE_{\bar{Y}} = 2.4904$. Test the hypothesis that the mean age at first transplant is 50. Use $\alpha = 0.05$.

As in the earlier analysis, define

μ = mean age at time of first transplant for population of patients.

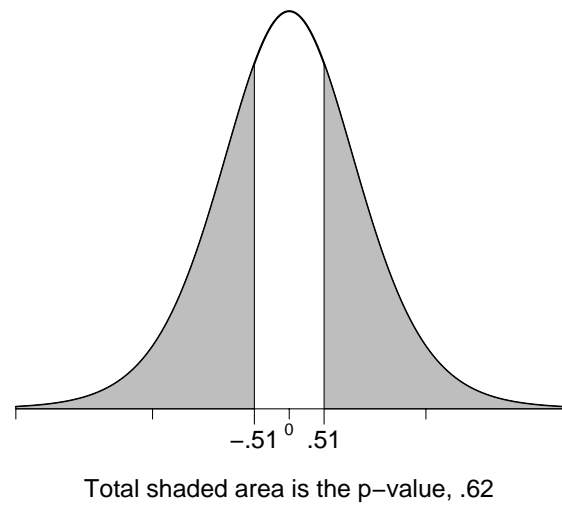
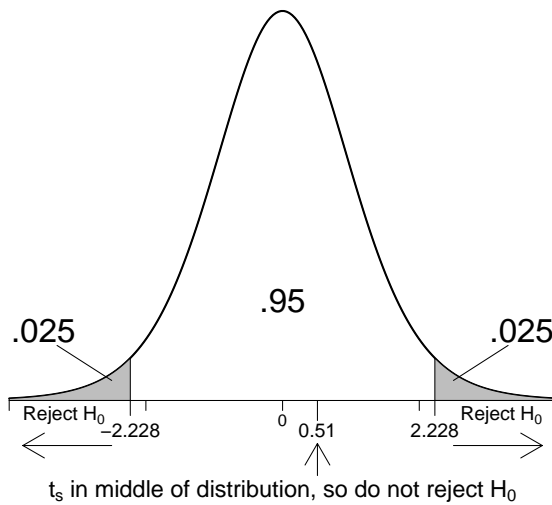
We are interested in testing $H_0 : \mu = 50$ against $H_A : \mu \neq 50$, so $\mu_0 = 50$.

The degrees of freedom are $df = 11 - 1 = 10$. The critical value for a 5% test is $t_{\text{crit}} = t_{0.025} = 2.228$. (Note $\alpha/2 = 0.05/2 = 0.025$). The same critical value was used with the 95% CI.

For the test,

$$t_s = \frac{\bar{Y} - \mu_0}{SE_{\bar{Y}}} = \frac{51.27 - 50}{2.4904} = 0.51.$$

Since $t_{\text{crit}} = 2.228$, we do not reject H_0 using a 5% test. Notice the placement of t_s relative to t_{crit} in the picture below. Equivalently, the p-value for the test is 0.62, thus we fail to reject H_0 because $0.62 > 0.05 = \alpha$. The results of the hypothesis test should not be surprising, since the CI tells you that 50 is a plausible value for the population mean age at transplant. Note: All you can say is that the data *could have* come from a distribution with a mean of 50 — this is not convincing evidence that μ actually *is* 50.



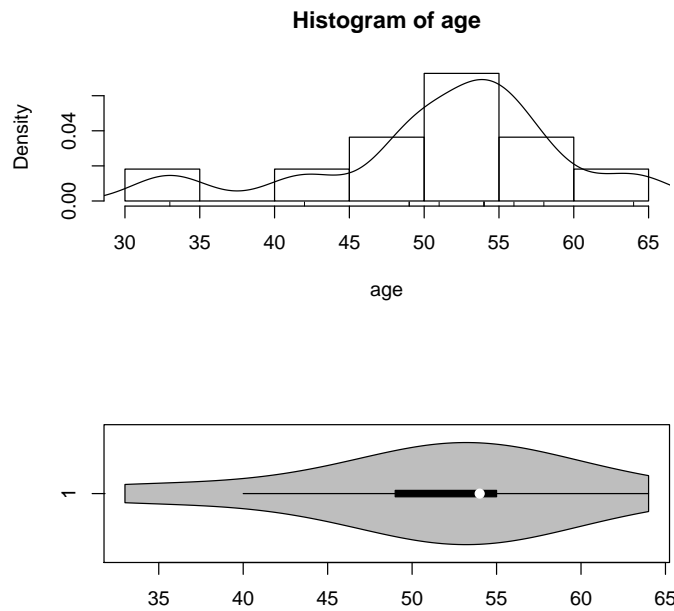
Example: Age at First Transplant R output for the heart transplant problem is given below. Let us look at the output and find all of the summaries we computed. Also, look at the graphical summaries to assess whether the *t*-test and CI are reasonable here.

```
#### Example: Age at First Transplant
# enter data as a vector
age <- c(54, 42, 51, 54, 49, 56, 33, 58, 54, 64, 49)
```

The age data is unimodal, skewed left, no extreme outliers.

```
par(mfrow=c(2,1))
# Histogram overlaid with kernel density curve
hist(age, freq = FALSE, breaks = 6)
points(density(age), type = "l")
rug(age)

# violin plot
library(vioplot)
vioplot(age, horizontal=TRUE, col="gray")
```



```
# stem-and-leaf plot
stem(age, scale=2)

##
##      The decimal point is 1 digit(s) to the right of the |
##
##      3 | 3
##      3 |
##      4 | 2
##      4 | 99
##      5 | 1444
##      5 | 68
##      6 | 4

# t.crit
qt(1 - 0.05/2, df = length(age) - 1)
## [1] 2.228

# look at help for t.test
?t.test
# defaults include: alternative = "two.sided", conf.level = 0.95
t.summary <- t.test(age, mu = 50)
t.summary

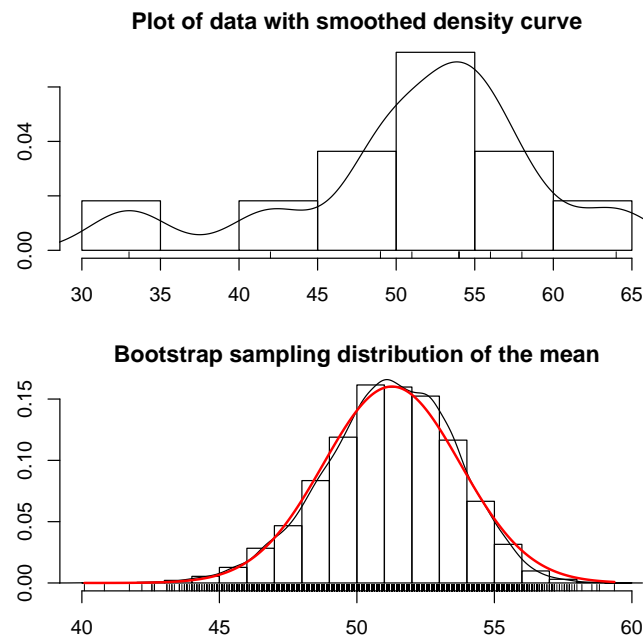
##
##  One Sample t-test
##
## data: age
## t = 0.5111, df = 10, p-value = 0.6204
## alternative hypothesis: true mean is not equal to 50
## 95 percent confidence interval:
##  45.72 56.82
## sample estimates:
```

```
## mean of x
##      51.27

summary(age)
##    Min. 1st Qu. Median     Mean 3rd Qu.     Max.
##    33.0    49.0    54.0    51.3    55.0    64.0
```

The assumption of normality of the sampling distribution appears reasonably close, using the bootstrap discussed earlier. Therefore, the results for the t -test above can be trusted.

```
bs.one.samp.dist(age)
```



Aside: To print the shaded region for the p-value, you can use the result of `t.test()` with the function `t.dist.pval()` defined here.

```
# Function to plot t-distribution with shaded p-value
t.dist.pval <- function(t.summary) {
  par(mfrow=c(1,1))
  lim.extreme <- max(4, abs(t.summary$statistic) + 0.5)
  lim.lower <- -lim.extreme;
  lim.upper <- lim.extreme;
  x.curve <- seq(lim.lower, lim.upper, length=200)
  y.curve <- dt(x.curve, df = t.summary$parameter)
  plot(x.curve, y.curve, type = "n"
       , ylab = paste("t-dist( df =", signif(t.summary$parameter, 3), ")"))
       , xlab = paste("t-stat =", signif(t.summary$statistic, 5)
                      , ", Shaded area is p-value =", signif(t.summary$p.value, 5)))
  if ((t.summary$alternative == "less")
      | (t.summary$alternative == "two.sided")) {
    x.pval.l <- seq(lim.lower, -abs(t.summary$statistic), length=200)
    y.pval.l <- dt(x.pval.l, df = t.summary$parameter)
    polygon(c(lim.lower, x.pval.l, -abs(t.summary$statistic))
            , c(0, y.pval.l, 0), col="gray")}
```

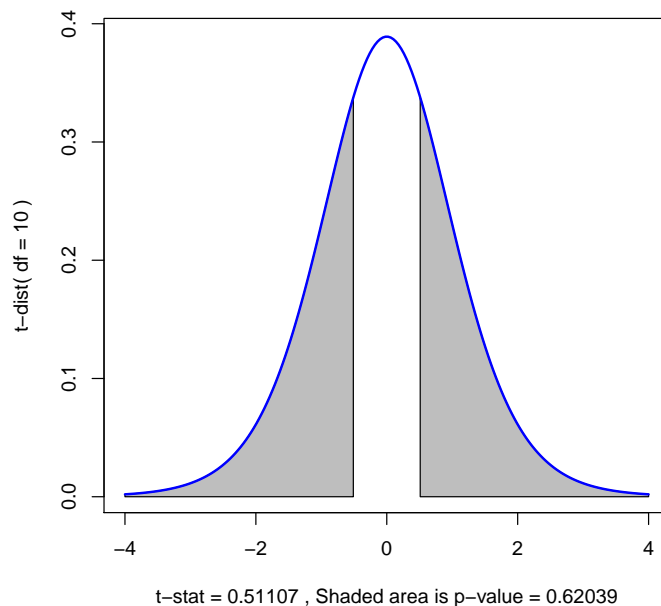
```

}

if ((t.summary$alternative == "greater")
  | (t.summary$alternative == "two.sided")) {
  x.pval.u <- seq(abs(t.summary$statistic), lim.upper, length=200)
  y.pval.u <- dt(x.pval.u, df = t.summary$parameter)
  polygon(c(abs(t.summary$statistic), x.pval.u, lim.upper)
    , c(0, y.pval.u, 0), col="gray")
}
points(x.curve, y.curve, type = "l", lwd = 2, col = "blue")
}

# for the age example
t.dist.pval(t.summary)

```



Aside: Note that the `t.summary` object returned from `t.test()` includes a number of quantities that might be useful for additional calculations.

```

names(t.summary)
## [1] "statistic"      "parameter"      "p.value"        "conf.int"
## [5] "estimate"       "null.value"     "alternative"   "method"
## [9] "data.name"

t.summary$statistic
##      t
## 0.5111

t.summary$parameter
## df
## 10

t.summary$p.value
## [1] 0.6204

t.summary$conf.int
## [1] 45.72 56.82
## attr(,"conf.level")
## [1] 0.95

t.summary$estimate
## mean of x
##      51.27

t.summary>null.value

```

```

## mean
## 50
t.summary$alternative
## [1] "two.sided"
t.summary$method
## [1] "One Sample t-test"
t.summary$data.name
## [1] "age"

```

Example: Meteorites One theory of the formation of the solar system states that all solar system meteorites have the same evolutionary history and thus have the same cooling rates. By a delicate analysis based on measurements of phosphide crystal widths and phosphide-nickel content, the cooling rates, in degrees Celsius per million years, were determined for samples taken from meteorites named in the accompanying table after the places they were found. The Walker² County (Alabama, US), Uwet³ (Cross River, Nigeria), and Tocopilla⁴ (Antofagasta, Chile) meteorite cooling rate data are below.

Suppose that a hypothesis of solar evolution predicted a mean cooling rate of $\mu = 0.54$ degrees per million year for the Tocopilla meteorite. Do the observed cooling rates support this hypothesis? Test at the 5% level. The boxplot and stem-and-leaf display (given below) show good symmetry. The assumption of a normal distribution of observations basic to the *t*-test appears to be realistic.

Meteorite	Cooling rates
Walker County	0.69 0.23 0.10 0.03 0.56 0.10 0.01 0.02 0.04 0.22
Uwet	0.21 0.25 0.16 0.23 0.47 1.20 0.29 1.10 0.16
Tocopilla	5.60 2.70 6.20 2.90 1.50 4.00 4.30 3.00 3.60 2.40 6.70 3.80

Let

μ = mean cooling rate over all pieces of the Tocopilla meteorite.

To answer the question of interest, we consider the test of $H_0 : \mu = 0.54$ against $H_A : \mu \neq 0.54$. Let us go carry out the test, compute the p-value, and calculate a 95% CI for μ . The sample summaries are $n = 12$, $\bar{Y} = 3.892$, $s = 1.583$. The standard error is $SE_{\bar{Y}} = s/\sqrt{n} = 0.457$.

²<http://www.lpi.usra.edu/meteor/metbull.php?code=24204>

³<http://www.lpi.usra.edu/meteor/metbull.php?code=24138>

⁴<http://www.lpi.usra.edu/meteor/metbull.php?code=17001>

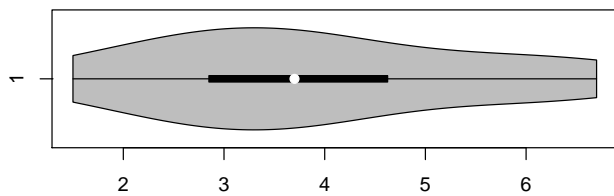
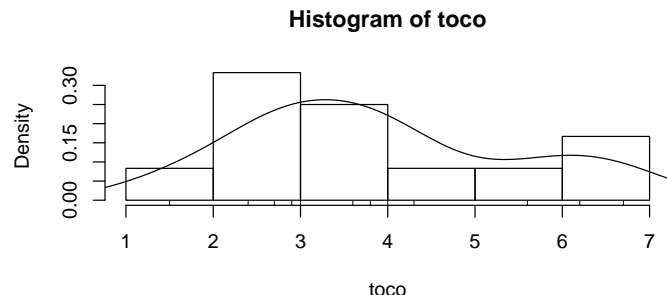
R output for this problem is given below. For a 5% test (i.e., $\alpha = 0.05$), you would reject H_0 in favor of H_A because the p-value ≤ 0.05 . The data strongly suggest that $\mu \neq 0.54$. The 95% CI says that you are 95% confident that the population mean cooling rate for the Tocopilla meteorite is between 2.89 and 4.90 degrees per million years. Note that the CI gives us a means to assess how different μ is from the hypothesized value of 0.54.

```
#### Example: Meteorites
# enter data as a vector
toco <- c(5.6, 2.7, 6.2, 2.9, 1.5, 4.0, 4.3, 3.0, 3.6, 2.4, 6.7, 3.8)
```

The Tocopilla data is unimodal, skewed right, no extreme outliers.

```
par(mfrow=c(2,1))
# Histogram overlaid with kernel density curve
hist(toco, freq = FALSE, breaks = 6)
points(density(toco), type = "l")
rug(toco)

# violin plot
library(vioplot)
vioplot(toco, horizontal=TRUE, col="gray")
```



```
# stem-and-leaf plot
stem(toco, scale=2)

##
##   The decimal point is at the |
##
##   1 | 5
##   2 | 479
##   3 | 068
##   4 | 03
```

```

##   5 | 6
##   6 | 27

# t.crit
qt(1 - 0.05/2, df = length(toco) - 1)
## [1] 2.201

t.summary <- t.test(toco, mu = 0.54)
t.summary

##
##  One Sample t-test
##
## data: toco
## t = 7.337, df = 11, p-value = 1.473e-05
## alternative hypothesis: true mean is not equal to 0.54
## 95 percent confidence interval:
##  2.886 4.897
## sample estimates:
## mean of x
##      3.892

summary(toco)
##    Min. 1st Qu. Median     Mean 3rd Qu. Max.
##    1.50    2.85    3.70    3.89    4.62   6.70

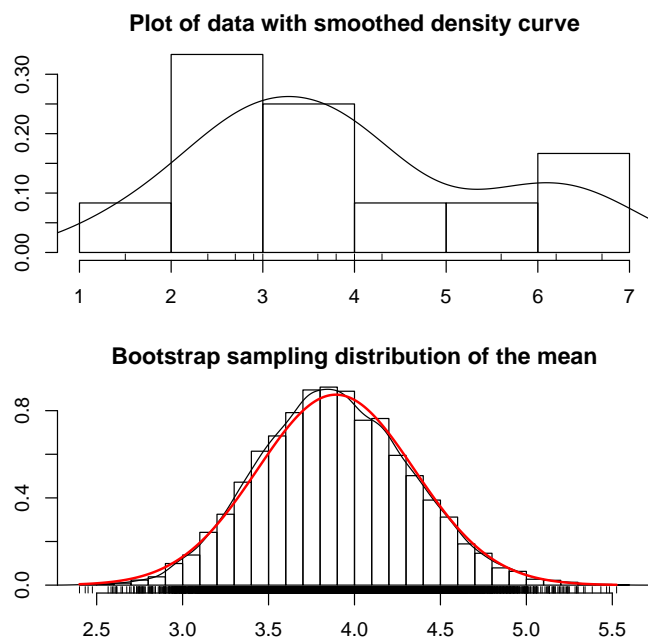
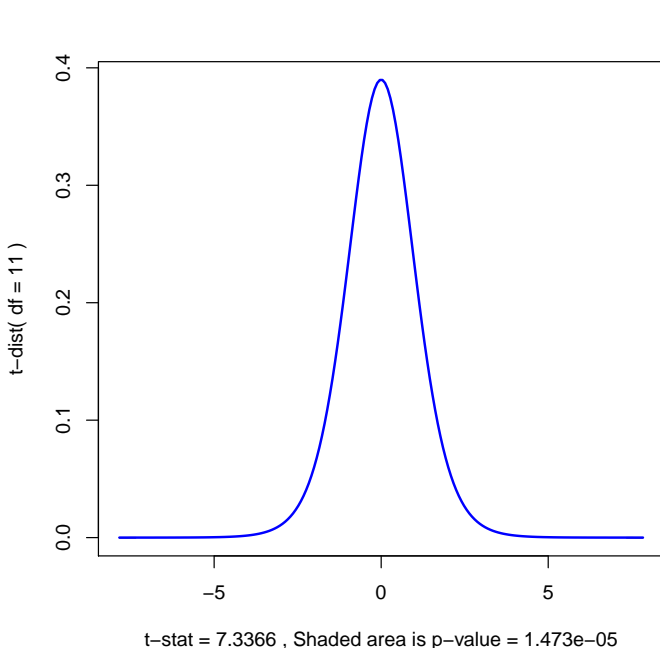
```

The assumption of normality of the sampling distribution appears reasonable. Therefore, the results for the t -test above can be trusted.

```

t.dist.pval(t.summary)
bs.one.samp.dist(toco)

```



2.3.3 The mechanics of setting up hypothesis tests

When setting up a test you should imagine you are the researcher conducting the experiment. In many studies, the researcher wishes to establish that there has been a change from the **status quo**, or that they have developed a method that produces a **change** (possibly in a specified direction) in the typical response. The researcher sets H_0 to be the **status quo** and H_A to be the **research hypothesis** — the claim the researcher wishes to make. In some studies you define the hypotheses so that H_A is the **take action** hypothesis — rejecting H_0 in favor of H_A leads one to take a radical action.

Some perspective on testing is gained by understanding the mechanics behind the tests. A hypothesis test is a decision process in the face of uncertainty. You are given data and asked which of two contradictory claims about a population parameter, say μ , is more reasonable. Two decisions are possible, but whether you make the correct decision depends on the true state of nature which is unknown to you.

Decision	State of nature	
	H_0 true	H_A true
Fail to reject [accept] H_0	correct decision	<i>Type-II error</i>
Reject H_0 in favor of H_A	<i>Type-I error</i>	correct decision

For a given problem, only one of these errors is possible. For example, if H_0 is true you can make a Type-I error but not a Type-II error. Any reasonable decision rule based on the data that tells us when to reject H_0 and when to not reject H_0 will have a certain probability of making a Type-I error if H_0 is true, and a corresponding probability of making a Type-II error if H_0 is false and H_A is true. For a given decision rule, define

$$\alpha = \text{Prob(Reject } H_0 \text{ given } H_0 \text{ is true)} = \text{Prob(Type-I error)}$$

and

$$\beta = \text{Prob(Fail to reject } H_0 \text{ when } H_A \text{ true)} = \text{Prob(Type-II error).}$$

The mathematics behind hypothesis tests allows you to prespecify or control α . For a given α , the tests we use (typically) have the smallest possible value of β . Given the researcher can control α , you set up the hypotheses so that committing a Type-I error is more serious than committing a Type-II error. The magnitude of α , also called the **size** or **level** of the test, should depend

on the seriousness of a Type-I error in the given problem. The more serious the consequences of a Type-I error, the smaller α should be. In practice α is often set to 0.10, 0.05, or 0.01, with $\alpha = 0.05$ being the scientific standard. By setting α to be a small value, you reject H_0 in favor of H_A only if the data **convincingly indicate** that H_0 is false.

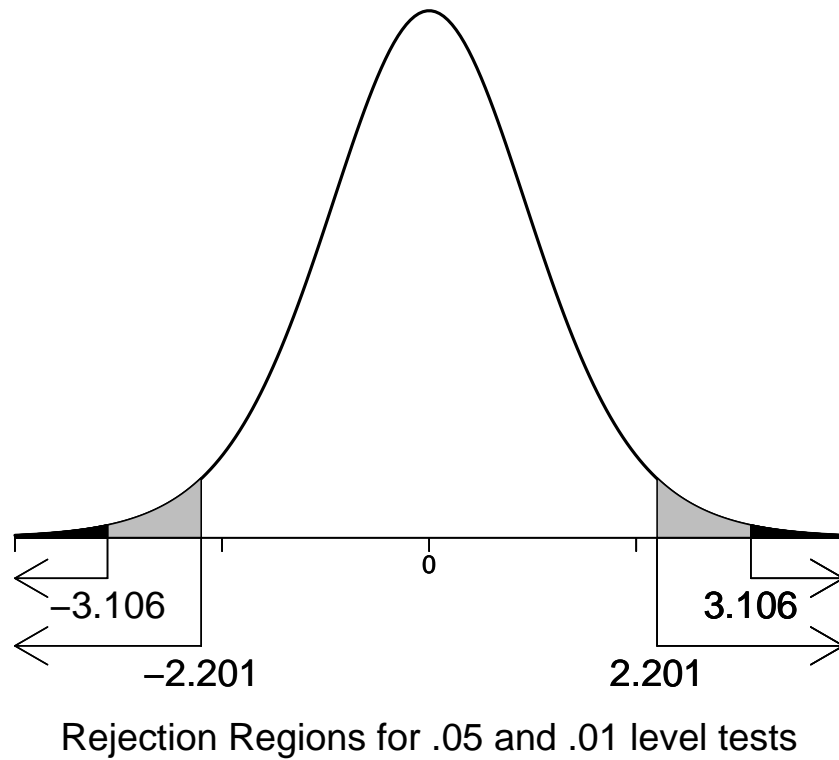
Let us piece together these ideas for the meteorite problem. Evolutionary history predicts $\mu = 0.54$. A scientist examining the validity of the theory is trying to decide whether $\mu = 0.54$ or $\mu \neq 0.54$. Good scientific practice dictates that rejecting another's claim when it is true is more serious than not being able to reject it when it is false. This is consistent with defining $H_0 : \mu = 0.54$ (the status quo) and $H_A : \mu \neq 0.54$. To convince yourself, note that the implications of a Type-I error would be to claim the evolutionary theory is false when it is true, whereas a Type-II error would correspond to not being able to refute the evolutionary theory when it is false. With this setup, the scientist will refute the theory only if the data overwhelmingly suggest that it is false.

2.3.4 The effect of α on the rejection region of a two-sided test

For a size α test, you reject $H_0 : \mu = \mu_0$ if

$$t_s = \frac{\bar{Y} - \mu_0}{SE_{\bar{Y}}}$$

satisfies $|t_s| > t_{\text{crit.}}$



The critical value is computed so that the area under the t -probability curve (with $df = n - 1$) outside $\pm t_{\text{crit}}$ is α , with 0.5α in each tail. Reducing α makes t_{crit} larger. That is, reducing the size of the test makes rejecting H_0 harder because the rejection region is smaller. A pictorial representation is given above for the Tocopilla data, where $\mu_0 = 0.54$, $n = 12$, and $df = 11$. Note that $t_{\text{crit}} = 2.201$ and 3.106 for $\alpha = 0.05$ and 0.01 , respectively.

The mathematics behind the test presumes that H_0 is true. Given the data, you use

$$t_s = \frac{\bar{Y} - \mu_0}{SE_{\bar{Y}}}$$

to measure how far \bar{Y} is from μ_0 , relative to the spread in the data given by $SE_{\bar{Y}}$. For t_s to be in the rejection region, \bar{Y} must be significantly above or below μ_0 , relative to the spread in the data. To see this, note that rejection

rule can be expressed as: **Reject** H_0 if

$$\bar{Y} < \mu_0 - t_{\text{crit}}SE_{\bar{Y}} \quad \text{or} \quad \bar{Y} > \mu_0 + t_{\text{crit}}SE_{\bar{Y}}.$$

The rejection rule is sensible because \bar{Y} is our best guess for μ . You would reject $H_0 : \mu = \mu_0$ only if \bar{Y} is so far from μ_0 that you would question the reasonableness of assuming $\mu = \mu_0$. How far \bar{Y} must be from μ_0 before you reject H_0 depends on α (i.e., how willing you are to reject H_0 if it is true), and on the value of $SE_{\bar{Y}}$. For a given sample, reducing α forces \bar{Y} to be further from μ_0 before you reject H_0 . For a given value of α and s , increasing n allows smaller differences between \bar{Y} and μ_0 to be **statistically significant** (i.e., lead to rejecting H_0). In problems where small differences between \bar{Y} and μ_0 lead to rejecting H_0 , you need to consider whether the observed differences are important.

In essence, the t -distribution provides an objective way to calibrate whether the observed \bar{Y} is typical of what sample means look like when sampling from a normal population where H_0 is true. If all other assumptions are satisfied, and \bar{Y} is inordinately far from μ_0 , then our only recourse is to conclude that H_0 must be incorrect.

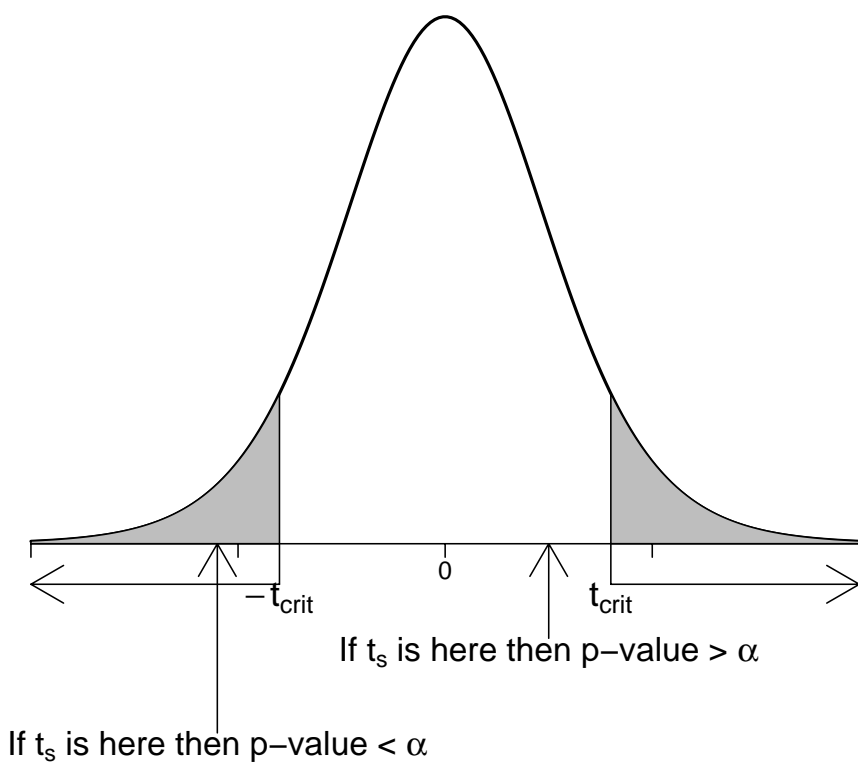
2.4 Two-sided tests, CI and p-values

An important relationship among two-sided tests of $H_0 : \mu = \mu_0$, CI, and p-values is that

$$\begin{aligned} \text{size } \alpha \text{ test rejects } H_0 &\Leftrightarrow 100(1 - \alpha)\% \text{ CI does not contain} \\ &\mu_0 \Leftrightarrow \text{p-value} \leq \alpha, \text{ and} \end{aligned}$$

$$\text{size } \alpha \text{ test fails to reject } H_0 \Leftrightarrow 100(1 - \alpha)\% \text{ CI contains } \mu_0 \Leftrightarrow \text{p-value} > \alpha.$$

For example, an $\alpha = 0.05$ test rejects $H_0 \Leftrightarrow 95\%$ CI does not contain $\mu_0 \Leftrightarrow \text{p-value} \leq 0.05$. The picture below illustrates the connection between p-values and rejection regions.



Either a CI or a test can be used to decide the plausibility of the claim that $\mu = \mu_0$. Typically, you use the test to answer the question **is there a difference?** If so, you use the CI to assess **how much of a difference exists**. I believe that scientists place too much emphasis on hypothesis testing.

2.5 Statistical versus practical significance

Suppose in the Tocopilla meteorite example, you rejected $H_0 : \mu = 0.54$ at the 5% level and found a 95% two-sided CI for μ to be 0.55 to 0.58. Although you have sufficient evidence to conclude that the population mean cooling rate μ differs from that suggested by evolutionary theory, the range of plausible values for μ is small and contains only values close to 0.54. Although you have shown statistical significance here, you need to ask yourself whether the actual difference between μ and 0.54 is large enough to be important. The answer to

such questions is always problem specific.

2.6 Design issues and power

An experiment may not be sensitive enough to pick up true differences. For example, in the Tocopilla meteorite example, suppose the true mean cooling rate is $\mu = 1.00$. To have a 50% chance of correctly rejecting $H_0 : \mu = 0.54$, you would need about $n = 48$ observations. If the true mean is $\mu = 0.75$, you would need about 221 observations to have a 50% chance of correctly rejecting H_0 . In general, the smaller the difference between the true and hypothesized mean (relative to the spread in the population), the more data that is needed to reject H_0 . If you have prior information on the expected difference between the true and hypothesized mean, you can design an experiment appropriately by choosing the sample size required to likely reject H_0 .

The **power** of a test is the probability of correctly rejecting H_0 when it is false. Equivalently,

$$\text{power} = 1 - \text{Prob}(\text{ failing to reject } H_0 \text{ when it is false }) = 1 - \text{Prob}(\text{ Type-II error}).$$

For a given sample size, the tests I have discussed have maximum power (or smallest probability of a Type-II error) among all tests with fixed size α . However, the actual power may be small, so sample size calculations, as briefly highlighted above, are important prior to collecting data. See your local statistician.

```
#### Power calculations (that you can do on your own)
# install.packages("pwr")
library(pwr)
?power.t.test
power.t.test(n = NULL, delta = 1.00 - 0.54, sd = sd(toco),
             , sig.level = 0.05, power = 0.50
             , type = "one.sample", alternative = "two.sided")
power.t.test(n = NULL, delta = 0.75 - 0.54, sd = sd(toco),
             , sig.level = 0.05, power = 0.50
             , type = "one.sample", alternative = "two.sided")
```

For more examples, try:

```
# install.packages("TeachingDemos")
library(TeachingDemos)
# Demonstrate concepts of Power.
?power.examp
```

2.7 One-sided tests on μ

There are many studies where a one-sided test is appropriate. The two common scenarios are the **lower one-sided test** $H_0 : \mu = \mu_0$ (or $\mu \geq \mu_0$) versus $H_A : \mu < \mu_0$ and the **upper one-sided test** $H_0 : \mu = \mu_0$ (or $\mu \leq \mu_0$) versus $H_A : \mu > \mu_0$. Regardless of the alternative hypothesis, the tests are based on the t -statistic:

$$t_s = \frac{\bar{Y} - \mu_0}{SE_{\bar{Y}}}.$$

For the **upper one-sided test**

1. Compute the critical value t_{crit} such that the area under the t -curve to the **right** of t_{crit} is the desired size α , that is $t_{\text{crit}} = t_\alpha$.
2. Reject H_0 if and only if $t_s \geq t_{\text{crit}}$.
3. The p-value for the test is the area under the t -curve to the **right** of the test statistic t_s .

The **upper one-sided test** uses the **upper tail** of the t -distribution for a rejection region. The p-value calculation reflects the form of the rejection region. You will reject H_0 only for large positive values of t_s which require \bar{Y} to be significantly greater than μ_0 . Does this make sense?

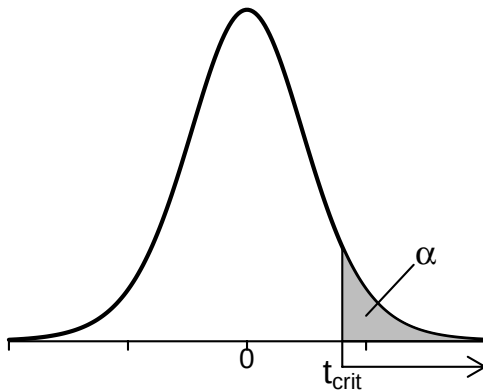
For the **lower one-sided test**

1. Compute the critical value t_{crit} such that the area under the t -curve to the **right** of t_{crit} is the desired size α , that is $t_{\text{crit}} = t_\alpha$.
2. Reject H_0 if and only if $t_s \leq -t_{\text{crit}}$.
3. The p-value for the test is the area under the t -curve to the **left** of the test statistic t_s .

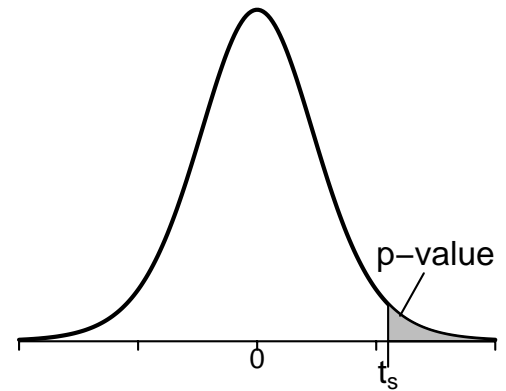
The **lower one-sided test** uses the **lower tail** of the t -distribution for a rejection region. The calculation of the rejection region in terms of $-t_{\text{crit}}$ is awkward but is necessary for hand calculations because many statistical tables only give upper tail percentiles. Note that here you will reject H_0 only for large negative values of t_s which require \bar{Y} to be significantly less than μ_0 .

As with two-sided tests, the p-value can be used to decide between rejecting or not rejecting H_0 for a test with a given size α . A picture of the rejection region and the p-value evaluation for one-sided tests is given below.

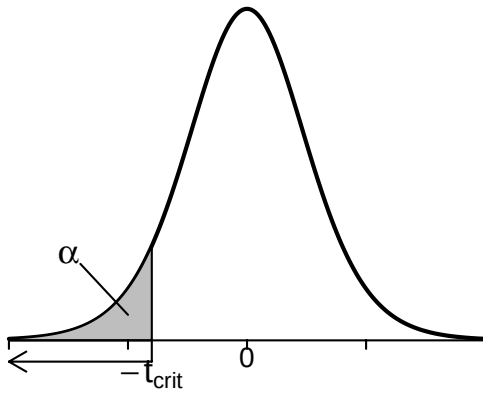
Upper One-Sided Rejection Region



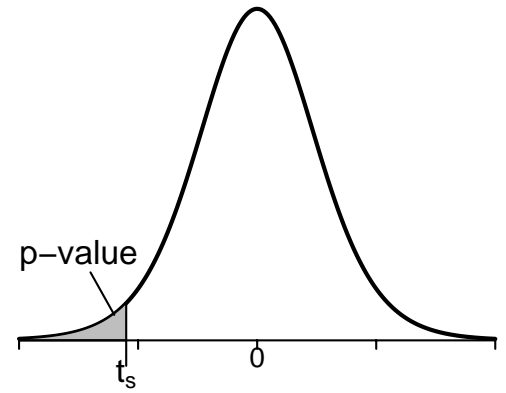
Upper One-Sided p-value



Lower One-Sided Rejection Region



Lower One-Sided p-value



Example: Weights of canned tomatoes A consumer group suspects that the average weight of canned tomatoes being produced by a large cannery is less than the advertised weight of 20 ounces. To check their conjecture, the group purchases 14 cans of the canner's tomatoes from various grocery stores. The weights of the contents of the cans to the nearest half ounce were as follows: 20.5, 18.5, 20.0, 19.5, 19.5, 21.0, 17.5, 22.5, 20.0, 19.5, 18.5, 20.0, 18.0, 20.5. Do the data confirm the group's suspicions? Test at the 5% level.

Let μ = the population mean weight for advertised 20 ounce cans of tomatoes produced by the cannery. The company claims that $\mu = 20$, but the consumer group believes that $\mu < 20$. Hence the consumer group wishes to test $H_0 : \mu = 20$ (or $\mu \geq 20$) against $H_A : \mu < 20$. The consumer group will reject H_0 only if the data overwhelmingly suggest that H_0 is false.

You should assess the normality assumption prior to performing the t -test. The stem-and-leaf display and the boxplot suggest that the distribution might be slightly skewed to the left. However, the skewness is not severe and no outliers are present, so the normality assumption is not unreasonable.

R output for the problem is given below. Let us do a hand calculation using the summarized data. The sample size, mean, and standard deviation are 14, 19.679, and 1.295, respectively. The standard error is $SE_{\bar{Y}} = s/\sqrt{n} = 0.346$. We see that the sample mean is less than 20. But is it sufficiently less than 20 for us to be willing to publicly refute the canner's claim? Let us carry out the test, first using the rejection region approach, and then by evaluating a p-value.

The test statistic is

$$t_s = \frac{\bar{Y} - \mu_0}{SE_{\bar{Y}}} = \frac{19.679 - 20}{0.346} = -0.93.$$

The critical value for a 5% one-sided test is $t_{0.05} = 1.771$, so we reject H_0 if $t_s < -1.771$ (you can get that value from r or from the table). The test statistic is not in the rejection region. Using the t -table, the p-value is between 0.15 and 0.20. I will draw a picture to illustrate the critical region and p-value calculation. The exact p-value from R is 0.185, which exceeds 0.05.

Both approaches lead to the conclusion that we do not have sufficient evidence to reject H_0 . That is, we do not have sufficient evidence to question the accuracy of the canner's claim. If you did reject H_0 , is there something about

how the data were recorded that might make you uncomfortable about your conclusions?

```
#### Example: Weights of canned tomatoes
tomato <- c(20.5, 18.5, 20.0, 19.5, 19.5, 21.0, 17.5
           , 22.5, 20.0, 19.5, 18.5, 20.0, 18.0, 20.5)

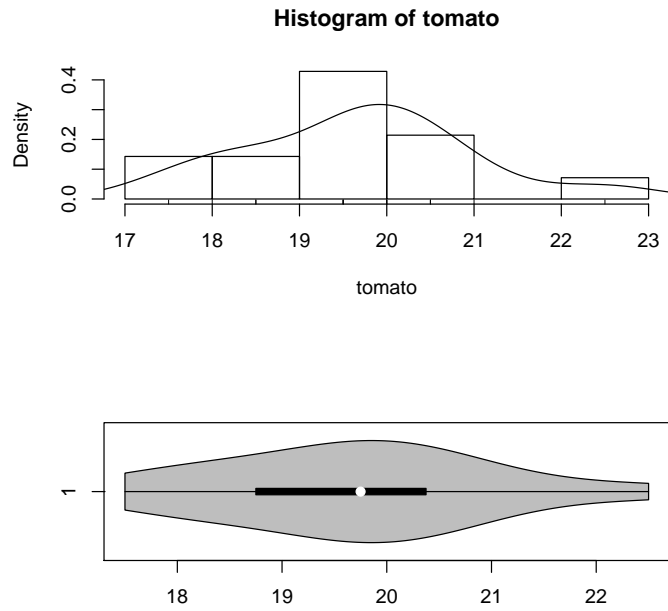
par(mfrow=c(2,1))
# Histogram overlaid with kernel density curve
hist(tomato, freq = FALSE, breaks = 6)
points(density(tomato), type = "l")
rug(tomato)

# violin plot
library(vioplot)
vioplot(tomato, horizontal=TRUE, col="gray")

# t.crit
qt(1 - 0.05/2, df = length(tomato) - 1)
## [1] 2.16

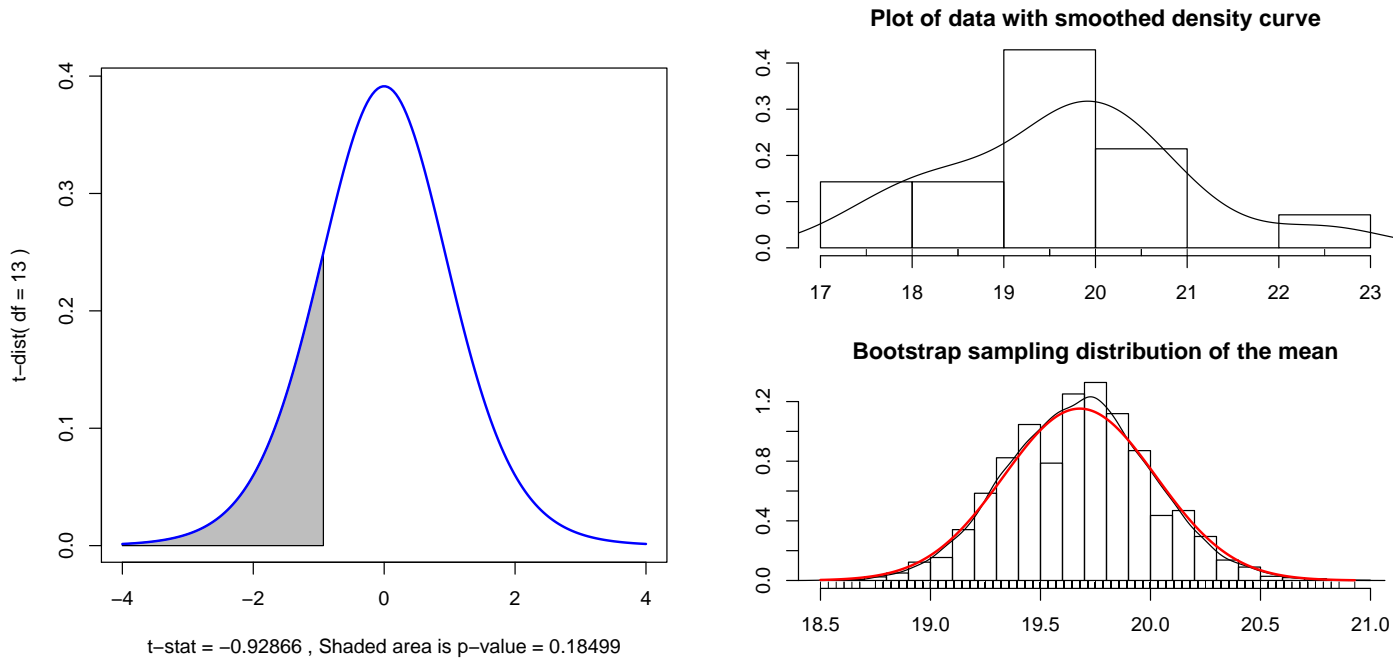
t.summary <- t.test(tomato, mu = 20, alternative = "less")
t.summary
##
##  One Sample t-test
##
## data: tomato
## t = -0.9287, df = 13, p-value = 0.185
## alternative hypothesis: true mean is less than 20
## 95 percent confidence interval:
##    -Inf 20.29
## sample estimates:
## mean of x
##      19.68

summary(tomato)
##   Min. 1st Qu. Median   Mean 3rd Qu.   Max.
## 17.5   18.8   19.8   19.7   20.4   22.5
```



The assumption of normality of the sampling distribution appears reasonable. Therefore, the results for the t -test above can be trusted.

```
t.dist.pval(t.summary)
bs.one.samp.dist(tomato)
```



2.7.1 One-sided CIs

How should you couple a one-sided test with a CI procedure? For a **lower one-sided test**, you are interested only in an **upper bound** on μ . Similarly, with an **upper one-sided test** you are interested in a **lower bound** on

μ . Computing these type of bounds maintains the consistency between tests and CI procedures. The general formulas for lower and upper $100(1 - \alpha)\%$ confidence bounds on μ are given by

$$\bar{Y} - t_{\text{crit}}SE_{\bar{Y}} \quad \text{and} \quad \bar{Y} + t_{\text{crit}}SE_{\bar{Y}}$$

respectively, where $t_{\text{crit}} = t_\alpha$.

In the cannery problem, to get an upper 95% bound on μ , the critical value is the same as we used for the one-sided 5% test: $t_{0.05} = 1.771$. The upper bound on μ is

$$\bar{Y} + t_{0.05}SE_{\bar{Y}} = 19.679 + 1.771 \times 0.346 = 19.679 + 0.613 = 20.292.$$

Thus, you are 95% confident that the population mean weight of the canner's 20oz cans of tomatoes is less than or equal to 20.29. As expected, this interval covers 20.

If you are doing a one-sided test in R, it will generate the correct one-sided bound. That is, a lower one-sided test will generate an upper bound, whereas an upper one-sided test generates a lower bound. If you only wish to compute a one-sided bound without doing a test, you need to specify the direction of the alternative which gives the type of bound you need. An upper bound was generated by R as part of the test we performed earlier. The result agrees with the hand calculation.

Quite a few packages, do not directly compute one-sided bounds so you have to fudge a bit. In the cannery problem, to get an upper 95% bound on μ , you take the upper limit from a 90% two-sided confidence limit on μ . The rational for this is that with the 90% two-sided CI, μ will fall above the upper limit 5% of the time and fall below the lower limit 5% of the time. Thus, you are 95% confident that μ falls below the upper limit of this interval, which gives us our one-sided bound. Here, you are 95% confident that the population mean weight of the canner's 20 oz cans of tomatoes is less than or equal to 20.29, which agrees with our hand calculation.

One-Sample T: Cans

Variable	N	Mean	StDev	SE Mean	90% CI
Cans	14	19.6786	1.2951	0.3461	(19.0656, 20.2915)

The same logic applies if you want to generalize the one-sided confidence bounds to arbitrary confidence levels and to lower one-sided bounds — always

double the error rate of the desired one-sided bound to get the error rate of the required two-sided interval! For example, if you want a lower 99% bound on μ (with a 1% error rate), use the lower limit on the 98% two-sided CI for μ (which has a 2% error rate).

■ CLICKER Qs — P-value ■

Chapter 3

Two-Sample Inferences

Learning objectives

After completing this topic, you should be able to:

select graphical displays that meaningfully compare independent populations.

assess the assumptions of the two-sample t-test visually.

decide whether the means between two populations are different.

recommend action based on a hypothesis test.

Achieving these goals contributes to mastery in these course learning outcomes:

1. organize knowledge.
5. define parameters of interest and hypotheses in words and notation.
6. summarize data visually, numerically, and descriptively.
8. use statistical software.
12. make evidence-based decisions.

3.1 Comparing Two Sets of Measurements

Suppose you have collected data on one variable from two (independent) samples and you are interested in “comparing” the samples. What tools are good to use?

Example: Head Breadths In this analysis, we will compare a physical feature of modern day Englishmen with the corresponding feature of some of

their ancient countrymen. The Celts were a vigorous race of people who once populated parts of England. It is not entirely clear whether they simply died out or merged with other people who were the ancestors of those who live in England today. A goal of this study might be to shed some light on possible genetic links between the two groups.

The study is based on the comparison of maximum head breadths (in millimeters) made on unearthed Celtic skulls and on a number of skulls of modern-day Englishmen. The data are given below. We have a sample of 18 Englishmen and an independent sample of 16 Celtic skulls.

```
#### Example: Head Breadths
# unstacked data as two vectors
english <- c(141, 148, 132, 138, 154, 142, 150, 146, 155, 158,
           150, 140, 147, 148, 144, 150, 149, 145)
celts   <- c(133, 138, 130, 138, 134, 127, 128, 138, 136, 131,
           126, 120, 124, 132, 125)
english
## [1] 141 148 132 138 154 142 150 146 155 158 150 140 147 148 144 150
## [17] 149 145
celts
## [1] 133 138 130 138 134 127 128 138 136 131 126 120 124 132 132 125
```

What features of these data would we likely be interested in comparing? The centers of the distributions, the spreads within each distribution, the distributional shapes, etc.

These data can be analyzed in R as either **unstacked** separate vectors or as **stacked** data where one column contains both samples, with a second column of labels or **subscripts** to distinguish the samples. It is easy to create stacked data from unstacked data and vice-versa. Many comparisons require the plots for the two groups to have the same scale, which is easiest to control when the data are stacked.

```
# stacked data as a vector of values and a vector of labels
HeadBreadth <- c(english, celts)
Group <- c(rep("English", length(english)), rep("Celts", length(celts)))
hb <- data.frame(HeadBreadth, Group)
hb
##   HeadBreadth   Group
## 1          141 English
## 2          148 English
## 3          132 English
## 4          138 English
## 5          154 English
```

```

## 6      142 English
## 7      150 English
## 8      146 English
## 9      155 English
## 10     158 English
## 11     150 English
## 12     140 English
## 13     147 English
## 14     148 English
## 15     144 English
## 16     150 English
## 17     149 English
## 18     145 English
## 19     133 Celts
## 20     138 Celts
## 21     130 Celts
## 22     138 Celts
## 23     134 Celts
## 24     127 Celts
## 25     128 Celts
## 26     138 Celts
## 27     136 Celts
## 28     131 Celts
## 29     126 Celts
## 30     120 Celts
## 31     124 Celts
## 32     132 Celts
## 33     132 Celts
## 34     125 Celts

```

3.1.1 Plotting head breadth data:

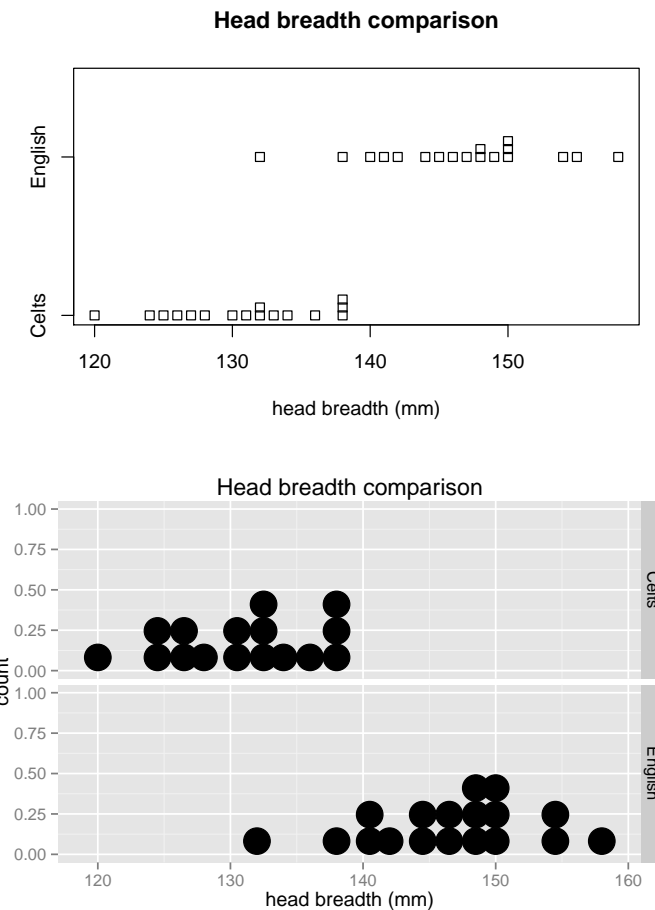
1. A dotplot with the same scale for both samples is obtained easily from the stacked data.

```

##### Plotting head breadth data
# stripchart (dotplot) using R base graphics
stripchart(HeadBreadth ~ Group, method = "stack", data = hb,
           main = "Head breadth comparison", xlab = "head breadth (mm)")

# stripchart (dotplot) using ggplot
library(ggplot2)
p <- ggplot(hb, aes(x = HeadBreadth))
p <- p + geom_dotplot(binwidth = 2)
p <- p + facet_grid(Group ~ .)      # rows are Group categories
p <- p + labs(title = "Head breadth comparison") + xlab("head breadth (mm)")
print(p)

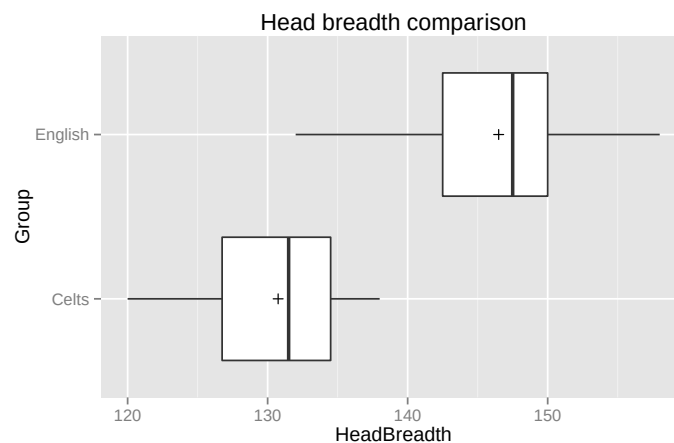
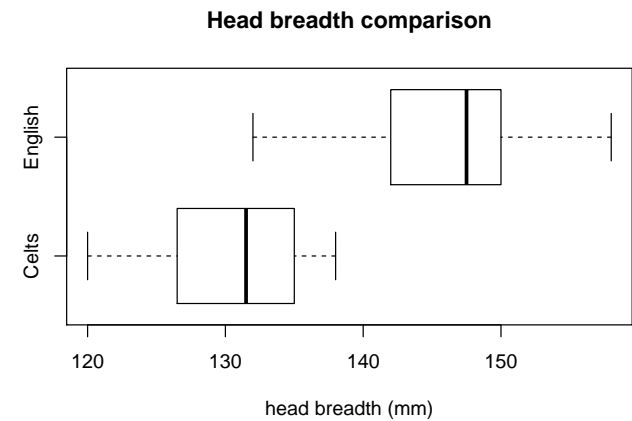
```



- Boxplots for comparison are most helpful when plotted in the same axes.

```
# boxplot using R base graphics
boxplot(HeadBreadth ~ Group, method = "stack", data = hb,
        horizontal = TRUE,
        main = "Head breadth comparison", xlab = "head breadth (mm)")

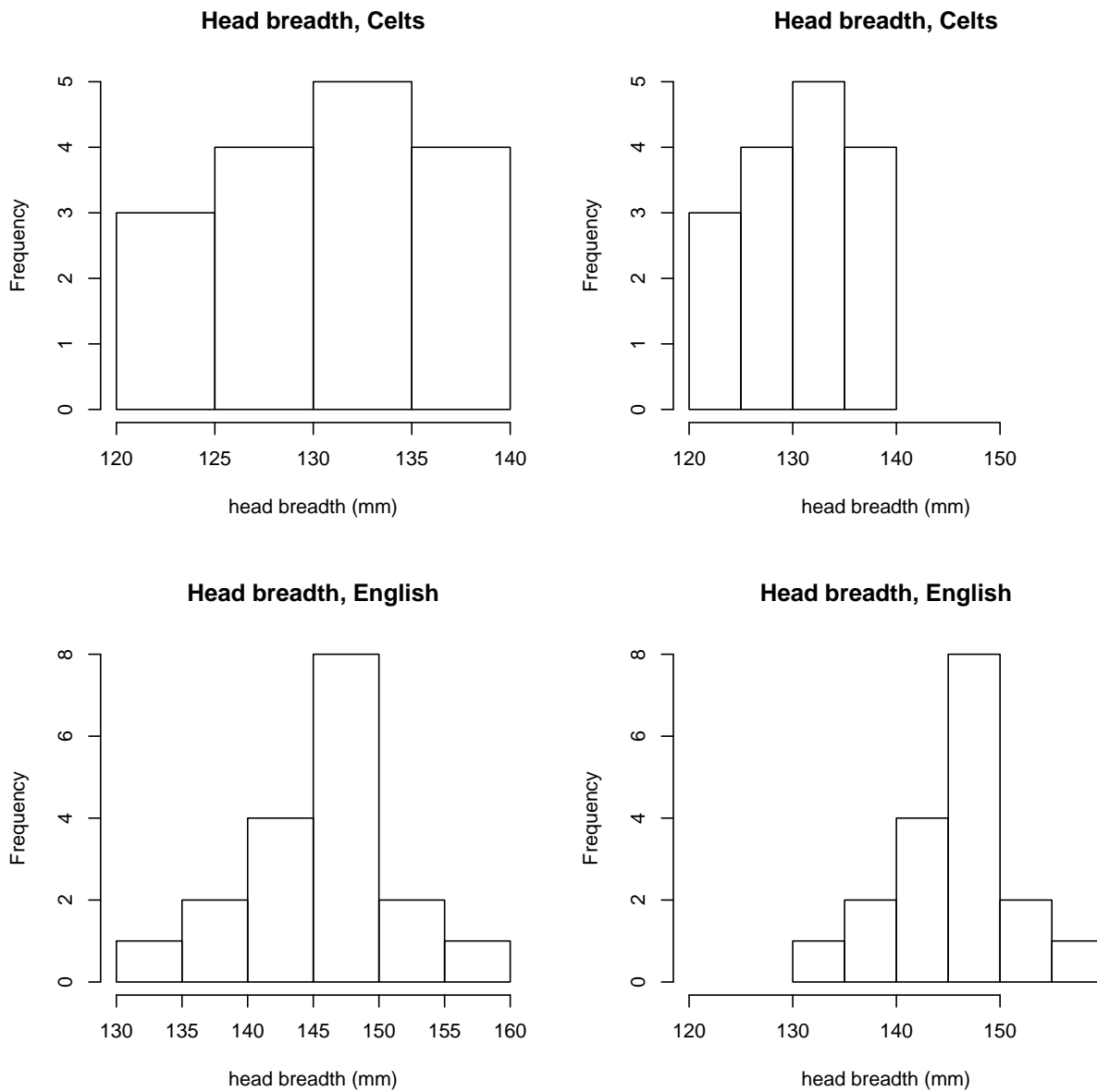
p <- ggplot(hb, aes(x = Group, y = HeadBreadth))
p <- p + geom_boxplot()
# add a "+" at the mean
p <- p + stat_summary(fun.y = mean, geom = "point", shape = 3, size = 2)
p <- p + coord_flip()
p <- p + labs(title = "Head breadth comparison")
print(p)
```



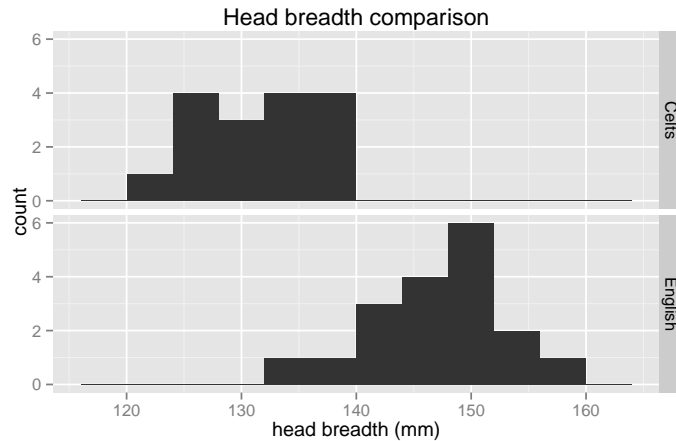
3. Histograms are hard to compare unless you make the scale and actual bins the same for both. Why is the pair on the right clearly preferable?

```
# histogram using R base graphics
par(mfcol=c(2,2))
hist(hb$HeadBreadth[(hb$Group == "Celts")],
     main = "Head breadth, Celts", xlab = "head breadth (mm)")
hist(hb$HeadBreadth[(hb$Group == "English")],
     main = "Head breadth, English", xlab = "head breadth (mm)")

# common x-axis limits based on the range of the entire data set
hist(hb$HeadBreadth[(hb$Group == "Celts")], xlim = range(hb$HeadBreadth),
      main = "Head breadth, Celts", xlab = "head breadth (mm)")
hist(hb$HeadBreadth[(hb$Group == "English")], xlim = range(hb$HeadBreadth),
      main = "Head breadth, English", xlab = "head breadth (mm)")
```

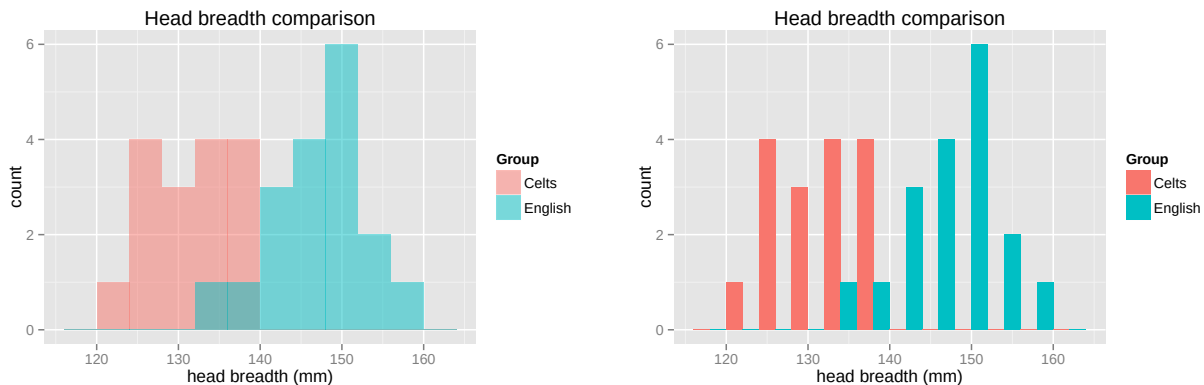


```
# histogram using ggplot
p <- ggplot(hb, aes(x = HeadBreadth))
p <- p + geom_histogram(binwidth = 4)
p <- p + facet_grid(Group ~ .)
p <- p + labs(title = "Head breadth comparison") + xlab("head breadth (mm)")
print(p)
```



```
p <- ggplot(hb, aes(x = HeadBreadth, fill=Group))
p <- p + geom_histogram(binwidth = 4, alpha = 0.5, position="identity")
p <- p + labs(title = "Head breadth comparison") + xlab("head breadth (mm)")
print(p)

p <- ggplot(hb, aes(x = HeadBreadth, fill=Group))
p <- p + geom_histogram(binwidth = 4, alpha = 1, position="dodge")
p <- p + labs(title = "Head breadth comparison") + xlab("head breadth (mm)")
print(p)
```



4. Stem-and-leaf displays for comparisons in R can be pretty useless. The stems are not forced to match (just like with histograms). It is pretty hard to make quick comparisons with the following:

```
stem(english, scale = 2)

##
## The decimal point is 1 digit(s) to the right of the |
##
## 13 | 2
## 13 | 8
## 14 | 0124
## 14 | 567889
## 15 | 0004
## 15 | 58

stem(celts, scale = 2)

##
## The decimal point is at the |
```

```
##  
## 120 | 0  
## 122 |  
## 124 | 00  
## 126 | 00  
## 128 | 0  
## 130 | 00  
## 132 | 000  
## 134 | 0  
## 136 | 0  
## 138 | 000
```

Using the `by()` function, you can get summaries by group.

```
##### summaries by group  
# summary for separate vectors  
summary(english)  
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.  
##      132    142    148    146    150    158  
summary(celts)  
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.  
##      120    127    132    131    134    138  
  
# comparing spreads, an assumption of equal variances seems reasonable  
sd(english)  
## [1] 6.382  
sd(celts)  
## [1] 5.434  
IQR(english)  
## [1] 7.5  
IQR(celts)  
## [1] 7.75  
  
# numerical summary of each column in data.frame hb by Group  
by(hb, Group, summary)  
## Group: Celts  
##   HeadBreadth      Group  
##   Min. :120  Celts :16  
##   1st Qu.:127  English: 0  
##   Median :132  
##   Mean   :131  
##   3rd Qu.:134  
##   Max.   :138  
## -----  
## Group: English  
##   HeadBreadth      Group  
##   Min. :132  Celts : 0  
##   1st Qu.:142  English:18  
##   Median :148  
##   Mean   :146
```

```
## 3rd Qu.:150
## Max. :158
```

3.1.2 Salient Features to Notice

The dotplots, boxplots, and histograms indicate that the English and Celt samples are slightly skewed to the left. There are no outliers in either sample. It is not unreasonable to operationally assume that the population frequency curves (i.e., the histograms for the populations from which the samples were selected) for the English and Celtic head breadths are normal. Therefore, the sampling distribution of the means will be reasonably normal.

The sample means and medians are close to each other in each sample, which is not surprising given the near symmetry and the lack of outliers.

The data suggest that the typical modern English head breadth is greater than that for Celts. The data sets have comparable spreads, as measured by either the standard deviation or the IQR.

3.2 Two-Sample Methods: Paired Versus Independent Samples

Suppose you have two populations of interest, say populations 1 and 2, and you are interested in comparing their (unknown) population means, μ_1 and μ_2 . Inferences on the unknown population means are based on samples from each population. In practice, most problems fall into one of two categories.

Independent samples where the sample taken from population 1 has no effect on which observations are selected from population 2, and vice versa.

Paired or dependent samples where experimental units are paired based on factors related or unrelated to the variable measured.

The distinction between paired and independent samples may be made clear through a series of examples.

Example The English and Celt head breadth samples are independent.

Example Suppose you are interested in whether the CaCO_3 (calcium carbonate) level in the Atrisco well field, which is the water source for Albuquerque, is changing over time. To answer this question, the CaCO_3 level was recorded at each of 15 wells at two time points. These data are paired. The two samples are the observations at Times 1 and 2.

Example To compare state incomes, a random sample of New Mexico households was selected, and an independent sample of Arizona households was obtained. It is reasonable to assume independent samples.

Example Suppose you are interested in whether the husband or wife is typically the heavier smoker among couples where both adults smoke. Data are collected on households. You measure the average number of cigarettes smoked by each husband and wife within the sample of households. These data are paired, i.e., you have selected husband wife pairs as the basis for the samples. It is reasonable to believe that the responses within a pair are related, or correlated.

Although the focus here will be on comparing population means, you should recognize that in paired samples you may also be interested, as in the problems above, in how observations compare within a pair. That is, a **paired comparison** might be interested in the **difference** between the two paired samples. These goals need not agree, depending on the questions of interest. Note that with paired data, the sample sizes are equal, and equal to the number of pairs.

■ CLICKER Qs — Independent or paired 1, STT.08.02.030 ■

■ CLICKER Qs — Independent or paired 2, STT.08.02.040 ■

3.3 Two Independent Samples: CI and Test Using Pooled Variance

These methods assume that the populations have normal frequency curves, with equal population standard deviations, i.e., $\sigma_1 = \sigma_2$. Let (n_1, \bar{Y}_1, s_1) and (n_2, \bar{Y}_2, s_2) be the sample sizes, means and standard deviations from the two samples. The standard CI for $\mu_1 - \mu_2$ is given by

$$\text{CI} = (\bar{Y}_1 - \bar{Y}_2) \pm t_{\text{crit}} SE_{\bar{Y}_1 - \bar{Y}_2}$$

The t -statistic for testing $H_0 : \mu_1 - \mu_2 = 0$ ($\mu_1 = \mu_2$) against $H_A : \mu_1 - \mu_2 \neq 0$ ($\mu_1 \neq \mu_2$) is given by

$$t_s = \frac{\bar{Y}_1 - \bar{Y}_2}{SE_{\bar{Y}_1 - \bar{Y}_2}}.$$

The standard error of $\bar{Y}_1 - \bar{Y}_2$ used in both the CI and the test is given by

$$SE_{\bar{Y}_1 - \bar{Y}_2} = s_{\text{pooled}} \sqrt{\frac{1}{n_1} + \frac{1}{n_2}}.$$

Here the **pooled variance estimator**,

$$s_{\text{pooled}}^2 = \frac{(n_1 - 1)s_1^2 + (n_2 - 1)s_2^2}{n_1 + n_2 - 2},$$

is our best estimate of the common population variance. The pooled estimator of variance is a weighted average of the two sample variances, with more weight given to the larger sample. If $n_1 = n_2$ then s_{pooled}^2 is the average of s_1^2 and s_2^2 . The critical value t_{crit} for CI and tests is obtained in usual way from a t -table with $df = n_1 + n_2 - 2$. For the test, follow the one-sample procedure, with the new t_s and t_{crit} .

The pooled CI and tests are sensitive to the normality and equal standard deviation assumptions. The observed data can be used to assess the reasonableness of these assumptions. You should look at boxplots and histograms to assess normality, and should check whether $s_1 \doteq s_2$ to assess the assumption $\sigma_1 = \sigma_2$. Formal tests of these assumptions will be discussed later.

3.4 Satterthwaite's Method, unequal variances

Satterthwaite's method assumes normality, but does not require equal population standard deviations. Satterthwaite's procedures are somewhat conservative, and adjust the SE and df to account for unequal population variances. Satterthwaite's method uses the same CI and test statistic formula, with a modified standard error:

$$SE_{\bar{Y}_1 - \bar{Y}_2} = \sqrt{\frac{s_1^2}{n_1} + \frac{s_2^2}{n_2}},$$

and degrees of freedom:

$$df = \frac{\left(\frac{s_1^2}{n_1} + \frac{s_2^2}{n_2}\right)^2}{\frac{s_1^4}{n_1^2(n_1-1)} + \frac{s_2^4}{n_2^2(n_2-1)}}.$$

Note that $df = n_1 + n_2 - 2$ when $n_1 = n_2$ and $s_1 = s_2$. The Satterthwaite and pooled variance procedures usually give similar results when $s_1 \doteq s_2$.

The df formula for Satterthwaite's method is fairly complex, so when done by hand some use a conservative df formula that uses the minimum of $n_1 - 1$ and $n_2 - 1$ instead.

3.4.1 R Implementation

R does the pooled and Satterthwaite (Welch) analyses, either on stacked or unstacked data. The output will contain a p-value for a two-sided test of equal population means and a CI for the difference in population means. If you include `var.equal = TRUE` you will get the pooled method, otherwise the output is for Satterthwaite's method.

Example: Head Breadths The English and Celts are independent samples. We looked at boxplots and histograms, which suggested that the normality assumption for the t -test is reasonable. The R output shows the English and

Celt sample standard deviations and IQRs are fairly close, so the pooled and Satterthwaite results should be comparable. The pooled analysis is preferable here, but either is appropriate.

We are interested in difference in mean head breadths between Celts and English.

1. Define the population parameters and hypotheses in words and notation

Let μ_1 and μ_2 be the mean head breadth for the Celts and English, respectively.

In words: “The difference in population means between Celts and English is different from zero mm.”

In notation: $H_0 : \mu_1 = \mu_2$ versus $H_A : \mu_1 \neq \mu_2$.

Alternatively: $H_0 : \mu_1 - \mu_2 = 0$ versus $H_A : \mu_1 - \mu_2 \neq 0$.

2. Calculate summary statistics from sample

Mean, standard deviation, sample size:

```
#### Calculate summary statistics
m1 <- mean(celts)
s1 <- sd(celts)
n1 <- length(celts)
m2 <- mean(english)
s2 <- sd(english)
n2 <- length(english)
c(m1, s1, n1)

## [1] 130.750 5.434 16.000

c(m2, s2, n2)

## [1] 146.500 6.382 18.000
```

The pooled-standard deviation, standard error, and degrees-of-freedom are:

```
sdpool <- sqrt(((n1 - 1) * s1^2 + (n2 - 1) * s2^2) / (n1 + n2 - 2))
sdpool

## [1] 5.957

SEpool <- sdpool * sqrt(1 / n1 + 1 / n2)
SEpool

## [1] 2.047

dfpool <- n1 + n2 - 2
dfpool

## [1] 32

t_pool <- (m1 - m2) / SEpool
t_pool

## [1] -7.695
```

The Satterthwaite SE and degrees-of-freedom are:

```
SE_Sat <- sqrt(s1^2 / n1 + s2^2 / n2)
SE_Sat

## [1] 2.027

df_Sat <- (SE_Sat^2)^2 / (s1^4 / (n1^2 * (n1 - 1)) + s2^4 / (n2^2 * (n2 - 1)))
df_Sat

## [1] 31.95

t_Sat <- (m1 - m2) / SE_Sat
t_Sat

## [1] -7.77
```

3. Specify confidence level, calculate t-stat, CI limits, p-value

Let us calculate a 95% CI for $\mu_1 - \mu_2$.

Assuming equal variances, using pooled-variance procedure:

```
## Equal variances
# var.equal = FALSE is the default

# two-sample t-test specifying two separate vectors
t.summary.eqvar <- t.test(celts, english, var.equal = TRUE)
t.summary.eqvar

##
## Two Sample t-test
##
## data: celts and english
## t = -7.695, df = 32, p-value = 9.003e-09
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -19.92 -11.58
## sample estimates:
## mean of x mean of y
## 130.8 146.5
```

Not assuming equal variances, Satterthwaite (Welch):

```
# two-sample t-test with unequal variances (Welch = Satterthwaite)
# specified using data.frame and a formula, HeadBreadth by Group
t.summary.uneqvar <- t.test(HeadBreadth ~ Group, data = hb, var.equal = FALSE)
t.summary.uneqvar

##
## Welch Two Sample t-test
##
## data: HeadBreadth by Group
## t = -7.77, df = 31.95, p-value = 7.414e-09
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -19.88 -11.62
## sample estimates:
## mean in group Celts mean in group English
```

##	130.8	146.5
----	-------	-------

The form of the output will tell you which sample corresponds to population 1 and which corresponds to population 2.

4. Summarize in words (Using the pooled-variance results.)

The pooled analysis strongly suggests that $H_0 : \mu_1 - \mu_2 = 0$ is false, given the large t -statistic of -7.7 and two-sided p-value of 9×10^{-9} . Because the p-value < 0.05 we reject the Null hypothesis in favor of the Alternative hypothesis concluding that the difference in population mean head breadths between the Celts and English are different.

We are 95% confident that the difference in population means, $\mu_1 - \mu_2$, is between -19.9 and -11.6 mm. That is, we are 95% confident that the population mean head breadth for Englishmen (μ_2) exceeds the population mean head breadth for Celts (μ_1) by between 11.6 and 19.9 mm.

The CI interpretation is made easier by recognizing that we concluded the population means are different, so the direction of difference must be consistent with that seen in the observed data, where the sample mean head breadth for Englishmen exceeds that for the Celts. Thus, the limits on the CI for $\mu_1 - \mu_2$ tells us how much smaller the mean is for the Celts (that is, between -19.9 and -11.6 mm).

5. Check assumptions

The assumption of equal population variances will be left to a later chapter. We can test the assumption that the distribution of $\bar{Y}_1 - \bar{Y}_2$ is normal using the bootstrap in the following function.

```
#### Visual comparison of whether sampling distribution is close to Normal via Bootstrap
# a function to compare the bootstrap sampling distribution
#   of the difference of means from two samples with
#   a normal distribution with mean and SEM estimated from the data
bs.two.samp.diff.dist <- function(dat1, dat2, N = 1e4) {
  n1 <- length(dat1);
  n2 <- length(dat2);
  # resample from data
  sam1 <- matrix(sample(dat1, size = N * n1, replace = TRUE), ncol=N);
  sam2 <- matrix(sample(dat2, size = N * n2, replace = TRUE), ncol=N);
  # calculate the means and take difference between populations
  sam1.mean <- colMeans(sam1);
  sam2.mean <- colMeans(sam2);
  diff.mean <- sam1.mean - sam2.mean;
  # save par() settings
  old.par <- par(no.readonly = TRUE)
  # make smaller margins
  par(mfrow=c(3,1), mar=c(3,2,2,1), oma=c(1,1,1,1))
  # Histogram overlaid with kernel density curve
  hist(dat1, freq = FALSE, breaks = 6
       , main = paste("Sample 1", "\n"
                     , "n =", n1
                     , ", mean =", signif(mean(dat1), digits = 5)
                     , ", sd =", signif(sd(dat1), digits = 5)))
  hist(dat2, freq = FALSE, breaks = 6
       , main = paste("Sample 2", "\n"
                     , "n =", n2
                     , ", mean =", signif(mean(dat2), digits = 5)
                     , ", sd =", signif(sd(dat2), digits = 5)))
  # Plot the difference of means
  plot(density(sam1 - sam2), xlab = "Difference in Means")
  lines(density(sam1 - sam2))
}
```

```

, xlim = range(c(dat1, dat2)))
points(density(dat1), type = "l")
rug(dat1)

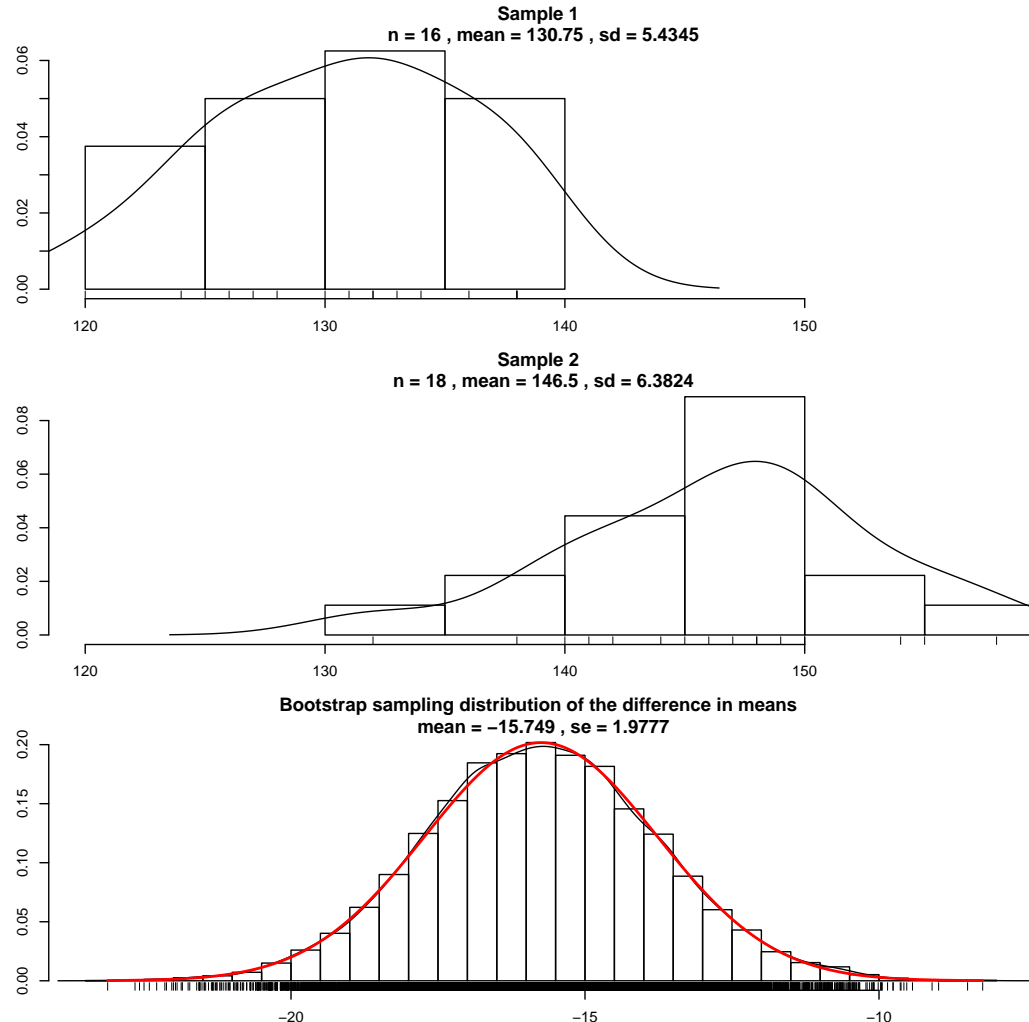
hist(dat2, freq = FALSE, breaks = 6
    , main = paste("Sample 2", "\n"
        , "n =", n2
        , ", mean =", signif(mean(dat2), digits = 5)
        , ", sd =", signif(sd(dat2), digits = 5))
    , xlim = range(c(dat1, dat2)))
points(density(dat2), type = "l")
rug(dat2)

hist(diff.mean, freq = FALSE, breaks = 25
    , main = paste("Bootstrap sampling distribution of the difference in means", "\n"
        , "mean =", signif(mean(diff.mean), digits = 5)
        , ", se =", signif(sd(diff.mean), digits = 5)))
# overlay a density curve for the sample means
points(density(diff.mean), type = "l")
# overlay a normal distribution, bold and red
x <- seq(min(diff.mean), max(diff.mean), length = 1000)
points(x, dnorm(x, mean = mean(diff.mean), sd = sd(diff.mean))
    , type = "l", lwd = 2, col = "red")
# place a rug of points under the plot
rug(diff.mean)
# restore par() settings
par(old.par)
}

```

The distribution of difference in means in the third plot looks very close to normal.

```
bs.two.samp.diff.dist(celts, english)
```



Example: Androstenedione levels in diabetics The data consist of independent samples of diabetic men and women. For each individual, the scientist recorded their androstenedione level (ng/dL) (a hormone, and Mark McGwire's favorite dietary supplement). Let μ_1 = mean androstenedione level for the population of diabetic men, and μ_2 = mean androstenedione level for the population of diabetic women. We are interested in comparing the population means given the observed data.

The raw data and R output are given below. The boxplots suggest that the distributions are reasonably symmetric. However, the normality assumption for the women is unreasonable due to the presence of outliers. The equal population standard deviation assumption also appears unreasonable. The sample standard deviation for men is noticeably larger than the women's standard deviation, even with outliers in the women's sample.

```
#### Example: Androstenedione levels in diabetics
# Data and numerical summaries
men <- c(217, 123, 80, 140, 115, 135, 59, 126, 70, 63,
       147, 122, 108, 70)
women <- c( 84, 87, 77, 84, 73, 66, 70, 35, 77, 73,
          56, 112, 56, 84, 80, 101, 66, 84)
level <- c(men, women)
sex <- c(rep("men", length(men)), rep("women", length(women)))
andro <- data.frame(level, sex)
andro

##      level    sex
## 1     217   men
## 2     123   men
## 3      80   men
## 4     140   men
## 5     115   men
## 6     135   men
## 7      59   men
## 8     126   men
## 9      70   men
## 10     63   men
## 11     147   men
## 12     122   men
## 13     108   men
## 14      70   men
## 15      84 women
```

```
## 16    87 women
## 17    77 women
## 18    84 women
## 19    73 women
## 20    66 women
## 21    70 women
## 22    35 women
## 23    77 women
## 24    73 women
## 25    56 women
## 26    112 women
## 27    56 women
## 28    84 women
## 29    80 women
## 30    101 women
## 31    66 women
## 32    84 women

# numerical summaries
by(andro, sex, summary)

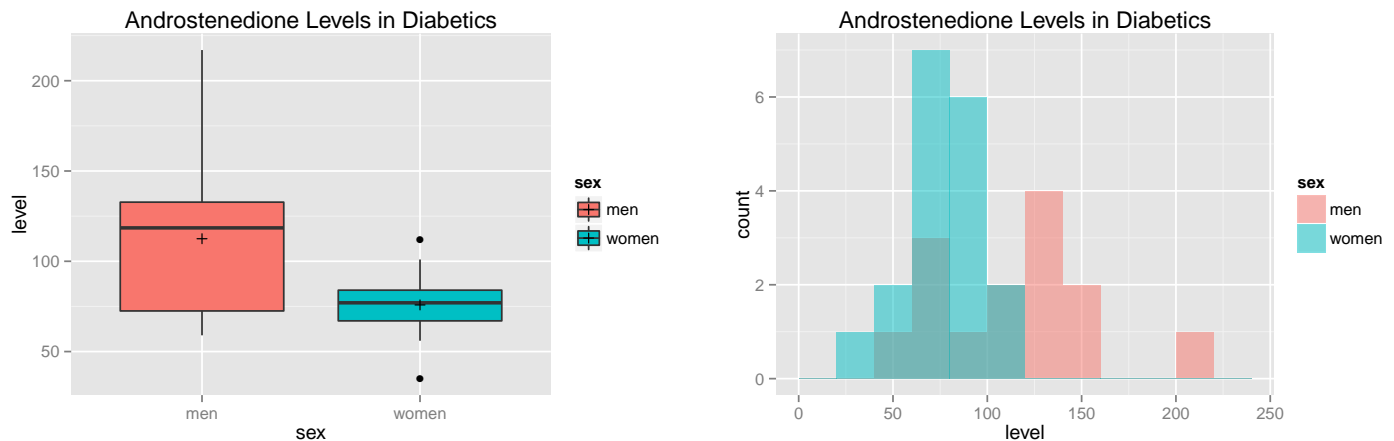
## sex: men
##      level      sex
## Min.   : 59.0   men   :14
## 1st Qu.: 72.5   women: 0
## Median :118.5
## Mean   :112.5
## 3rd Qu.:132.8
## Max.   :217.0
## -----
## sex: women
##      level      sex
## Min.   : 35.0   men   : 0
## 1st Qu.: 67.0   women:18
## Median : 77.0
## Mean   : 75.8
## 3rd Qu.: 84.0
## Max.   :112.0

c(sd(men), sd(women), IQR(men), IQR(women), length(men), length(women))
## [1] 42.75 17.24 60.25 17.00 14.00 18.00

p <- ggplot(andro, aes(x = sex, y = level, fill=sex))
p <- p + geom_boxplot()
# add a "+" at the mean
p <- p + stat_summary(fun.y = mean, geom = "point", shape = 3, size = 2)
#p <- p + coord_flip()
p <- p + labs(title = "Androstenedione Levels in Diabetics")
print(p)

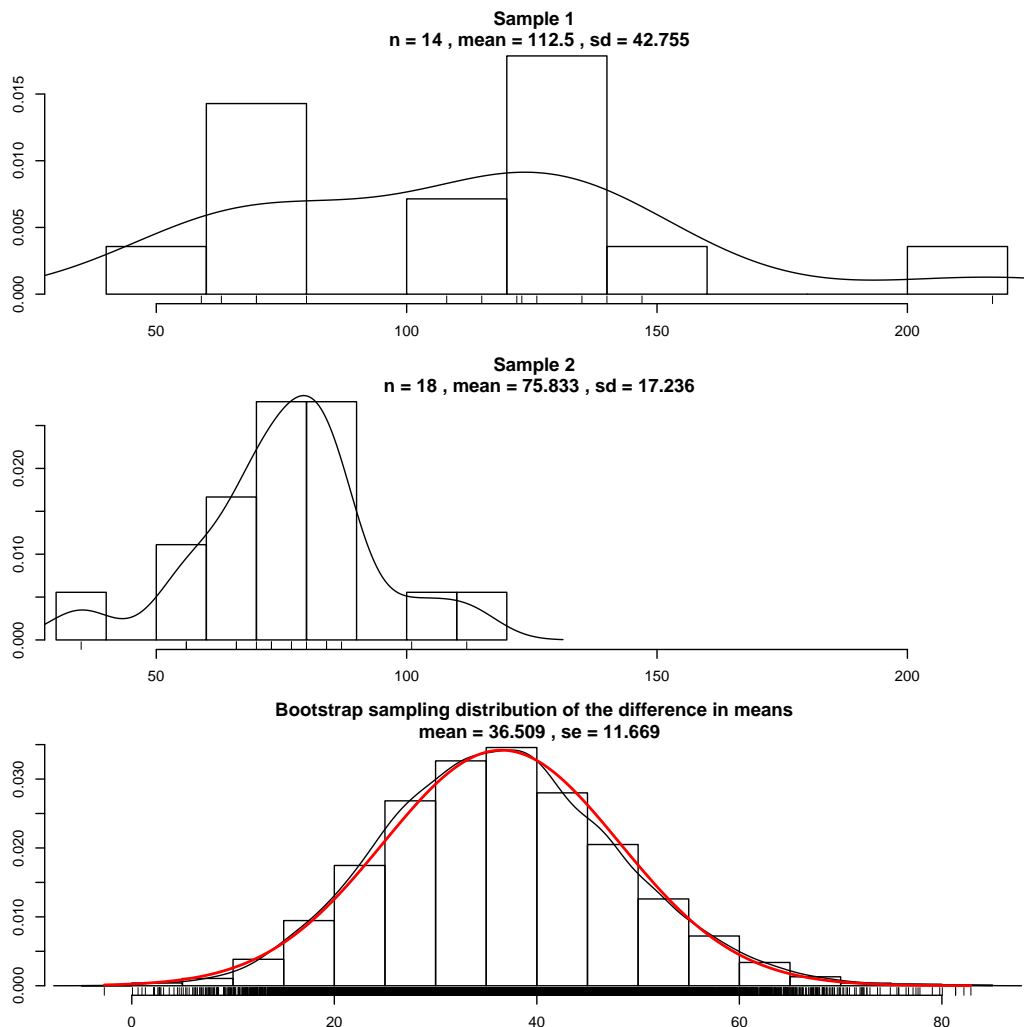
p <- ggplot(andro, aes(x = level, fill=sex))
p <- p + geom_histogram(binwidth = 20, alpha = 0.5, position="identity")
```

```
p <- p + labs(title = "Androstenedione Levels in Diabetics")
print(p)
```



Because of the large difference in variances, I will be more comfortable with the Satterthwaite analysis here than the pooled variance analysis. The normality assumption of the difference in means appears to be met using the bootstrap assessment. The distribution of difference in means in the third plot looks very close to normal.

```
bs.two.samp.diff.dist(men, women)
```



1. Define the population parameters and hypotheses in words and notation

Let μ_1 and μ_2 be the mean androstenedione level for diabetic men and women, respectively.

In words: “The difference in population mean androstenedione levels between diabetic men and women is different from zero.”

In notation: $H_0 : \mu_1 - \mu_2 = 0$ versus $H_A : \mu_1 - \mu_2 \neq 0$.

2. Calculate summary statistics from sample

(see above)

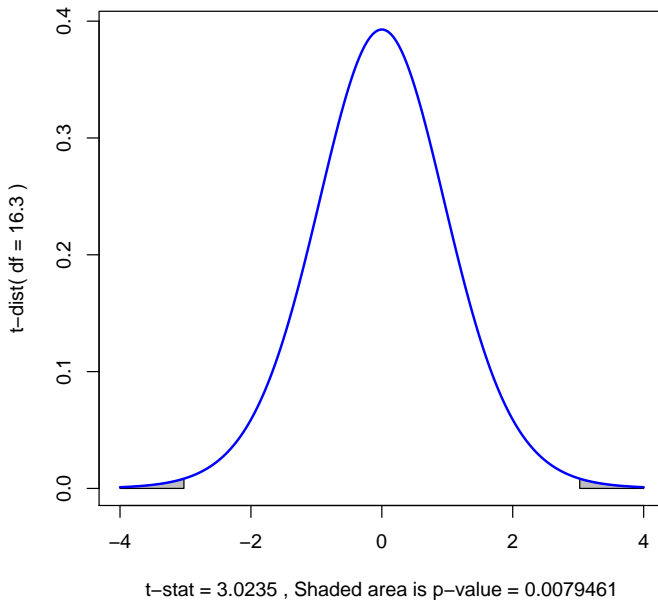
3. Specify confidence level, calculate t-stat, CI limits, p-value

Not assuming equal variances, Satterthwaite (Welch):

```
# two-sample t-test with unequal variances (Welch = Satterthwaite)
#   specified using data.frame and a formula, level by sex
t.summary <- t.test(level ~ sex, data = andro, var.equal = FALSE)
t.summary

##
## Welch Two Sample t-test
##
## data: level by sex
## t = 3.023, df = 16.3, p-value = 0.007946
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 11.00 62.34
## sample estimates:
## mean in group men mean in group women
## 112.50 75.83
```

```
# plot t-distribution with shaded p-value
t.dist.pval(t.summary)
```



4. Summarize in words The unequal-variance analysis suggests that $H_0 : \mu_1 - \mu_2 = 0$ is false, given the large t -statistic of 3.02 and two-sided p-value of 0.008. Because the p-value < 0.05 we reject the Null hypothesis in favor of the Alternative hypothesis concluding that the difference in population mean androstenedione levels between diabetic men and women are different.

We are 95% confident that the difference in population means, $\mu_1 - \mu_2$, is between 11 and 62.3 ng/dL.

5. Check assumptions

As checked before, while the assumption of equal population variances is not met, the assumption that the distribution of $\bar{Y}_1 - \bar{Y}_2$ is normal using the bootstrap appeared reasonable.

As a comparison, let us examine the output for the pooled procedure (which is inappropriate since variances are unequal). The p-value for the pooled t -test is 0.002, whereas the 95% confidence limits are 14.1 and 59.2. That is, we are 95% confident that the population mean andro level for men exceeds that for women by at least 14.1 but by no more than 59.2. These results are qualitatively similar to the Satterthwaite conclusions.

3.5 One-Sided Tests

One-sided tests for two-sample problems are where the null hypothesis is $H_0 : \mu_1 - \mu_2 = 0$ but the alternative is directional, either $H_A : \mu_1 - \mu_2 < 0$ (i.e., $\mu_1 < \mu_2$) or $H_A : \mu_1 - \mu_2 > 0$ (i.e., $\mu_1 > \mu_2$). Once you understand the general form of rejection regions and p-values for one-sample tests, the one-sided two-sample tests do not pose any new problems. Use the t -statistic, with the appropriate tail of the t -distribution to define critical values and p-values. One-sided two-sample tests are directly implemented in R, by specifying the type of test with `alternative = "less"` or `alternative = "greater"`. One-sided confidence bounds are given with the one-sided tests.

3.6 Paired Analysis

With paired data, inferences on $\mu_1 - \mu_2$ are based on the sample of differences within pairs. By taking differences within pairs, two dependent samples are transformed into one sample, which contains the relevant information for inferences on $\mu_d = \mu_1 - \mu_2$. To see this, suppose the observations within a pair are Y_1 and Y_2 . Then within each pair, compute the difference $d = Y_1 - Y_2$:

$$\begin{aligned} d_1 &= Y_{11} - Y_{21} \\ d_2 &= Y_{12} - Y_{22} \\ &\vdots \\ d_n &= Y_{1n} - Y_{2n} \end{aligned}$$

If the Y_1 data are from a population with mean μ_1 and the Y_2 data are from a population with mean μ_2 , then the ds are a sample from a population with mean $\mu_d = \mu_1 - \mu_2$. Furthermore, if the sample of differences comes from a normal population, then we can use standard one-sample techniques on d_1, \dots, d_n to test $\mu_d = 0$ (that is, $\mu_1 = \mu_2$), and to get a CI for $\mu_d = \mu_1 - \mu_2$.

Let $\bar{d} = n^{-1} \sum_i d_i = \bar{Y}_1 - \bar{Y}_2$ be the sample mean of the differences (which is also the mean difference), and let s_d be the sample standard deviation of the differences. The standard error of \bar{d} is $SE_{\bar{d}} = s_d / \sqrt{n}$, where n is the number

of pairs. The paired t -test (two-sided) CI for μ_d is given by $\bar{d} \pm t_{\text{crit}}SE_{\bar{d}}$. To test $H_0 : \mu_d = 0$ ($\mu_1 = \mu_2$) against $H_A : \mu_d \neq 0$ ($\mu_1 \neq \mu_2$), use

$$t_s = \frac{\bar{d} - 0}{SE_{\bar{d}}}$$

to compute a p-value as in a two-sided one-sample test. One-sided tests are evaluated in the usual way for one-sample tests on means.

A graphical analysis of paired data focuses on the **sample of differences**, and not on the original samples. In particular, the normality assumption is assessed on the sample of differences.

3.6.1 R Analysis

The most natural way to enter paired data is as two columns, one for each treatment group. You can then create a new column of differences, and do the usual one-sample graphical and inferential analysis on this column of differences, or you can do the paired analysis directly without this intermediate step.

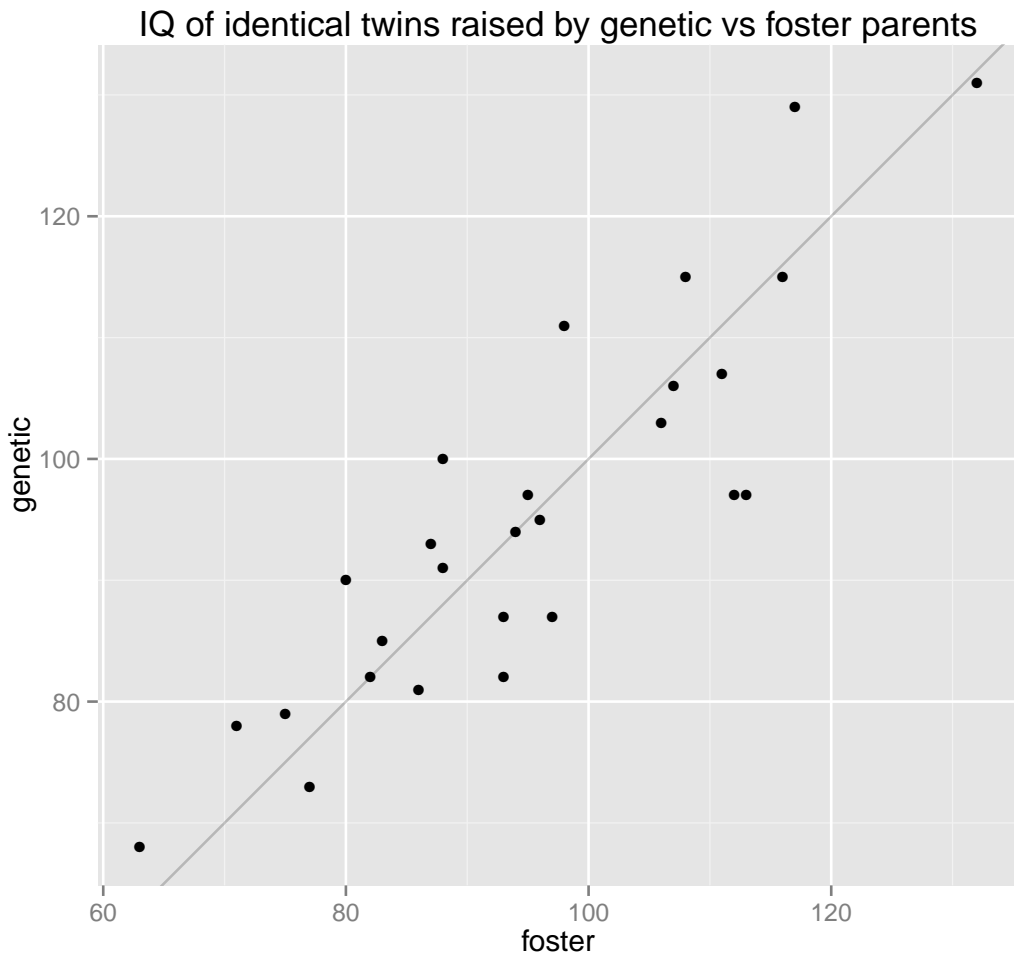
Example: Paired Analysis of Data on Twins Burt (1966) presented data on IQ scores for identical twins that were raised apart, one by foster parents and one by the genetic parents. Assuming the data are a random sample of twin pairs, consider comparing the population mean IQs for twins raised at home to those raised by foster parents. Let μ_f =population mean IQ for twin raised by foster parents, and μ_g =population mean IQ for twin raised by genetic parents.

I created the data in the worksheet (c1=foster; c2=genetic), and computed the differences between the IQ scores for the children raised by the genetic and foster parents (c3=diff=genetic-foster). I also made a scatter plot of the genetic versus foster IQ scores.

```
#### Example: Paired Analysis of Data on Twins
# Data and numerical summaries
foster <- c(82, 80, 88, 108, 116, 117, 132, 71, 75, 93,
           95, 88, 111, 63, 77, 86, 83, 93, 97, 87,
           94, 96, 112, 113, 106, 107, 98)
genetic <- c(82, 90, 91, 115, 115, 129, 131, 78, 79, 82,
            97, 100, 107, 68, 73, 81, 85, 87, 87, 93,
            94, 95, 97, 97, 103, 106, 111)
diff <- genetic - foster;
```

```
iq <- data.frame(foster, genetic, diff)

# scatterplot of foster and genetic IQs, with 1:1 line
p <- ggplot(iq, aes(x = foster, y = genetic))
# draw a 1:1 line, dots above line indicate "genetic > foster"
p <- p + geom_abline(intercept=0, slope=1, alpha=0.2)
p <- p + geom_point()
# make the axes square so it's a fair visual comparison
p <- p + coord_equal()
p <- p + labs(title = "IQ of identical twins raised by genetic vs foster parents")
print(p)
```



This plot of IQ scores shows that scores are related within pairs of twins. This is consistent with the need for a paired analysis.

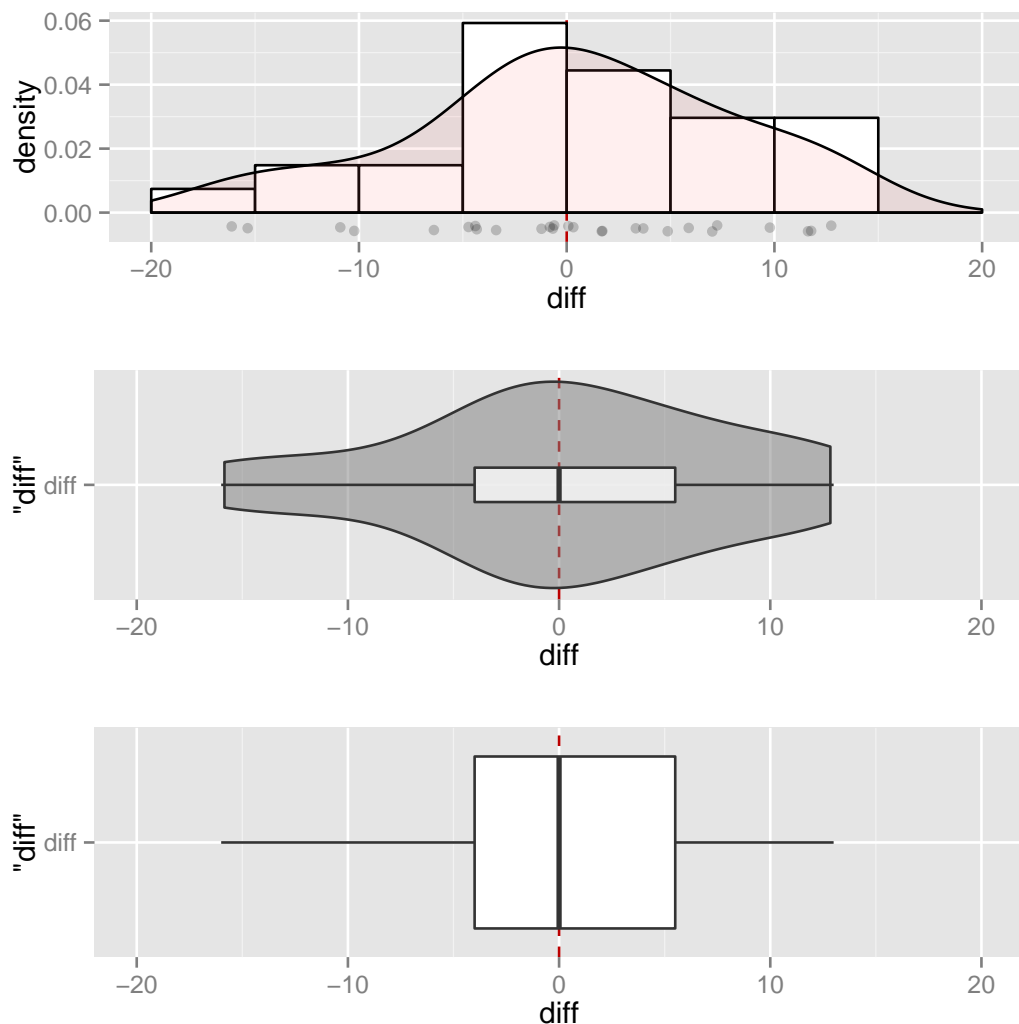
Given the sample of differences, I created a boxplot and a stem and leaf display, neither of which showed marked deviation from normality. The boxplot is centered at zero, so one would not be too surprised if the test result is insignificant.

```
p1 <- ggplot(iq, aes(x = diff))
p1 <- p1 + scale_x_continuous(limits=c(-20,+20))
# vertical line at 0
p1 <- p1 + geom_vline(xintercept=0, colour="#BB0000", linetype="dashed")
p1 <- p1 + geom_histogram(aes(y=..density..)
                           , binwidth=5
                           , colour="black", fill="white")
# Overlay with transparent density plot
p1 <- p1 + geom_density(alpha=0.1, fill="#FF6666")
p1 <- p1 + geom_point(aes(y = -0.005)
                      , position = position_jitter(height = 0.001)
                      , alpha = 1/5)

# violin plot
p2 <- ggplot(iq, aes(x = "diff", y = diff))
p2 <- p2 + scale_y_continuous(limits=c(-20,+20))
p2 <- p2 + geom_hline(yintercept=0, colour="#BB0000", linetype="dashed")
p2 <- p2 + geom_violin(fill = "gray50", alpha=1/2)
p2 <- p2 + geom_boxplot(width = 0.2, alpha = 3/4)
p2 <- p2 + coord_flip()

# boxplot
p3 <- ggplot(iq, aes(x = "diff", y = diff))
p3 <- p3 + scale_y_continuous(limits=c(-20,+20))
p3 <- p3 + geom_hline(yintercept=0, colour="#BB0000", linetype="dashed")
p3 <- p3 + geom_boxplot()
p3 <- p3 + coord_flip()

library(gridExtra)
grid.arrange(p1, p2, p3, ncol=1)
```



The normality assumption of the sample mean for a one-sample test is satisfied (below, left).

Given the sample of differences, I generated a one-sample CI and test. The hypothesis under test is $\mu_d = \mu_g - \mu_f = 0$. The p-value for this test is large. We do not have sufficient evidence to claim that the population mean IQs for twins raised apart are different. This is consistent with the CI for μ_d given below, which covers zero.

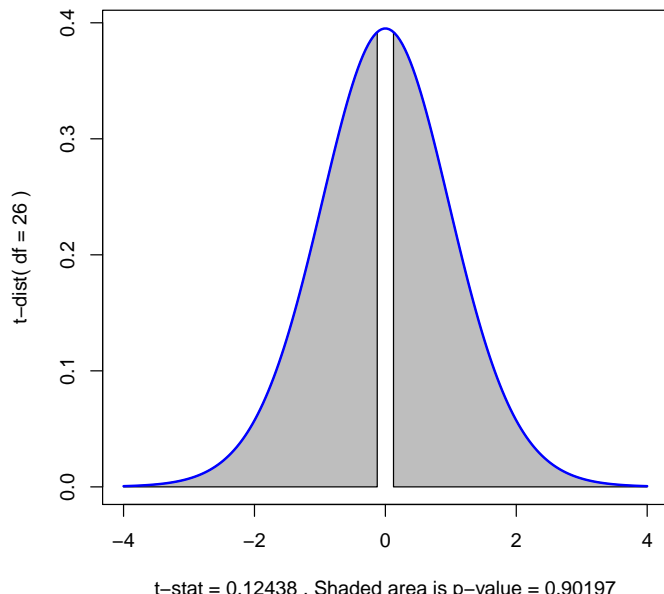
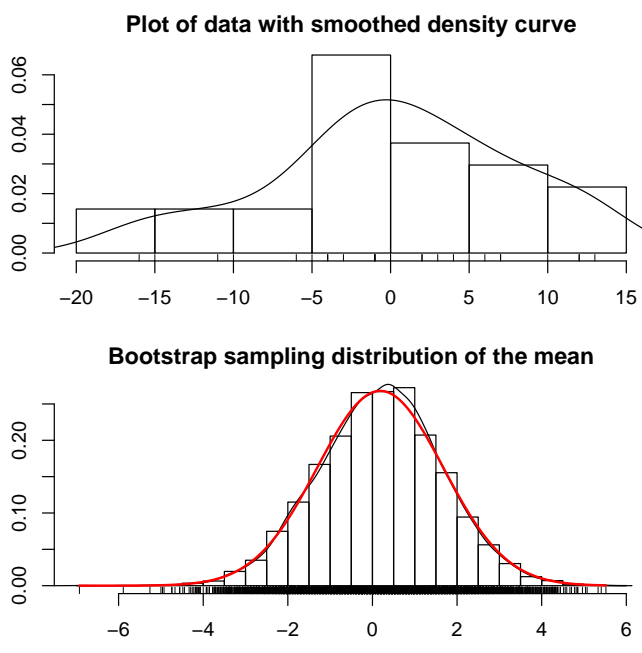
```
bs.one.samp.dist(iq$diff)

# one-sample t-test of differences (paired t-test)
t.summary <- t.test(iq$diff)
t.summary

##
## One Sample t-test
##
## data: iq$diff
## t = 0.1244, df = 26, p-value = 0.902
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
```

```
## -2.875 3.246
## sample estimates:
## mean of x
## 0.1852

# plot t-distribution with shaded p-value
t.dist.pval(t.summary)
```



Alternatively, I can generate the test and CI directly from the raw data in two columns, specifying `paired=TRUE`. This gives the following output, which leads to identical conclusions to the earlier analysis.

```
# two-sample paired t-test
t.summary <- t.test(iq$genetic, iq$foster, paired=TRUE)
t.summary

##
## Paired t-test
##
## data: iq$genetic and iq$foster
## t = 0.1244, df = 26, p-value = 0.902
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -2.875 3.246
## sample estimates:
## mean of the differences
## 0.1852
```

You might ask why I tortured you by doing the first analysis, which required creating and analyzing the sample of differences, when the alternative and equivalent second analysis is so much easier. (A later topic deals with

non-parametric analyses of paired data for which the differences must be first computed.)

Remark: I could have defined the difference to be the foster IQ score minus the genetic IQ score. How would this change the conclusions?

Example: Paired Comparisons of Two Sleep Remedies The following data give the amount of sleep gained in hours from two sleep remedies, A and B, applied to 10 individuals who have trouble sleeping an adequate amount. Negative values imply sleep loss. In 9 of the 10 individuals, the sleep gain on B exceeded that on A.

Let μ_A = population mean sleep gain (among troubled sleepers) on remedy A, and μ_B = population mean sleep gain (among troubled sleepers) on remedy B. Consider testing $H_0 : \mu_B - \mu_A = 0$ or equivalently $\mu_d = 0$, where $\mu_d = \mu_B - \mu_A$.

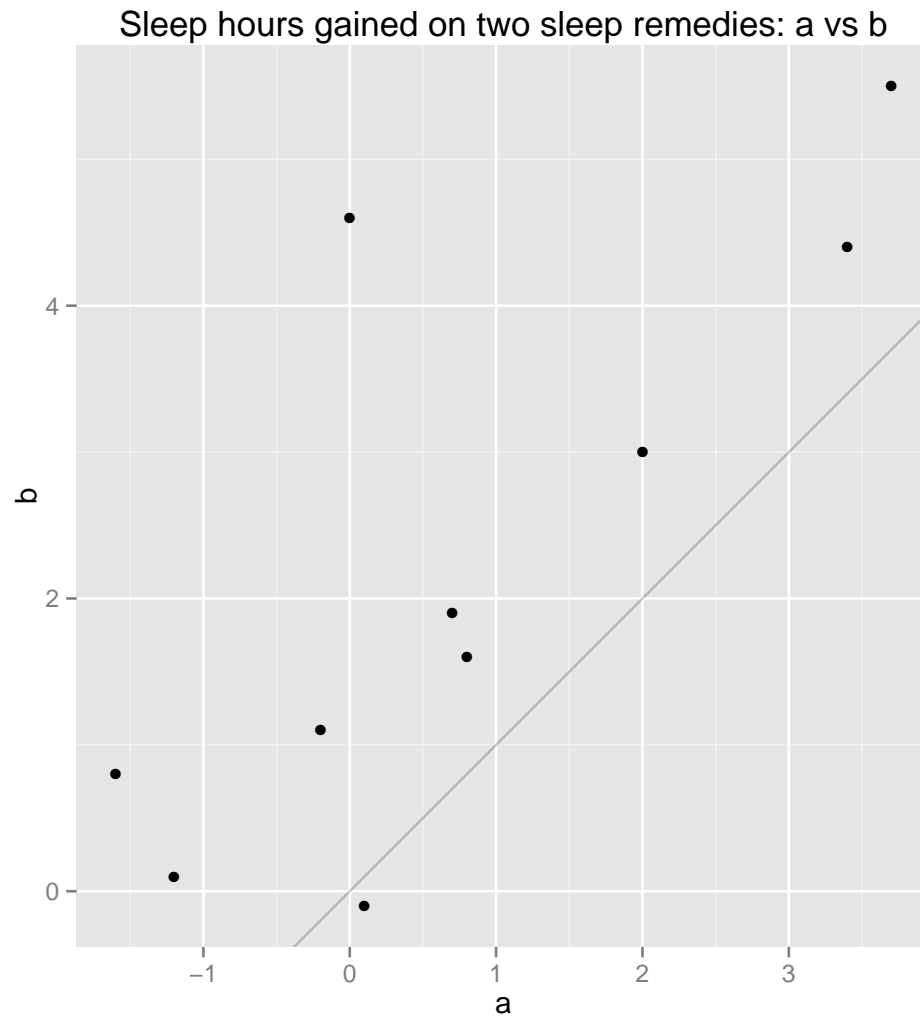
The observed distribution of differences between B and A is slightly skewed to the right, with a single outlier in the upper tail. The normality assumption of the standard one-sample t -test and CI are suspect here. I will continue with the analysis, anyways.

```
#### Example: Paired Comparisons of Two Sleep Remedies
# Data and numerical summaries
a <- c( 0.7, -1.6, -0.2, -1.2,  0.1,  3.4,  3.7,  0.8,  0.0,  2.0)
b <- c( 1.9,  0.8,  1.1,  0.1, -0.1,  4.4,  5.5,  1.6,  4.6,  3.0)
d <- b - a;
sleep <- data.frame(a, b, d)
sleep

##      a     b     d
## 1  0.7  1.9  1.2
## 2 -1.6  0.8  2.4
## 3 -0.2  1.1  1.3
## 4 -1.2  0.1  1.3
## 5  0.1 -0.1 -0.2
## 6  3.4  4.4  1.0
## 7  3.7  5.5  1.8
## 8  0.8  1.6  0.8
## 9  0.0  4.6  4.6
## 10 2.0   3.0  1.0

# scatterplot of a and b IQs, with 1:1 line
p <- ggplot(sleep, aes(x = a, y = b))
```

```
# draw a 1:1 line, dots above line indicate "b > a"
p <- p + geom_abline(intercept=0, slope=1, alpha=0.2)
p <- p + geom_point()
# make the axes square so it's a fair visual comparison
p <- p + coord_equal()
p <- p + labs(title = "Sleep hours gained on two sleep remedies: a vs b")
print(p)
```



There is evidence here against the normality assumption of the sample mean. We'll continue anyway (in practice we'd use a nonparametric method, instead, in a later chapter).

```
p1 <- ggplot(sleep, aes(x = d))
p1 <- p1 + scale_x_continuous(limits=c(-5,+5))
# vertical line at 0
p1 <- p1 + geom_vline(xintercept=0, colour="#BB0000", linetype="dashed")
p1 <- p1 + geom_histogram(aes(y=..density..)
                          , binwidth=1
                          , colour="black", fill="white")
# Overlay with transparent density plot
p1 <- p1 + geom_density(alpha=0.1, fill="#FF6666")
p1 <- p1 + geom_point(aes(y = -0.01))
```

```

        , position = position_jitter(height = 0.005)
        , alpha = 1/5)
p1 <- p1 + labs(title = "Difference of sleep hours gained: d = b - a")

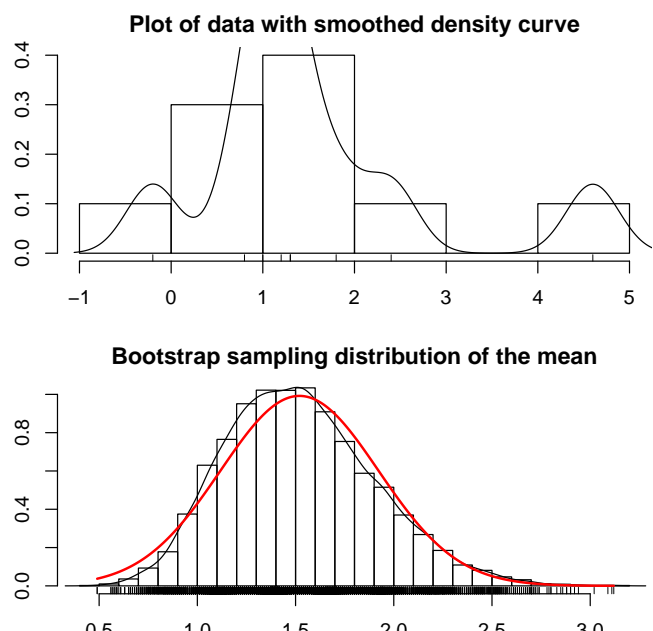
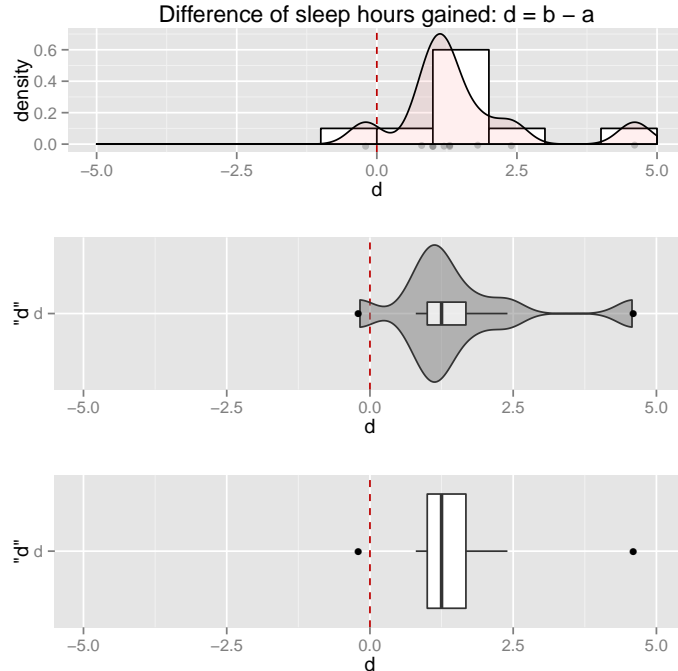
# violin plot
p2 <- ggplot(sleep, aes(x = "d", y = d))
p2 <- p2 + scale_y_continuous(limits=c(-5,+5))
p2 <- p2 + geom_hline(yintercept=0, colour="#BB0000", linetype="dashed")
p2 <- p2 + geom_violin(fill = "gray50", alpha=1/2)
p2 <- p2 + geom_boxplot(width = 0.2, alpha = 3/4)
p2 <- p2 + coord_flip()

# boxplot
p3 <- ggplot(sleep, aes(x = "d", y = d))
p3 <- p3 + scale_y_continuous(limits=c(-5,+5))
p3 <- p3 + geom_hline(yintercept=0, colour="#BB0000", linetype="dashed")
p3 <- p3 + geom_boxplot()
p3 <- p3 + coord_flip()

library(gridExtra)
grid.arrange(p1, p2, p3, ncol=1)

```

```
bs.one.samp.dist(sleep$d)
```



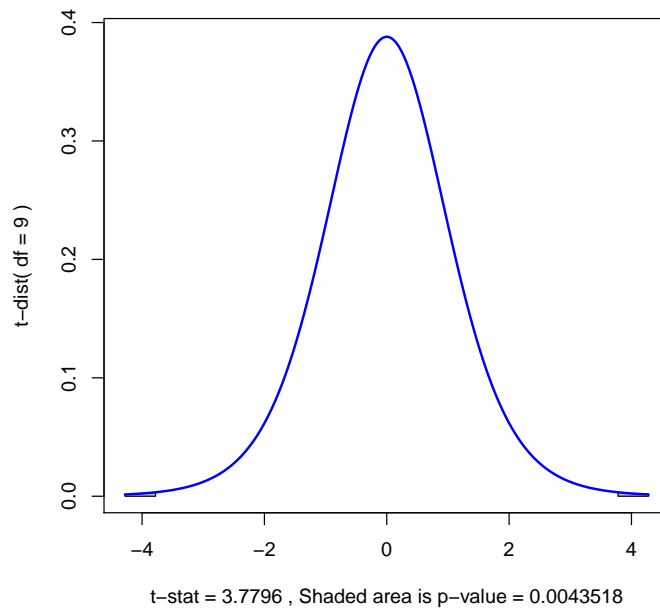
The p-value for testing H_0 is 0.004. We'd reject H_0 at the 5% or 1% level, and conclude that the population mean sleep gains on the remedies are different. We are 95% confident that μ_B exceeds μ_A by between 0.61 and 2.43 hours. Again, these results must be reported with caution, because the normality

assumption is unreasonable. However, the presence of outliers tends to make the *t*-test and CI conservative, so we'd expect to find similar conclusions if we used the nonparametric methods discussed later in the semester.

```
# one-sample t-test of differences (paired t-test)
t.summary <- t.test(sleep$d)
t.summary

##
## One Sample t-test
##
## data: sleep$d
## t = 3.78, df = 9, p-value = 0.004352
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
## 0.6102 2.4298
## sample estimates:
## mean of x
## 1.52

# plot t-distribution with shaded p-value
t.dist.pval(t.summary)
```



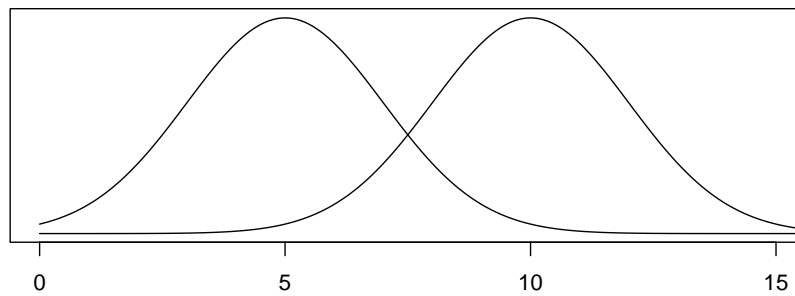
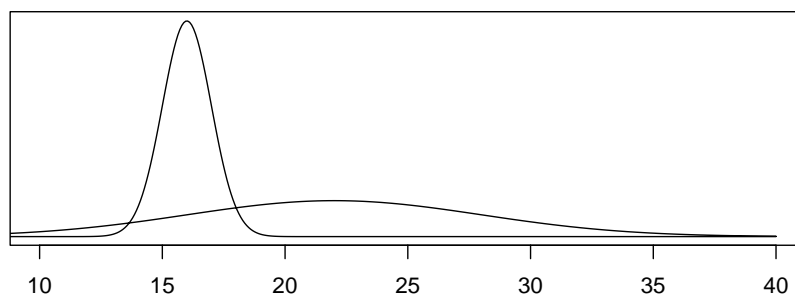
Question: In what order should the remedies be given to the patients?



3.7 Should You Compare Means?

The mean is the most common feature on which two distributions are compared. You should not, however, blindly apply the two-sample tests (paired or unpaired) without asking yourself whether the means are the relevant feature to compare. This issue is not a big concern when, as highlighted in the first graph below, the two (normal) populations have equal spreads or standard deviations. In such cases the difference between the two population means is equal to the difference between any fixed percentile for the two distributions, so the mean difference is a natural measure of difference.

Consider instead the hypothetical scenario depicted in the bottom pane below, where the population mean lifetimes using two distinct drugs for a fatal disease are $\mu_1 = 16$ months from time of diagnosis and $\mu_2 = 22$ months from time of diagnosis, respectively. The standard deviations under the two drugs are $\sigma_1 = 1$ and $\sigma_2 = 6$, respectively. The second drug has the higher mean lifetime, but at the expense of greater risk. For example, the first drug gives you a 97.7% chance of living at least 14 months, whereas the second drug only gives you a 90.8% chance of living at least 14 months. Which drug is best? It depends on what is important to you, a higher expected lifetime or a lower risk of dying early.

Normal Distributions with Identical Variances**Normal Distributions with Different Variances**

Chapter 4

Checking Assumptions

Learning objectives

After completing this topic, you should be able to:

assess the assumptions visually and via formal tests.

Achieving these goals contributes to mastery in these course learning outcomes:

10. Model assumptions.

4.1 Introduction

Almost all statistical methods make assumptions about the data collection process and the shape of the population distribution. If you reject the null hypothesis in a test, then a reasonable conclusion is that the null hypothesis is false, provided all the distributional assumptions made by the test are satisfied. If the assumptions are not satisfied then that alone might be the cause of rejecting H_0 . Additionally, if you fail to reject H_0 , that could be caused solely by failure to satisfy assumptions also. Hence, you should always check assumptions to the best of your abilities.

Two assumptions that underly the tests and CI procedures that I have discussed are that the data are a random sample, and that the population frequency curve is normal. For the pooled variance two-sample test the population variances are also required to be equal.

The random sample assumption can often be assessed from an understanding of the data collection process. Unfortunately, there are few general tests for

checking this assumption. I have described exploratory (mostly visual) methods to assess the normality and equal variance assumption. I will now discuss formal methods to assess these assumptions.

4.2 Testing Normality

An informal test of normality can be based on a **normal scores plot**, sometimes called a **rankit plot** or a **normal probability plot** or a **normal QQ plot** ($QQ = \text{quantile-quantile}$). You plot the quantiles of the data against the quantiles of the normal distribution, or **expected normal order statistics** (in a standard normal distribution) for a sample with the given number of observations. The normality assumption is plausible if the plot is fairly linear. I give below several plots often seen with real data, and what they indicate about the underlying distribution.

There are multiple ways to produce QQ plots in R. The shape can depend upon whether you plot the normal scores on the x-axis or the y-axis. It is conventional to plot the data on the y -axis and the normal scores on the x -axis.

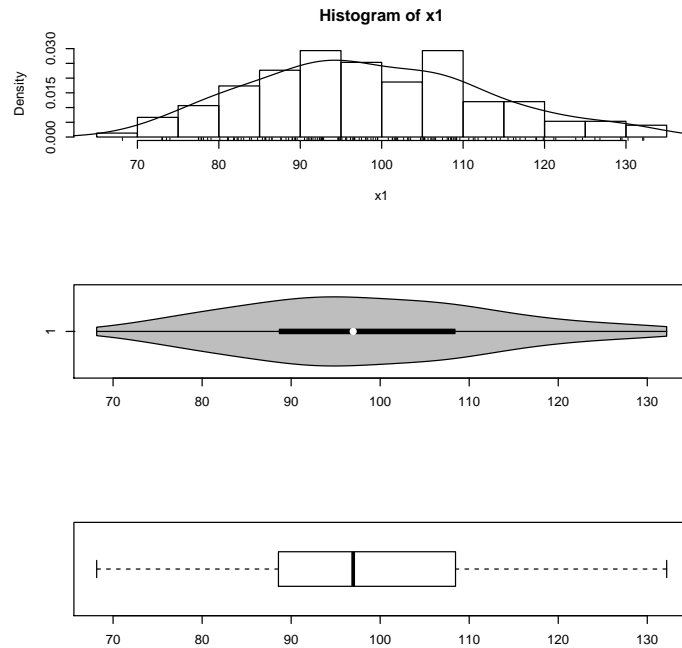
Let's start with some data from a normal distribution.

```
#### sample from normal distribution
x1 <- rnorm(150, mean = 100, sd = 15)

par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(x1, freq = FALSE, breaks = 20)
points(density(x1), type = "l")
rug(x1)

# violin plot
library(vioplot)
vioplot(x1, horizontal=TRUE, col="gray")

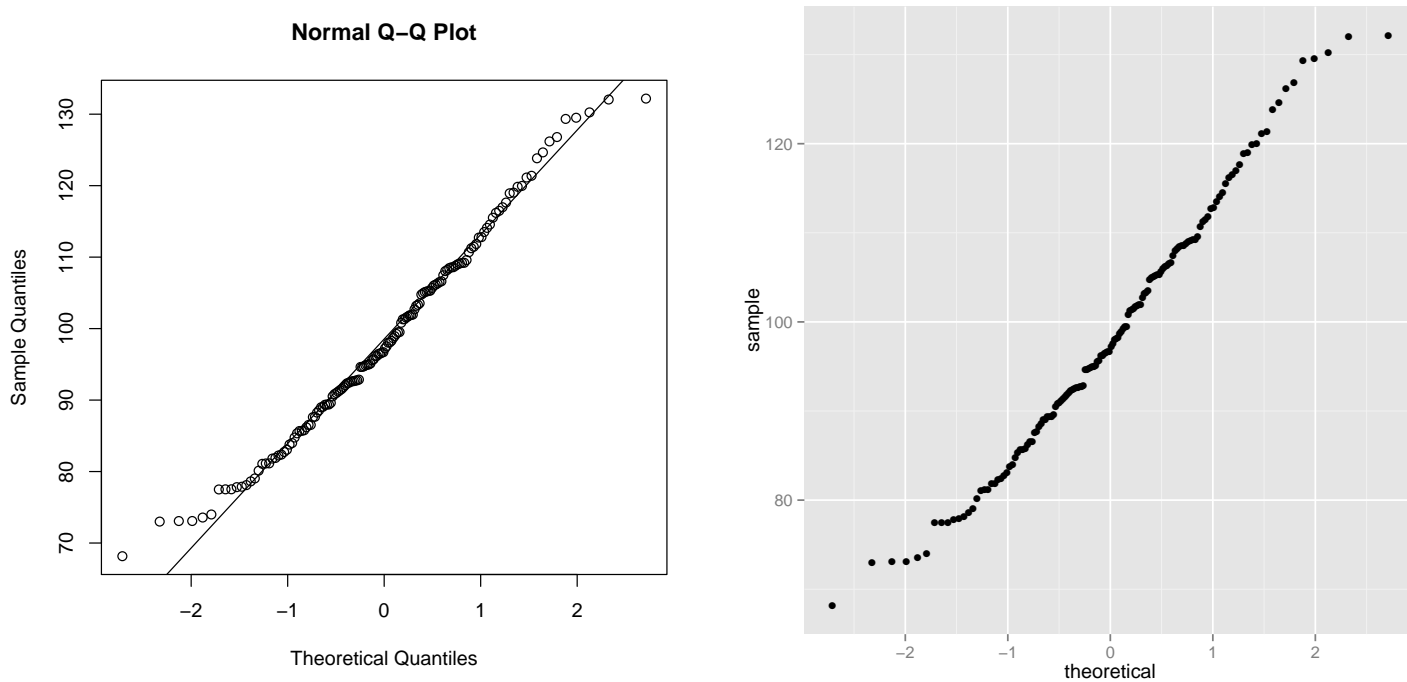
# boxplot
boxplot(x1, horizontal=TRUE)
```



There are many ways to get adequate QQ plots. Consider how outliers shows up in the QQ plot. There may be isolated points on ends of the QQ plot, but only on the right side is there an outlier. How could you have identified that the right tail looks longer than the left tail from the QQ plot?

```
#### QQ plots
# R base graphics
par(mfrow=c(1,1))
# plots the data vs their normal scores
qqnorm(x1)
# plots the reference line
qqline(x1)

# ggplot2 graphics
library(ggplot2)
# http://had.co.nz/ggplot2/stat_qq.html
df <- data.frame(x1)
# stat_qq() below requires "sample" to be assigned a data.frame column
p <- ggplot(df, aes(sample = x1))
# plots the data vs their normal scores
p <- p + stat_qq()
print(p)
```



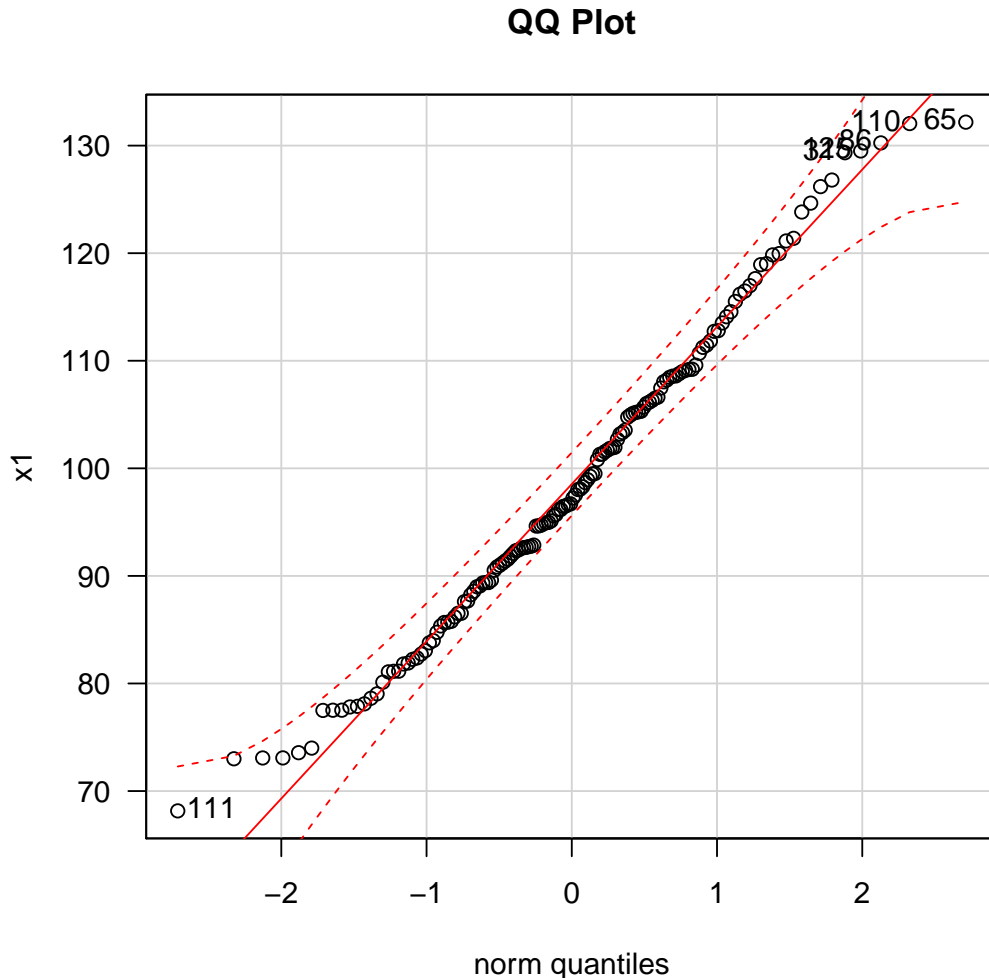
If you lay a straightedge along the bulk of the plot (putting in a regression line is not the right way to do it, even if it is easy), you see that the most extreme point on the right is a little below the line, and the last few points on the left a little above the line. What does this mean? The point on the right corresponds to a data value *more extreme* than expected from a normal distribution (the straight line is where expected and actual coincide). Extreme points on the right are above the line. What about the left? Extreme points there should be *above* the line — since the deviations from the line are above it on the left, those points are also *more extreme* than expected.

Even more useful is to add confidence intervals (point-wise, not family-wise — you will learn the meaning of those terms in the ANOVA section). You don't expect a sample from a normally distributed population to have a normal scores plot that falls exactly on the line, and the amount of deviation depends upon the sample size.

The best QQ plot I could find is available in the `car` package called `qqPlot`. Note that with the `dist=` option you can use this technique to see if the data appear from lots of possible distributions, not just normal.

```
par(mfrow=c(1,1))
# Normality of Residuals
library(car)
# qq plot for studentized resid
#   las      = 1 : turns labels on y-axis to read horizontally
#   id.n    = n : labels n most extreme observations, and outputs to console
#   id.cex = 1 : is the size of those labels
```

```
# lwd      = 1 : line width
qqPlot(x1, las = 1, id.n = 6, id.cex = 1, lwd = 1, main="QQ Plot")
## 65 110 86 125 31 111
## 150 149 148 147 146 1
```



In this case the x -axis is labelled “norm quantiles”. You only see a couple of data values outside the limits (in the tails, where it usually happens). You expect around 5% outside the limits, so there is no indication of non-normality here. I *did* sample from a normal population.

4.2.1 Normality tests on non-normal data

Let’s turn to examples of sampling from other, non-normal distributions to see how the normal QQ plot identifies important features.

Light-tailed symmetric (Uniform)

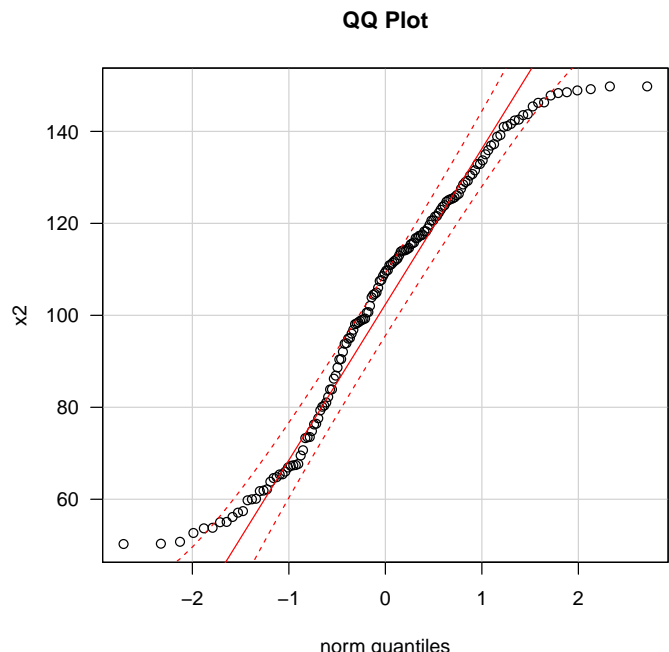
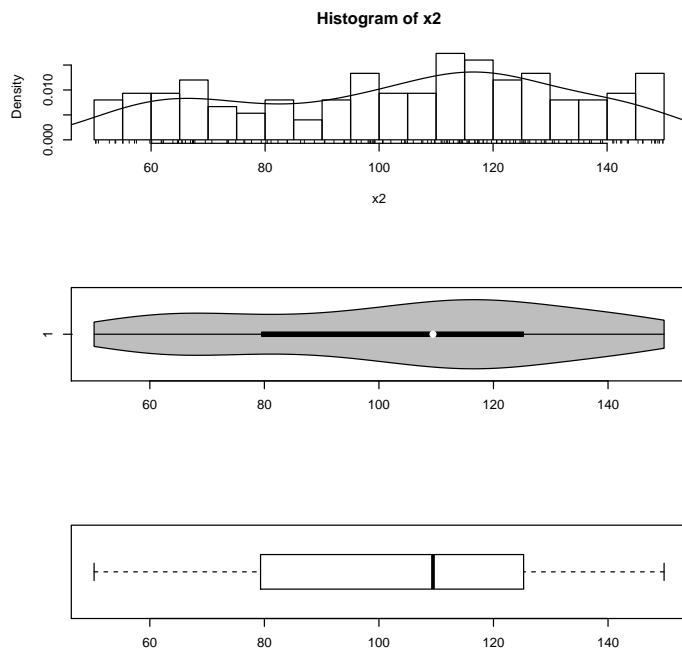
```
#### Light-tailed symmetric (Uniform)
# sample from uniform distribution
x2 <- runif(150, min = 50, max = 150)

par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(x2, freq = FALSE, breaks = 20)
points(density(x2), type = "l")
rug(x2)

# violin plot
library(vioplot)
vioplot(x2, horizontal=TRUE, col="gray")

# boxplot
boxplot(x2, horizontal=TRUE)

par(mfrow=c(1,1))
qqPlot(x2, las = 1, id.n = 0, id.cex = 1, lwd = 1, main="QQ Plot")
```



Heavy-tailed (fairly) symmetric (Normal-squared)

```
#### Heavy-tailed (fairly) symmetric (Normal-squared)
# sample from normal distribution
x3.temp <- rnorm(150, mean = 0, sd = 1)
x3 <- sign(x3.temp)*x3.temp^2 * 15 + 100

par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
```

```

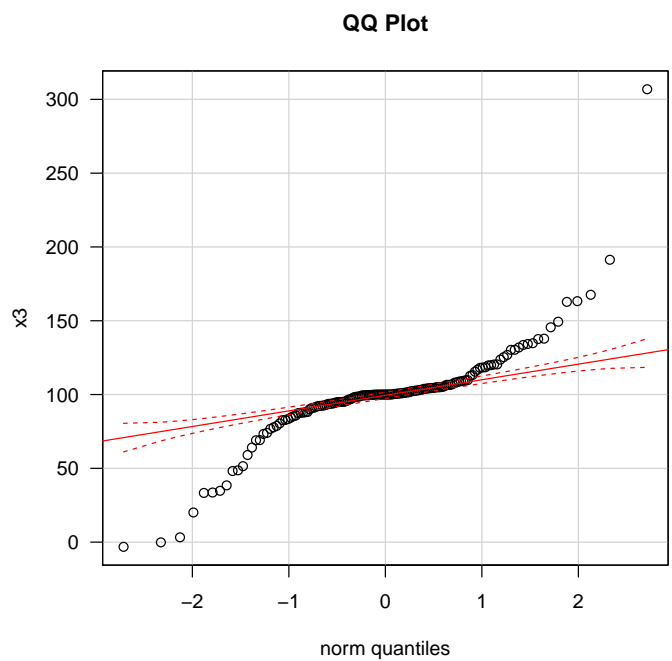
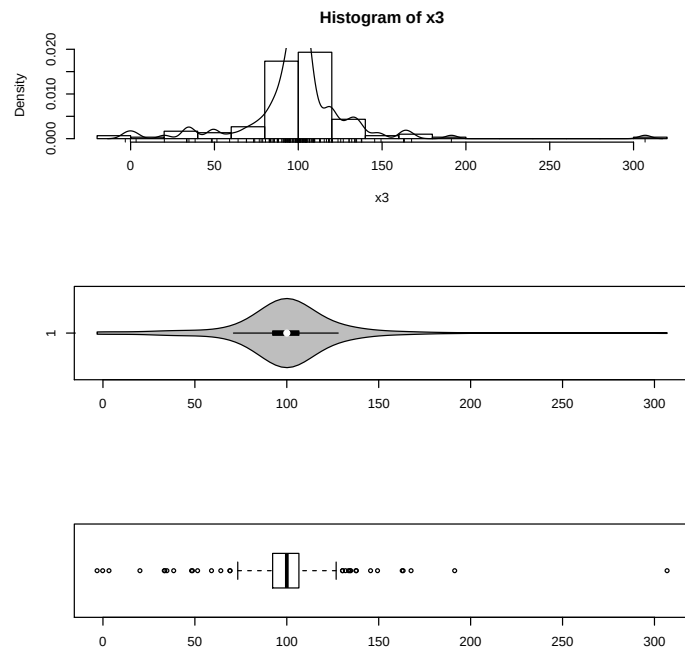
hist(x3, freq = FALSE, breaks = 20)
points(density(x3), type = "l")
rug(x3)

# violin plot
library(vioplot)
vioplot(x3, horizontal=TRUE, col="gray")

# boxplot
boxplot(x3, horizontal=TRUE)

par(mfrow=c(1,1))
qqPlot(x3, las = 1, id.n = 0, id.cex = 1, lwd = 1, main="QQ Plot")

```



Right-skewed (Exponential)

```

##### Right-skewed (Exponential)
# sample from exponential distribution
x4 <- rexp(150, rate = 1)

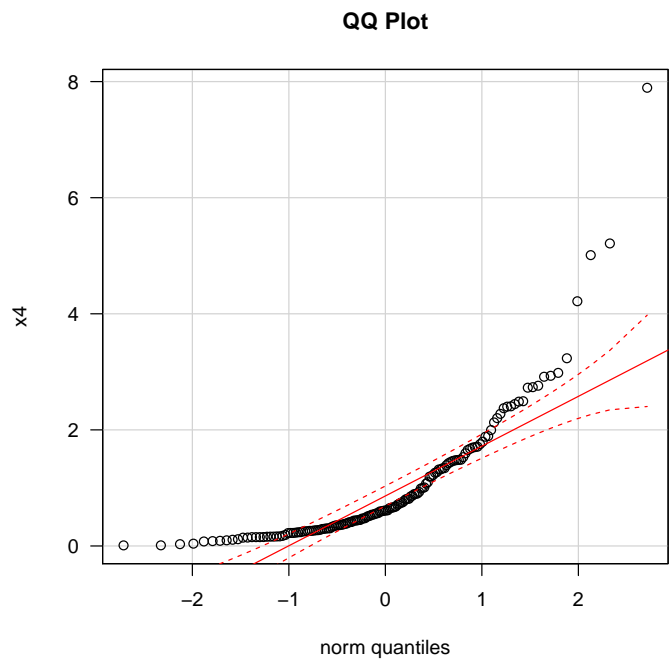
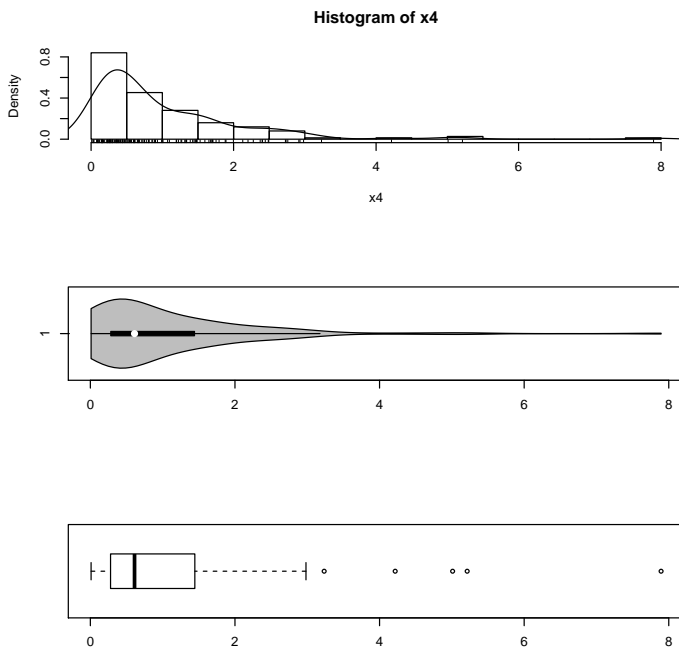
par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(x4, freq = FALSE, breaks = 20)
points(density(x4), type = "l")
rug(x4)

# violin plot
library(vioplot)
vioplot(x4, horizontal=TRUE, col="gray")

```

```
# boxplot
boxplot(x4, horizontal=TRUE)

par(mfrow=c(1,1))
qqPlot(x4, las = 1, id.n = 0, id.cex = 1, lwd = 1, main="QQ Plot")
```



Left-skewed (Exponential, reversed)

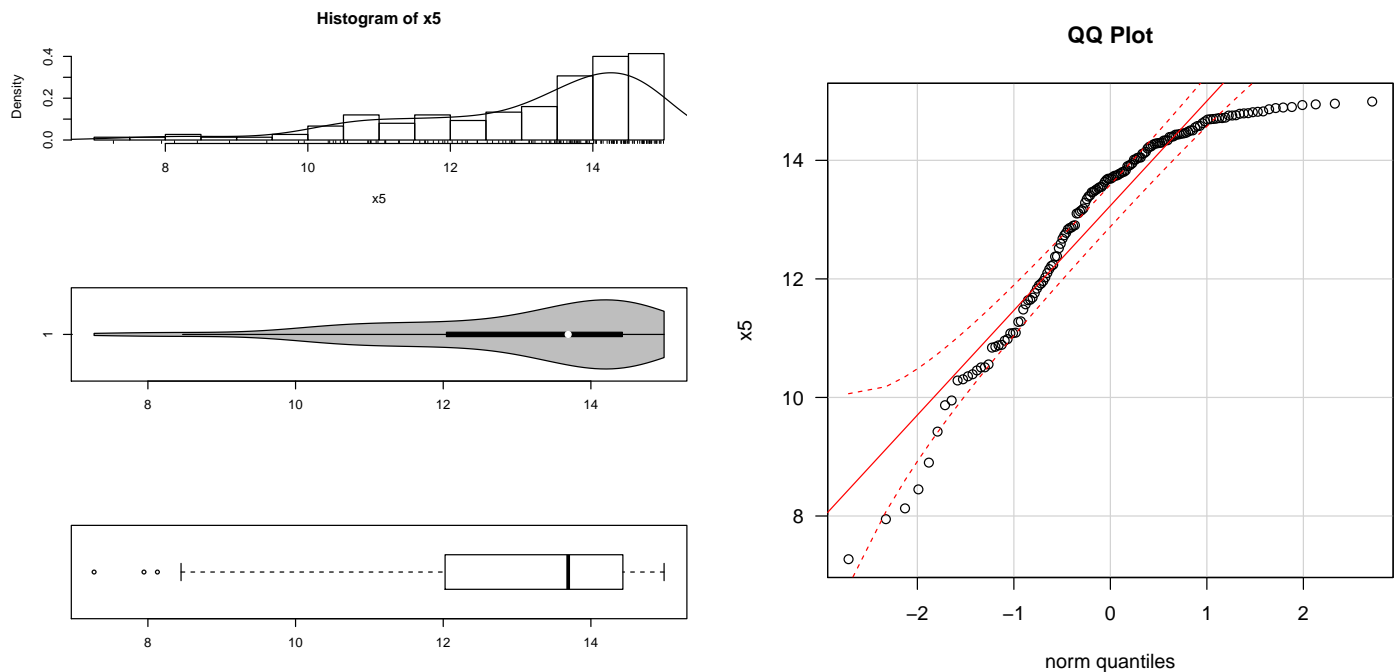
```
#### Left-skewed (Exponential, reversed)
# sample from exponential distribution
x5 <- 15 - rexp(150, rate = 0.5)

par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(x5, freq = FALSE, breaks = 20)
points(density(x5), type = "l")
rug(x5)

# violin plot
library(vioplot)
vioplot(x5, horizontal=TRUE, col="gray")

# boxplot
boxplot(x5, horizontal=TRUE)

par(mfrow=c(1,1))
qqPlot(x5, las = 1, id.n = 0, id.cex = 1, lwd = 1, main="QQ Plot")
```



Notice how striking is the lack of linearity in the QQ plot for all the non-normal distributions, particularly the symmetric light-tailed distribution where the boxplot looks fairly good. The QQ plot is a sensitive measure of normality. Let us summarize the patterns we see regarding tails in the plots:

	Tail	
Tail Weight	Left	Right
Light	Left side of plot points left	Right side of plot points right
Heavy	Left side of plot points down	Right side of plot points up

4.3 Formal Tests of Normality

A formal test of normality is based on the **correlation** between the data and the normal scores. The correlation is a measure of the strength of a linear relationship, with the sign of the correlation indicating the direction of the relationship (that is, + for increasing relationship, and - for decreasing). The correlation varies from -1 to $+1$. In a normal scores plot, you are looking for a correlation close to $+1$. Normality is rejected if the correlation is too small. Critical values for the correlation test of normality, which is commonly called the **Shapiro-Wilk** test, can be found in many texts.

R has several tests of normality. The **Shapiro-Wilk** test `shapiro.test()` is a base function. The R package `nortest` has five others: the Anderson-Darling

test `ad.test()` is useful, related to the **Kolmogorov-Smirnov** (Lilliefors) test `lillie.test()` which is commonly used in many scientific disciplines but is essentially useless, the Cramer-von Mises test `cvm.test()`, and two more. Some packages also have the **Ryan-Joiner** test (closely related to the Shapiro-Wilk test).

Extreme outliers and skewness have the biggest effects on standard methods based on normality. The Shapiro-Wilk (SW) test is better at picking up these problems than the Kolmogorov-Smirnov (KS) test. The KS test tends to highlight deviations from normality in the center of the distribution. These types of deviations are rarely important because they do not have a noticeable effect on the operating characteristics of the standard methods. The AD and RJ tests are modifications designed to handle some of these objections.

Tests for normality may have low power in small to moderate sized samples. Visual assessment of normality is often more valuable than a formal test. The tests for the distributions of data above are below and in Figure 4.1.

Normal distribution

```
#### Formal Tests of Normality
shapiro.test(x1)

##
##  Shapiro-Wilk normality test
##
## data: x1
## W = 0.9858, p-value = 0.1289

library(nortest)
ad.test(x1)

##
##  Anderson-Darling normality test
##
## data: x1
## A = 0.4073, p-value = 0.3446

# lillie.test(x1)
cvm.test(x1)

##
##  Cramer-von Mises normality test
##
## data: x1
## W = 0.0567, p-value = 0.4159
```

Heavy-tailed (fairly) symmetric

Light-tailed symmetric

```
shapiro.test(x2)

##
##  Shapiro-Wilk normality test
##
## data: x2
## W = 0.9525, p-value = 5.336e-05

library(nortest)
ad.test(x2)

##
## Anderson-Darling normality test
##
## data: x2
## A = 1.943, p-value = 5.644e-05

# lillie.test(x2)
cvm.test(x2)

##
## Cramer-von Mises normality test
##
## data: x2
## W = 0.2957, p-value = 0.0003642
```

```
shapiro.test(x3)

##
##  Shapiro-Wilk normality test
##
## data: x3
## W = 0.7963, p-value = 3.587e-13

library(nortest)
ad.test(x3)

##
## Anderson-Darling normality test
##
## data: x3
## A = 9.143, p-value < 2.2e-16

# lillie.test(x3)
cvm.test(x3)

## Warning: p-value is smaller than 7.37e-10,
## cannot be computed more accurately

##
## Cramer-von Mises normality test
##
## data: x3
## W = 1.825, p-value = 7.37e-10
```

Right-skewed

```
shapiro.test(x4)

##
##  Shapiro-Wilk normality test
##
## data: x4
## W = 0.7412, p-value = 5.872e-15

library(nortest)
ad.test(x4)

##
## Anderson-Darling normality test
##
## data: x4
## A = 9.371, p-value < 2.2e-16

# lillie.test(x4)
cvm.test(x4)

## Warning: p-value is smaller than 7.37e-10,
## cannot be computed more accurately

##
## Cramer-von Mises normality test
##
## data: x4
## W = 1.654, p-value = 7.37e-10
```

Left-skewed

```
shapiro.test(x5)

##
##  Shapiro-Wilk normality test
##
## data: x5
## W = 0.8743, p-value = 5.933e-10

library(nortest)
ad.test(x5)

##
## Anderson-Darling normality test
##
## data: x5
## A = 6.002, p-value = 7.938e-15

# lillie.test(x5)
cvm.test(x5)

##
## Cramer-von Mises normality test
##
## data: x5
## W = 1.055, p-value = 9.648e-10
```

Figure 4.1: Normality tests for non-normal distributions

Example: Paired Differences on Sleep Remedies The following boxplot and normal scores plots suggest that the underlying distribution of differences (for the paired sleep data taken from the previous chapter) is reasonably symmetric, but heavy tailed. The p-value for the SW test of normality is 0.042, and for the AD test is 0.029, both of which call into question a normality assumption. A non-parametric test comparing the sleep remedies (one that does not assume normality) is probably more appropriate here. We will return to these data later.

```
# Normality tests
shapiro.test(sleep$d)

##
##  Shapiro-Wilk normality test
##
## data: sleep$d
## W = 0.838, p-value = 0.04173

library(nortest)
ad.test(sleep$d)

##
##  Anderson-Darling normality test
##
## data: sleep$d
## A = 0.7738, p-value = 0.02898

# lillie.test(sleep$d)
cvm.test(sleep$d)

##
##  Cramer-von Mises normality test
##
## data: sleep$d
## W = 0.1382, p-value = 0.02769

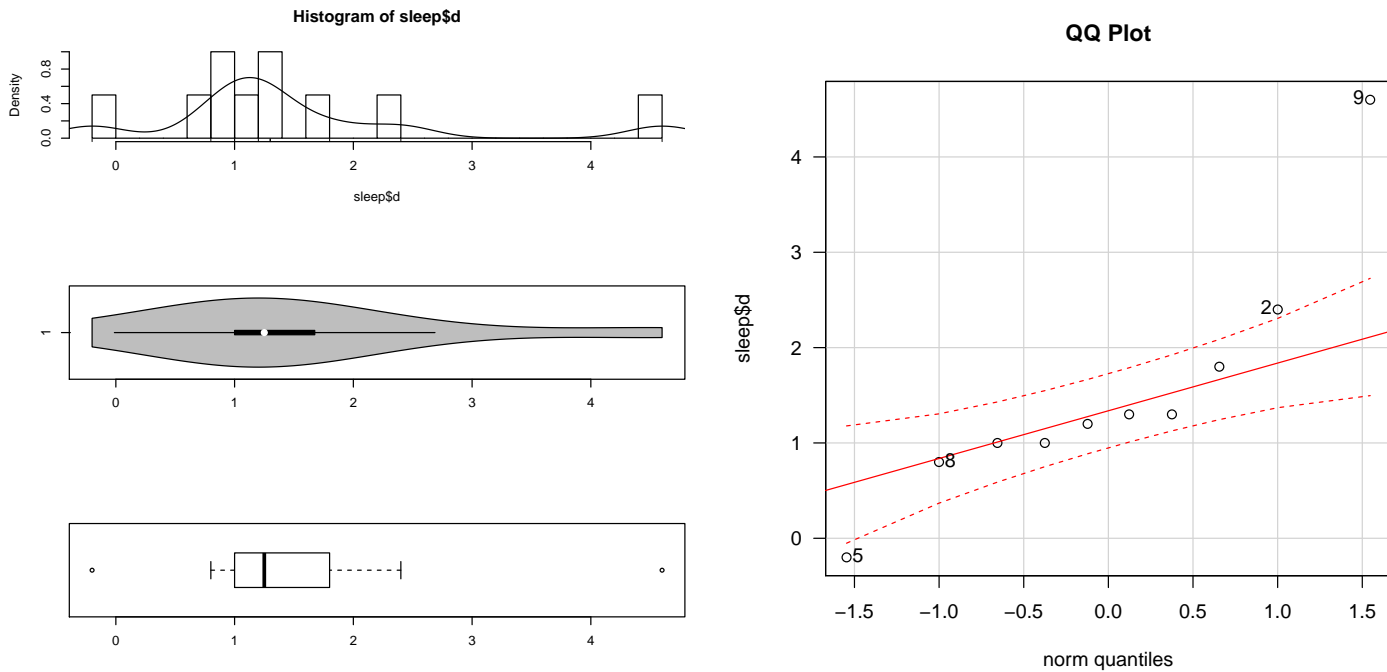
# plot of data
par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(sleep$d, freq = FALSE, breaks = 20)
points(density(sleep$d), type = "l")
rug(sleep$d)

# violin plot
library(vioplot)
vioplot(sleep$d, horizontal=TRUE, col="gray")

# boxplot
boxplot(sleep$d, horizontal=TRUE)

# QQ plot
```

```
par(mfrow=c(1,1))
qqPlot(sleep$d, las = 1, id.n = 4, id.cex = 1, lwd = 1, main="QQ Plot")
##  9  5  2  8
## 10  1  9  2
```



Example: Androstenedione Levels This is an independent two-sample problem, so you must look at normal scores plots for males and females.

The AD test p-value and the SW test p-value for testing normality exceeds 0.10 in each sample. Thus, given the sample sizes (14 for men, 18 for women), we have insufficient evidence (at $\alpha = 0.05$) to reject normality in either population.

The women's boxplot contains two mild outliers, which is highly unusual when sampling from a normal distribution. The tests are possibly not powerful enough to pick up this type of deviation from normality in such a small sample. In practice, this may not be a big concern. The two mild outliers probably have a small effect on inferences in the sense that non-parametric methods would probably lead to similar conclusions here.

```
p1 <- ggplot(andro, aes(x = sex, y = level, fill=sex))
p1 <- p1 + geom_boxplot()
# add a "+" at the mean
p1 <- p1 + stat_summary(fun.y = mean, geom = "point", shape = 3, size = 2)
#p1 <- p1 + coord_flip()
p1 <- p1 + labs(title = "Androstenedione Levels in Diabetics")
#print(p1)
```

Men

```

shapiro.test(men)

##
##  Shapiro-Wilk normality test
##
## data: men
## W = 0.9059, p-value = 0.1376

library(nortest)
ad.test(men)

##
## Anderson-Darling normality test
##
## data: men
## A = 0.4718, p-value = 0.2058

# lillie.test(men)
cvm.test(men)

##
## Cramer-von Mises normality test
##
## data: men
## W = 0.0631, p-value = 0.3221

```

Women

```

shapiro.test(women)

##
## Shapiro-Wilk normality test
##
## data: women
## W = 0.9598, p-value = 0.5969

library(nortest)
ad.test(women)

##
## Anderson-Darling normality test
##
## data: women
## A = 0.3947, p-value = 0.3364

# lillie.test(women)
cvm.test(women)

##
## Cramer-von Mises normality test
##
## data: women
## W = 0.0652, p-value = 0.3057

```

```

p2 <- ggplot(andro, aes(x = level, fill=sex))
p2 <- p2 + geom_histogram(binwidth = 20, alpha = 0.5, position="identity")
p2 <- p2 + labs(title = "Androstenedione Levels in Diabetics")
#print(p2)

```

```

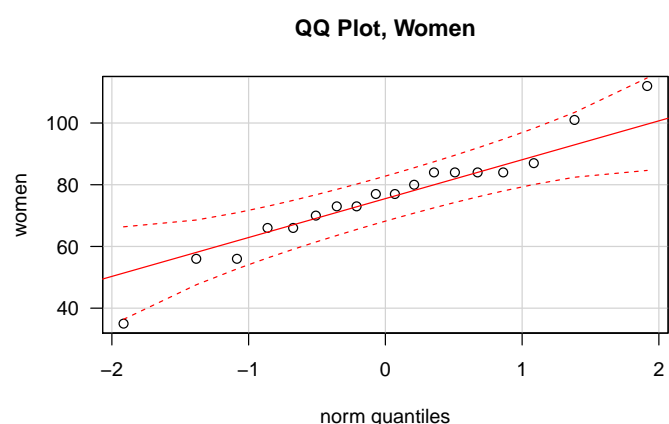
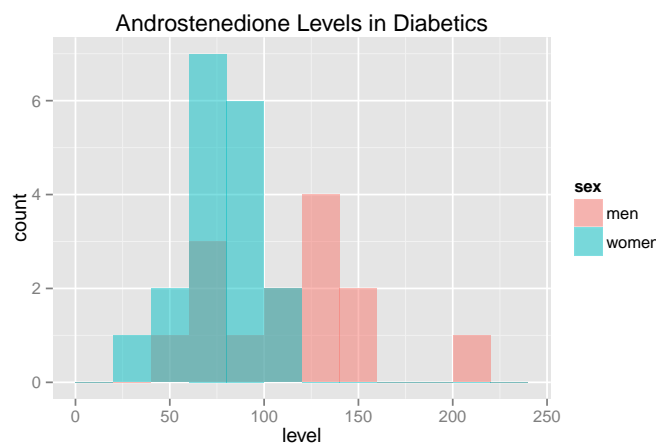
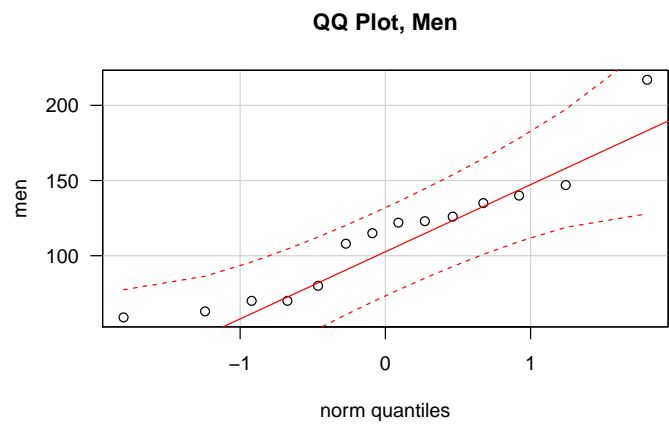
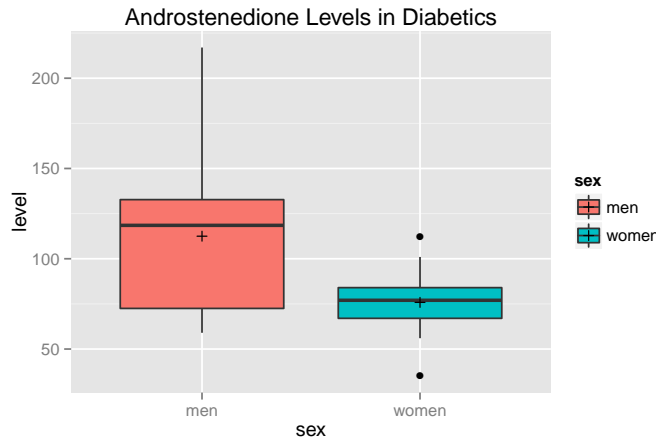
library(gridExtra)
grid.arrange(p1, p2, ncol=1)

```

```

# QQ plot
par(mfrow=c(2,1))
qqPlot(men, las = 1, id.n = 0, id.cex = 1, lwd = 1, main="QQ Plot, Men")
qqPlot(women, las = 1, id.n = 0, id.cex = 1, lwd = 1, main="QQ Plot, Women")

```



Most statisticians use graphical methods (boxplot, normal scores plot) to assess normality, and do not carry out formal tests.

4.4 Testing Equal Population Variances

In the independent two sample t -test, some researchers test $H_0 : \sigma_1^2 = \sigma_2^2$ as a means to decide between using the pooled-variance procedure or Satterthwaite's methods. They suggest the pooled t -test and CI if H_0 is not rejected, and Satterthwaite's methods otherwise.

There are a number of well-known tests for equal population variances, of which Bartlett's test and Levene's test are probably the best known. Bartlett's test assumes the population distributions are normal, and is the best test when this is true. In practice, unequal variances and non-normality often go hand-in-hand, so you should check normality prior to using Bartlett's test. It is sensitive to data which is not non-normally distributed, thus it is more likely to return a "false positive" (reject H_0 of equal variances) when the data is non-normal. Levene's test is more robust to departures from normality than Bartlett's test;

it is in the `car` package. Fligner-Killeen test is a non-parametric test which is very robust against departures from normality.

I will now define **Bartlett's test**, which assumes normally distributed data. As above, let $n^* = n_1 + n_2 + \dots + n_k$, where the n_i s are the sample sizes from the k groups, and define

$$v = 1 + \frac{1}{3(k-1)} \left(\sum_{i=1}^k \frac{1}{n_i - 1} - \frac{1}{n^* - k} \right).$$

Bartlett's statistic for testing $H_0 : \sigma_1^2 = \dots = \sigma_k^2$ is given by

$$B_{obs} = \frac{2.303}{v} \left\{ (n - k) \log(s_{\text{pooled}}^2) - \sum_{i=1}^k (n_i - 1) \log(s_i^2) \right\},$$

where s_{pooled}^2 is the pooled estimator of variance and s_i^2 is the estimated variance based on the i^{th} sample.

Large values of B_{obs} suggest that the population variances are unequal. For a size α test, we reject H_0 if $B_{\text{obs}} \geq \chi_{k-1, \text{crit}}^2$, where $\chi_{k-1, \text{crit}}^2$ is the upper- α percentile for the χ_{k-1}^2 (chi-squared) probability distribution with $k - 1$ degrees of freedom. A generic plot of the χ^2 distribution is given below. A p-value for the test is given by the area under the chi-squared curve to the right of B_{obs} .

Example: Androstenedione Levels The sample standard deviations and samples sizes are: $s_1 = 42.8$ and $n_1 = 14$ for men and $s_2 = 17.2$ and $n_2 = 18$ for women. The sample standard deviations appear to be very different, so I would not be surprised if the test of equal population variances is highly significant. The output below confirms this: the p-values for Bartlett's F-test, Levene's Test, and Fligner-Killeen test are all much smaller than 0.05. An implication is that the standard pooled-CI and test on the population means is inappropriate.

```
#### Testing Equal Population Variances
# numerical summaries
c(mean(men), mean(women), sd(men), sd(women))
## [1] 112.50 75.83 42.75 17.24
c(IQR(men), IQR(women), length(men), length(women))
```

```
## [1] 60.25 17.00 14.00 18.00

## Test equal variance
# assumes populations are normal
bartlett.test(level ~ sex, data = andro)

##
##  Bartlett test of homogeneity of variances
##
## data: level by sex
## Bartlett's K-squared = 11.2, df = 1, p-value = 0.0008183

# does not assume normality, requires car package
library(car)
leveneTest(level ~ sex, data = andro)

## Levene's Test for Homogeneity of Variance (center = median)
##          Df F value Pr(>F)
## group    1   7.2  0.012 *
##            30
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

# nonparametric test
fligner.test(level ~ sex, data = andro)

##
## Fligner-Killeen test of homogeneity of variances
##
## data: level by sex
## Fligner-Killeen:med chi-squared = 5.892, df = 1, p-value =
## 0.01521
```

Chapter 5

One-Way Analysis of Variance

Learning objectives

After completing this topic, you should be able to:

select graphical displays that meaningfully compare independent populations.

assess the assumptions of the ANOVA visually and by formal tests.

decide whether the means between populations are different, and how.

Achieving these goals contributes to mastery in these course learning outcomes:

1. organize knowledge.
5. define parameters of interest and hypotheses in words and notation.
6. summarize data visually, numerically, and descriptively.
8. use statistical software.
12. make evidence-based decisions.

5.1 ANOVA

The one-way analysis of variance (**ANOVA**) is a generalization of the two sample t -test to $k \geq 2$ groups. Assume that the populations of interest have the following (unknown) population means and standard deviations:

	population 1	population 2	...	population k
mean	μ_1	μ_2	...	μ_k
std dev	σ_1	σ_2	...	σ_k

A usual interest in ANOVA is whether $\mu_1 = \mu_2 = \dots = \mu_k$. If not, then we wish to know which means differ, and by how much. To answer these questions we select samples from each of the k populations, leading to the following data summary:

	sample 1	sample 2	...	sample k
size	n_1	n_2	...	n_k
mean	\bar{Y}_1	\bar{Y}_2	...	\bar{Y}_k
std dev	s_1	s_2	...	s_k

A little more notation is needed for the discussion. Let Y_{ij} denote the j^{th} observation in the i^{th} sample and define the total sample size $n^* = n_1 + n_2 + \dots + n_k$. Finally, let $\bar{\bar{Y}}$ be the average response over all samples (combined), that is

$$\bar{\bar{Y}} = \frac{\sum_{ij} Y_{ij}}{n^*} = \frac{\sum_i n_i \bar{Y}_i}{n^*}.$$

Note that $\bar{\bar{Y}}$ is *not* the average of the sample means, unless the samples sizes n_i are equal.

An F -statistic is used to test $H_0 : \mu_1 = \mu_2 = \dots = \mu_k$ against $H_A : \text{not } H_0$ (that is, at least two means are different). The assumptions needed for the standard ANOVA F -test are analogous to the independent pooled two-sample t -test assumptions: (1) Independent random samples from each population. (2) The population frequency curves are normal. (3) The populations have equal standard deviations, $\sigma_1 = \sigma_2 = \dots = \sigma_k$.

The F -test is computed from the ANOVA table, which breaks the spread in the combined data set into two components, or **Sums of Squares** (SS). The **Within SS**, often called the **Residual SS** or the **Error SS**, is the portion of the total spread due to variability *within* samples:

$$\text{SS(Within)} = (n_1 - 1)s_1^2 + (n_2 - 1)s_2^2 + \dots + (n_k - 1)s_k^2 = \sum_{ij} (Y_{ij} - \bar{Y}_i)^2.$$

The **Between SS**, often called the Model SS, measures the spread between the sample means

$$\begin{aligned} \text{SS(Between)} &= \\ n_1(\bar{Y}_1 - \bar{\bar{Y}})^2 + n_2(\bar{Y}_2 - \bar{\bar{Y}})^2 + \dots + n_k(\bar{Y}_k - \bar{\bar{Y}})^2 &= \sum_i n_i (\bar{Y}_i - \bar{\bar{Y}})^2, \end{aligned}$$

weighted by the sample sizes. These two SS add to give

$$\text{SS}(\text{Total}) = \text{SS}(\text{Between}) + \text{SS}(\text{Within}) = \sum_{ij} (Y_{ij} - \bar{\bar{Y}})^2.$$

Each SS has its own degrees of freedom (df). The $df(\text{Between})$ is the number of groups minus one, $k - 1$. The $df(\text{Within})$ is the total number of observations minus the number of groups: $(n_1 - 1) + (n_2 - 1) + \dots + (n_k - 1) = n^* - k$. These two df add to give $df(\text{Total}) = (k - 1) + (n^* - k) = n^* - 1$.

The Sums of Squares and df are neatly arranged in a table, called the ANOVA table:

Source	df	SS	MS	F
Between Groups (Model)	$dfM = k - 1$	$SSM = \sum_i n_i (\bar{Y}_i - \bar{\bar{Y}})^2$	$MSM = SSM/dfM$	MSM/MSE
Within Groups (Error)	$dfE = n^* - k$	$SSE = \sum_i (n_i - 1)s_i^2$	$MSE = SSE/dfE$	
Total	$dfT = n^* - 1$	$SST = \sum_{ij} (Y_{ij} - \bar{\bar{Y}})^2$	$MST = SST/dfT$	

The Mean Square for each source of variation is the corresponding SS divided by its df . The Mean Squares can be easily interpreted.

The MS(Within)

$$\text{MS}(\text{Within}) = \frac{(n_1 - 1)s_1^2 + (n_2 - 1)s_2^2 + \dots + (n_k - 1)s_k^2}{n^* - k} = s_{\text{pooled}}^2$$

is a weighted average of the sample variances. The MS(Within) is known as the pooled estimator of variance, and estimates the assumed common population variance. If all the sample sizes are equal, the MS(Within) is the average sample variance. The MS(Within) is identical to the **pooled variance estimator** in a two-sample problem when $k = 2$.

The MS(Between)

$$\text{MS}(\text{Between}) = \frac{\sum_i n_i (\bar{Y}_i - \bar{\bar{Y}})^2}{k - 1}$$

is a measure of variability among the sample means. This MS is a multiple of the sample variance of $\bar{Y}_1, \bar{Y}_2, \dots, \bar{Y}_k$ when all the sample sizes are equal.

The MS(Total)

$$\text{MS}(\text{Total}) = \frac{\sum_{ij} (Y_{ij} - \bar{\bar{Y}})^2}{n^* - 1}$$

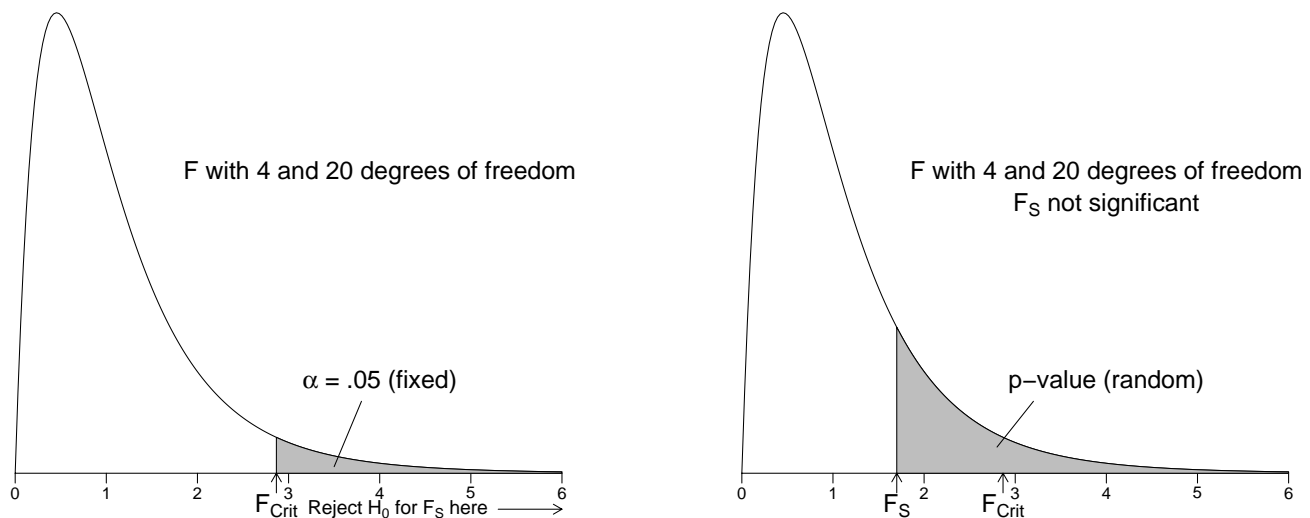
is the variance in the combined data set.

The decision on whether to reject $H_0 : \mu_1 = \mu_2 = \dots = \mu_k$ is based on the ratio of the MS(Between) and the MS(Within):

$$F_s = \frac{\text{MS(Between)}}{\text{MS(Within)}}.$$

Large values of F_s indicate large variability among the sample means $\bar{Y}_1, \bar{Y}_2, \dots, \bar{Y}_k$ relative to the spread of the data within samples. That is, large values of F_s suggest that H_0 is false.

Formally, for a size α test, reject H_0 if $F_s \geq F_{crit}$, where F_{crit} is the upper- α percentile from an F distribution with numerator degrees of freedom $k - 1$ and denominator degrees of freedom $n^* - k$ (i.e., the df for the numerators and denominators in the F -ratio). The p-value for the test is the area under the F -probability curve to the right of F_s :



For $k = 2$ the ANOVA F -test is equivalent to the pooled two-sample t -test.

The specification of a one-way analysis of variance is analogous to a regression analysis (discussed later, though we've seen the specification in some plotting functions). The only difference is that the descriptive (x) variable needs to be a factor and not a numeric variable. We calculate a model object using `lm()` and extract the analysis of variance table with `anova()`.

Example: Comparison of Fats During cooking, doughnuts absorb fat in various amounts. A scientist wished to learn whether the amount absorbed depends on the type of fat. For each of 4 fats, 6 batches of 24 doughnuts were prepared. The data are grams of fat absorbed per batch (minus 100).

Let

μ_i = pop mean grams of fat i absorbed per batch of 24 doughnuts (-100).

The scientist wishes to test $H_0 : \mu_1 = \mu_2 = \mu_3 = \mu_4$ against $H_A : \text{not } H_0$. There is no strong evidence against normality here. Furthermore the sample standard deviations (see output below) are close. The standard ANOVA appears to be appropriate here.

Row	fat1	fat2	fat3	fat4
1	64	78	75	55
2	72	91	86	66
3	68	97	78	49
4	77	82	71	64
5	90	85	63	70
6	76	77	76	68

Let's take a short detour to read the wide table and convert it into long format.

R skills: wide to long table format Many functions in R expect data to be in a long format rather than a wide format. Let's use the fat data as an example of how to read a table as text, convert the wide format to long, and then back to wide format.

```
#### Example: Comparison of Fats
fat <- read.table(text="
Row  fat1  fat2  fat3  fat4
 1    64    78    75    55
 2    72    91    86    66
 3    68    97    78    49
 4    77    82    71    64
 5    90    85    63    70
 6    76    77    76    68
", header=TRUE)
fat

##   Row fat1 fat2 fat3 fat4
## 1    1    64    78    75    55
## 2    2    72    91    86    66
## 3    3    68    97    78    49
## 4    4    77    82    71    64
## 5    5    90    85    63    70
## 6    6    76    77    76    68
```

From wide to long: Use `melt()` from the `reshape2` package.

```
#### From wide to long format
library(reshape2)
fat.long <- melt(fat,
  # id.vars: ID variables
  #   all variables to keep but not split apart on
  id.vars=c("Row"),
  # measure.vars: The source columns
  #   (if unspecified then all other variables are measure.vars)
  measure.vars = c("fat1", "fat2", "fat3", "fat4"),
  # variable.name: Name of the destination column identifying each
  #   original column that the measurement came from
  variable.name = "type",
  # value.name: column name for values in table
  value.name = "amount"
)
# naming variables manually, the variable.name and value.name not working 11/2012
names(fat.long) <- c("Row", "type", "amount")
fat.long
##      Row type amount
## 1     1 fat1    64
## 2     2 fat1    72
## 3     3 fat1    68
## 4     4 fat1    77
## 5     5 fat1    90
## 6     6 fat1    76
## 7     1 fat2    78
## 8     2 fat2    91
## 9     3 fat2    97
## 10    4 fat2    82
## 11    5 fat2    85
## 12    6 fat2    77
## 13    1 fat3    75
## 14    2 fat3    86
## 15    3 fat3    78
## 16    4 fat3    71
## 17    5 fat3    63
## 18    6 fat3    76
## 19    1 fat4    55
## 20    2 fat4    66
## 21    3 fat4    49
## 22    4 fat4    64
## 23    5 fat4    70
## 24    6 fat4    68
# or as simple as:
# melt(fat, "Row")
```

If you don't specify variable.name, it will name that column "variable", and if you leave out value.name, it will name that column "value".

From long to wide: Use dcast() from the reshape2 package.

```
##### From long to wide format
fat.wide <- dcast(fat.long, Row ~ type, value.var = "amount")
fat.wide

##   Row fat1 fat2 fat3 fat4
## 1   1    64    78    75    55
## 2   2    72    91    86    66
## 3   3    68    97    78    49
## 4   4    77    82    71    64
## 5   5    90    85    63    70
## 6   6    76    77    76    68
```

Now that we've got our data in the long format, let's return to the ANOVA.

Back to ANOVA: Let's look at the numerical summaries. We've seen other ways of computing these so I'll show you another way.

```
##### Back to ANOVA
# Calculate the mean, sd, n, and se for the four fats

# The plyr package is an advanced way to apply a function to subsets of data
#   "Tools for splitting, applying and combining data"
library(plyr)
# ddply "dd" means the input and output are both data.frames
fat.summary <- ddply(fat.long,
                      "type",
                      function(X) {
                        data.frame(m = mean(X$amount),
                                   s = sd(X$amount),
                                   n = length(X$amount))
                      })
# standard errors
fat.summary$se <- fat.summary$s/sqrt(fat.summary$n)
# individual confidence limits
fat.summary$ci.l <- fat.summary$m - qt(1-.05/2, df=fat.summary$n-1) * fat.summary$se
fat.summary$ci.u <- fat.summary$m + qt(1-.05/2, df=fat.summary$n-1) * fat.summary$se
fat.summary

##   type     m     s     n     se   ci.l   ci.u
## 1 fat1 74.50 9.028 6 3.686 65.03 83.97
## 2 fat2 85.00 7.772 6 3.173 76.84 93.16
## 3 fat3 74.83 7.627 6 3.114 66.83 82.84
## 4 fat4 62.00 8.222 6 3.357 53.37 70.63
```

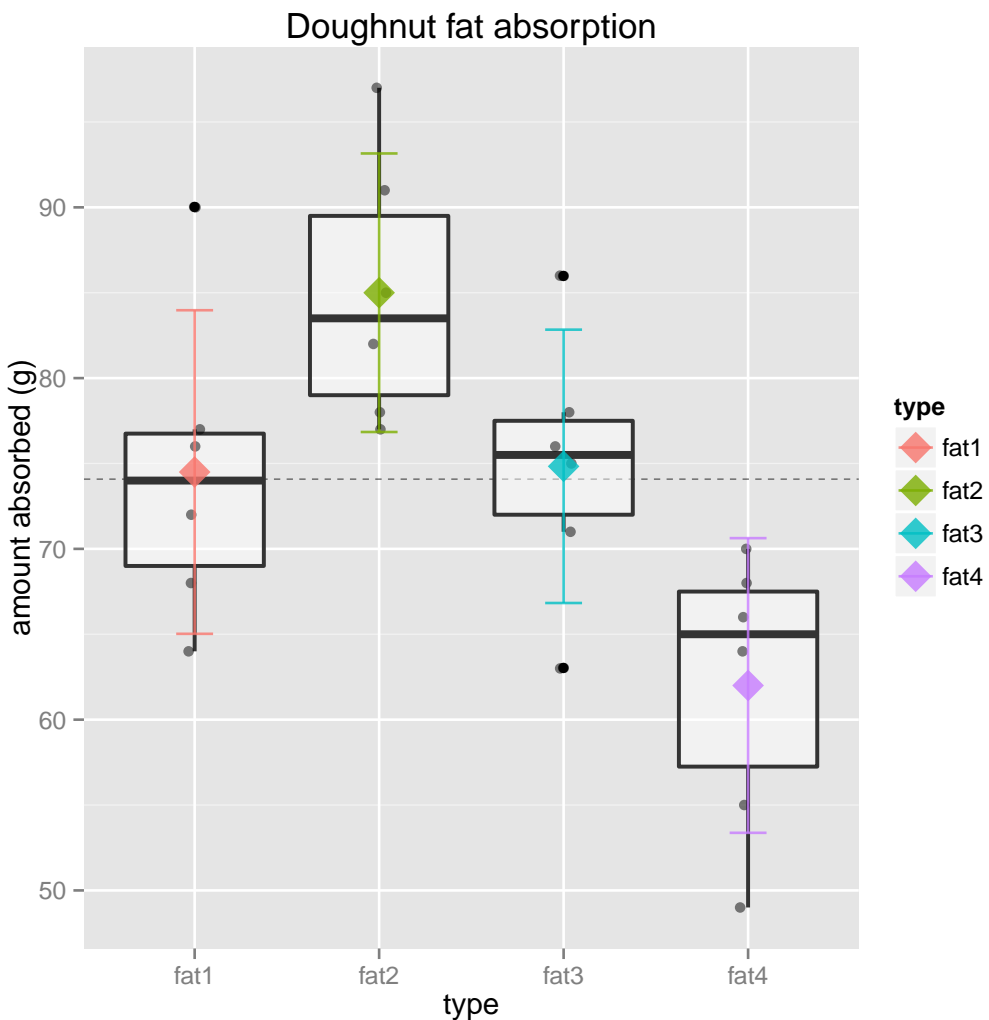
Let's plot the data with boxplots, individual points, mean, and CI by fat type.

```
# Plot the data using ggplot
library(ggplot2)
p <- ggplot(fat.long, aes(x = type, y = amount))
# plot a reference line for the global mean (assuming no groups)
p <- p + geom_hline(yintercept = mean(fat.long$amount),
```

```

            colour = "black", linetype = "dashed", size = 0.3, alpha = 0.5)
# boxplot, size=.75 to stand out behind CI
p <- p + geom_boxplot(size = 0.75, alpha = 0.5)
# points for observed data
p <- p + geom_point(position = position_jitter(w = 0.05, h = 0), alpha = 0.5)
# diamond at mean for each group
p <- p + stat_summary(fun.y = mean, geom = "point", shape = 18, size = 6,
                      aes(colour = type), alpha = 0.8)
# confidence limits based on normal distribution
p <- p + stat_summary(fun.data = "mean_cl_normal", geom = "errorbar",
                      width = .2, aes(colour=type), alpha = 0.8)
p <- p + labs(title = "Doughnut fat absorption") + ylab("amount absorbed (g)")
print(p)

```



The p-value for the F -test is 0.001. The scientist would reject H_0 at any of the usual test levels (such as, 0.05 or 0.01). The data suggest that the population mean absorption rates differ across fats *in some way*. The F -test does not say *how* they differ. The pooled standard deviation $s_{\text{pooled}} = 8.18$ is the “Residual standard error”. We’ll ignore the rest of this output for now.

```

fit.f <- aov(amount ~ type, data = fat.long)
summary(fit.f)
##          Df Sum Sq Mean Sq F value Pr(>F)

```

```

## type      3   1595     532    7.95 0.0011 ***
## Residuals 20   1338      67
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
fit.f
## Call:
##   aov(formula = amount ~ type, data = fat.long)
##
## Terms:
##           type Residuals
## Sum of Squares 1596     1338
## Deg. of Freedom 3         20
##
## Residual standard error: 8.18
## Estimated effects may be unbalanced

```

■ CLICKER Qs — ANOVA, Fat 1/2 ■

■ CLICKER Qs — ANOVA, Fat 1/2 ■

5.2 Multiple Comparison Methods: Fisher's Method

The ANOVA F -test checks whether all the population means are equal. **Multiple comparisons** are often used as a follow-up to a significant ANOVA F -test to determine which population means are different. I will discuss Fisher's, Bonferroni's, and Tukey's methods for comparing all pairs of means.

Fisher's least significant difference method (**LSD or FSD**) is a two-step process:

1. Carry out the ANOVA F -test of $H_0 : \mu_1 = \mu_2 = \dots = \mu_k$ at the α level. If H_0 is not rejected, stop and conclude that there is insufficient evidence to claim differences among population means. If H_0 is rejected, go to step 2.

2. Compare each pair of means using a pooled two sample t -test at the α level. Use s_{pooled} from the ANOVA table and $df = dfE$ (Residual).

To see where the name LSD originated, consider the t -test of $H_0 : \mu_i = \mu_j$ (i.e., populations i and j have same mean). The t -statistic is

$$t_s = \frac{\bar{Y}_i - \bar{Y}_j}{s_{\text{pooled}} \sqrt{\frac{1}{n_i} + \frac{1}{n_j}}}.$$

You reject H_0 if $|t_s| \geq t_{\text{crit}}$, or equivalently, if

$$|\bar{Y}_i - \bar{Y}_j| \geq t_{\text{crit}} s_{\text{pooled}} \sqrt{\frac{1}{n_i} + \frac{1}{n_j}}.$$

The minimum absolute difference between \bar{Y}_i and \bar{Y}_j needed to reject H_0 is the LSD, the quantity on the right hand side of this inequality. If all the sample sizes are equal $n_1 = n_2 = \dots = n_k$ then the LSD is the same for each comparison:

$$LSD = t_{\text{crit}} s_{\text{pooled}} \sqrt{\frac{2}{n_1}},$$

where n_1 is the common sample size.

I will illustrate Fisher's method on the doughnut data, using $\alpha = 0.05$. At the first step, you reject the hypothesis that the population mean absorptions are equal because p-value= 0.001. At the second step, compare all pairs of fats at the 5% level. Here, $s_{\text{pooled}} = 8.18$ and $t_{\text{crit}} = 2.086$ for a two-sided test based on 20 df (the dfE for Residual SS). Each sample has six observations, so the LSD for each comparison is

$$LSD = 2.086 \times 8.18 \times \sqrt{\frac{2}{6}} = 9.85.$$

Any two sample means that differ by at least 9.85 in magnitude are **significantly different** at the 5% level.

An easy way to compare all pairs of fats is to order the samples by their sample means. The samples can then be grouped easily, noting that two fats are in the same group if the absolute difference between their sample means is smaller than the LSD.

Fats	Sample Mean
2	85.00
3	74.83
1	74.50
4	62.00

There are six comparisons of two fats. From this table, you can visually assess which sample means differ by at least the $LSD=9.85$, and which ones do not. For completeness, the table below summarizes each comparison:

Comparison	Absolute difference in means	Exceeds LSD?
Fats 2 and 3	10.17	Yes
2 and 1	10.50	Yes
2 and 4	23.00	Yes
Fats 3 and 1	0.33	No
3 and 4	12.83	Yes
Fats 1 and 4	12.50	Yes

The end product of the multiple comparisons is usually presented as a collection of **groups**, where a group is defined to be a set of populations with sample means that are not significantly different from each other. Overlap among groups is common, and occurs when one or more populations appears in two or more groups. Any overlap requires a more careful interpretation of the analysis.

There are three groups for the doughnut data, with no overlap. Fat 2 is in a group by itself, and so is Fat 4. Fats 3 and 1 are in a group together. This information can be summarized by ordering the samples from lowest to highest average, and then connecting the fats in the same group using an underscore:

FAT 4 FAT 1 FAT 3 FAT 2
----- ----- -----

The results of a multiple comparisons must be interpreted carefully. At the 5% level, you have sufficient evidence to conclude that the population mean absorption for Fat 2 and Fat 4 are each different than the other population means. However, there is insufficient evidence to conclude that the population mean absorptions for Fats 1 and 3 differ.

Be Careful with Interpreting Groups in Multiple Comparisons!

To see why you must be careful when interpreting groupings, suppose you obtain two groups in a three sample problem. One group has samples 1 and 3. The other group has samples 3 and 2:

1	3	2
-----		-----

This occurs, for example, when $|\bar{Y}_1 - \bar{Y}_2| \geq LSD$, but both $|\bar{Y}_1 - \bar{Y}_3|$ and $|\bar{Y}_3 - \bar{Y}_2|$ are less than the LSD. There is a tendency to conclude, and please try to avoid this line of attack, that populations 1 and 3 have the same mean, populations 2 and 3 have the same mean, but populations 1 and 2 have different means. This conclusion is illogical. The groupings imply that we have sufficient evidence to conclude that population means 1 and 2 are different, but insufficient evidence to conclude that population mean 3 differs from either of the other population means.

5.2.1 FSD Multiple Comparisons in R

One way to get Fisher comparisons in R uses `pairwise.t.test()` with `p.adjust.method`. The resulting summary of the multiple comparisons is in terms of p-values for all pairwise two-sample t-tests using the pooled standard deviation from the ANOVA using `pool.sd = TRUE`. This output can be used to generate groupings. A summary of the p-values is given below. Let us see that we can recover the groups from this output.

```
#### Multiple Comparisons
# all pairwise comparisons among levels of fat
# Fisher's LSD (FSD) uses "none"
pairwise.t.test(fat.long$amount, fat.long$type,
                 pool.sd = TRUE, p.adjust.method = "none")

##
##  Pairwise comparisons using t tests with pooled SD
##
## data: fat.long$amount and fat.long$type
##
##      fat1   fat2    fat3
## fat2 0.038  -
## fat3 0.944 0.044  -
## fat4 0.015 9.3e-05 0.013
##
## P value adjustment method: none
```

Discussion of the FSD Method: family error rate

There are $c = k(k - 1)/2$ pairs of means to compare in the second step of the FSD method. Each comparison is done at the α level, where for a generic comparison of the i^{th} and j^{th} populations

$$\alpha = \text{probability of rejecting } H_0 : \mu_i = \mu_j \text{ when } H_0 \text{ is true.}$$

The individual error rate is not the only error rate that is important in multiple comparisons. The **family error rate** (FER), or the **experimentwise error rate**, is defined to be *the probability of at least one false rejection of a true hypothesis $H_0 : \mu_i = \mu_j$ over all comparisons*. When many comparisons are made, you *may* have a large probability of making one or more false rejections of true null hypotheses. In particular, when all c comparisons of two population means are performed, each at the α level, then

$$\alpha < FER < c\alpha.$$

For example, in the doughnut problem where $k = 4$, there are $c = 4(3)/2 = 6$ possible comparisons of pairs of fats. If each comparison is carried out at the 5% level, then $0.05 < FER < 0.30$. At the second step of the FSD method, you could have up to a 30% chance of claiming one or more pairs of population means are different if no differences existed between population means. Most statistical packages do not evaluate the FER, so the upper bound is used.

The first step of the FSD method is the ANOVA “screening” test. The multiple comparisons are carried out only if the F -test suggests that not all population means are equal. This screening test tends to deflate the FER for the two-step FSD procedure. However, the FSD method is commonly criticized for being extremely liberal (too many false rejections of true null hypotheses) when some, but not many, differences exist — especially when the number of comparisons is large. This conclusion is fairly intuitive. When you do a large number of tests, each, say, at the 5% level, then sampling variation alone will suggest differences in 5% of the comparisons where the H_0 is true. The number of false rejections could be enormous with a large number of comparisons. For example, chance variation alone would account for an average of 50 significant differences in 1000 comparisons (about 45 populations) each at the 5% level.

5.2.2 Bonferroni Comparisons

The Bonferroni method controls the FER by reducing the individual comparison error rate. The FER is guaranteed to be no larger than a prespecified amount, say α , by setting the individual error rate for each of the c comparisons of interest to α/c . Therefore, $\alpha/c < FER < c\alpha/c = \alpha$, thus the upper bound for FER is α . Larger differences in the sample means are needed before declaring statistical significance using the Bonferroni adjustment than when using the FSD method at the α level.

■ CLICKER Qs — ANOVA, Bonferroni ■

Assuming all comparisons are of interest, you can implement the Bonferroni adjustment in R by specifying `p.adjust.method = "bonf"`. A by-product of the Bonferroni adjustment is that we have at least $100(1 - \alpha)\%$ confidence that all pairwise t -test statements hold simultaneously!

```
# Bonferroni 95% Individual p-values
# All Pairwise Comparisons among Levels of fat
pairwise.t.test(fat.long$amount, fat.long$type,
                 pool.sd = TRUE, p.adjust.method = "bonf")

##
##  Pairwise comparisons using t tests with pooled SD
##
##  data: fat.long$amount and fat.long$type
##
##      fat1     fat2     fat3
## fat2 0.22733 -       -
## fat3 1.00000 0.26241 -
## fat4 0.09286 0.00056 0.07960
##
## P value adjustment method: bonferroni
```

Looking at the output, can you create the groups? You should get the groups given below, which implies you have sufficient evidence to conclude that the population mean absorption for Fat 2 is different than that for Fat 4.

FAT 4	FAT 1	FAT 3	FAT 2
-----	-----		

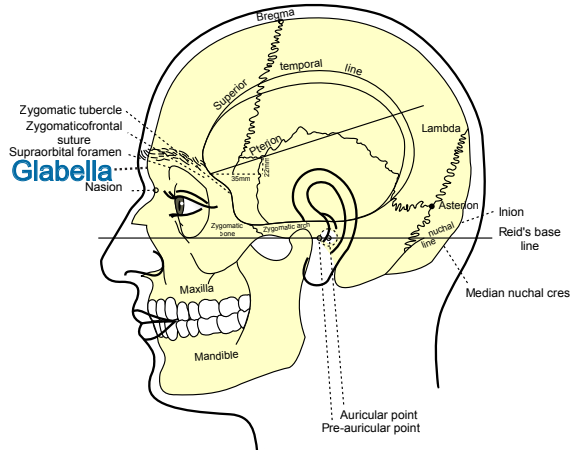
The Bonferroni method tends to produce “coarser” groups than the FSD method, because the individual comparisons are conducted at a lower (alpha/error) level. Equivalently, the minimum significant difference is inflated for the Bonferroni method. For example, in the doughnut problem with $FER \leq 0.05$, the critical

value for the individual comparisons at the 0.0083 level is $t_{\text{crit}} = 2.929$ with $df = 20$. The minimum significant difference for the Bonferroni comparisons is

$$LSD = 2.929 \times 8.18 \times \sqrt{\frac{2}{6}} = 13.824$$

versus an LSD=9.85 for the FSD method. Referring back to our table of sample means on page 158, we see that the sole comparison where the absolute difference between sample means exceeds 13.824 involves Fats 2 and 4.

Example from Koopmans: glabella facial tissue thickness In an anthropological study of facial tissue thickness for different racial groups, data were taken during autopsy at several points on the faces of deceased individuals. The Glabella measurements taken at the bony ridge for samples of individuals from three racial groups (cauc = Caucasian, afam = African American, and naaa = Native American and Asian) follow. The data values are in mm.



```
#### Example from Koopmans: glabella facial tissue thickness
```

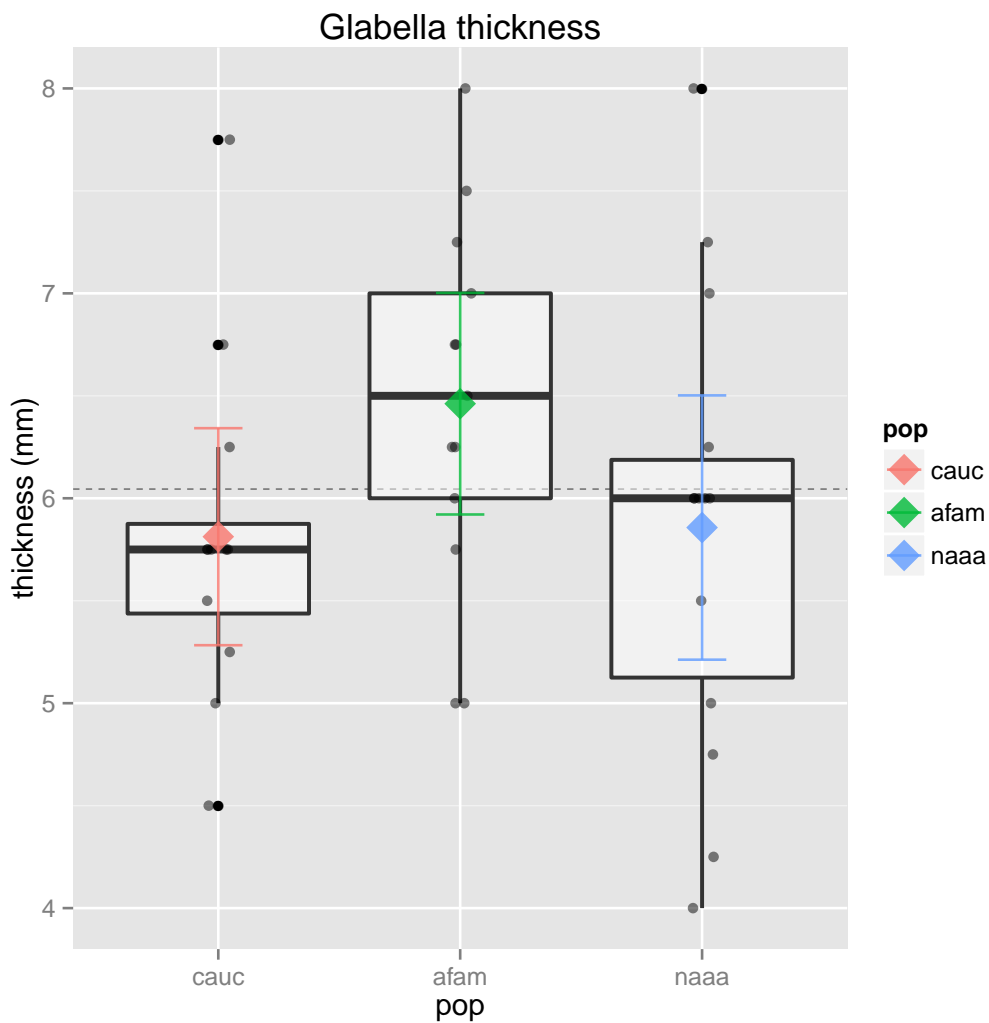
```
glabella <- read.table(text="
```

Row	cauc	afam	naaa
1	5.75	6.00	8.00
2	5.50	6.25	7.00
3	6.75	6.75	6.00
4	5.75	7.00	6.25
5	5.00	7.25	5.50
6	5.75	6.75	4.00
7	5.75	8.00	5.00
8	7.75	6.50	6.00
9	5.75	7.50	7.25
10	5.25	6.25	6.00
11	4.50	5.00	6.00
12	6.25	5.75	4.25

```
13     NA  5.00  4.75
14     NA     NA  6.00
", header=TRUE)

glabella.long <- melt(glabella,
                      id.vars=c("Row"),
                      variable.name = "pop",
                      value.name = "thickness",
                      # remove NAs
                      na.rm = TRUE
)
# naming variables manually, the variable.name and value.name not working 11/2012
names(glabella.long) <- c("Row", "pop", "thickness")
# another way to remove NAs:
#glabella.long <- subset(glabella.long, !is.na(thickness))

# Plot the data using ggplot
library(ggplot2)
p <- ggplot(glabella.long, aes(x = pop, y = thickness))
# plot a reference line for the global mean (assuming no groups)
p <- p + geom_hline(yintercept = mean(glabella.long$thickness),
                     colour = "black", linetype = "dashed", size = 0.3, alpha = 0.5)
# boxplot, size=.75 to stand out behind CI
p <- p + geom_boxplot(size = 0.75, alpha = 0.5)
# points for observed data
p <- p + geom_point(position = position_jitter(w = 0.05, h = 0), alpha = 0.5)
# diamond at mean for each group
p <- p + stat_summary(fun.y = mean, geom = "point", shape = 18, size = 6,
                      aes(colour=pop), alpha = 0.8)
# confidence limits based on normal distribution
p <- p + stat_summary(fun.data = "mean_cl_normal", geom = "errorbar",
                      width = .2, aes(colour=pop), alpha = 0.8)
p <- p + labs(title = "Glabella thickness") + ylab("thickness (mm)")
print(p)
```



There are 3 groups, so there are 3 possible pairwise comparisons. If you want a Bonferroni analysis with FER of no greater than 0.05, you should do the individual comparisons at the $0.05/3 = 0.0167$ level. Except for the mild outlier in the Caucasian sample, the observed distributions are fairly symmetric, with similar spreads. I would expect the standard ANOVA to perform well here.

Let μ_c = population mean Glabella measurement for Caucasians, μ_a = population mean Glabella measurement for African Americans, and μ_n = population mean Glabella measurement for Native Americans and Asians.

```
glabella.summary <- ddply(glabella.long, "pop",
  function(X) { data.frame( m = mean(X$thickness),
    s = sd(X$thickness),
    n = length(X$thickness) ) } )
glabella.summary
##   pop      m      s   n
## 1 cauc 5.812 0.8334 12
## 2 afam 6.462 0.8947 13
## 3 naaa 5.857 1.1168 14
```

```

fit.g <- aov(thickness ~ pop, data = glabella.long)
summary(fit.g)

##           Df Sum Sq Mean Sq F value Pr(>F)
## pop          2    3.4   1.699   1.83   0.18
## Residuals   36   33.5   0.929

fit.g
## Call:
##   aov(formula = thickness ~ pop, data = glabella.long)
##
## Terms:
##           pop Residuals
## Sum of Squares 3.40     33.46
## Deg. of Freedom 2         36
##
## Residual standard error: 0.9641
## Estimated effects may be unbalanced

```

At the 5% level, you would not reject the hypothesis that the population mean Glabella measurements are identical. That is, you do not have sufficient evidence to conclude that these racial groups differ with respect to their average Glabella measurement. **This is the end of the analysis!**

The Bonferroni intervals reinforce this conclusion, all the p-values are greater than 0.05. If you were to calculate CIs for the difference in population means, each would contain zero. You can think of the Bonferroni intervals as simultaneous CI. We're (at least) 95% confident that all of the following statements hold simultaneously: $-1.62 \leq \mu_c - \mu_a \leq 0.32$, $-0.91 \leq \mu_n - \mu_c \leq 1.00$, and $-1.54 \leq \mu_n - \mu_a \leq 0.33$. The individual CIs have level $100(1 - 0.0167)\% = 98.33\%$.

```

# Bonferroni 95% Individual p-values
# All Pairwise Comparisons among Levels of glabella
pairwise.t.test(glabella.long$thickness, glabella.long$pop,
                 pool.sd = TRUE, p.adjust.method = "bonf")

##
## Pairwise comparisons using t tests with pooled SD
##
## data: glabella.long$thickness and glabella.long$pop
##
##      cauc afam
## afam 0.30 -
## naaa 1.00 0.34
##
## P value adjustment method: bonferroni

```

5.3 Further Discussion of Multiple Comparisons

The FSD and Bonferroni methods comprise the ends of the spectrum of multiple comparisons methods. Among multiple comparisons procedures, the FSD method is most likely to find differences, whether real or due to sampling variation, whereas Bonferroni is often the most conservative method. You can be reasonably sure that differences suggested by the Bonferroni method will be suggested by almost all other methods, whereas differences not significant under FSD will not be picked up using other approaches.

The Bonferroni method is conservative, but tends to work well when the number of comparisons is small, say 4 or less. A smart way to use the Bonferroni adjustment is to focus attention only on the comparisons of interest (generated independently of looking at the data!), and ignore the rest. I will return to this point later.

A commonly-used alternative is **Tukey's** honest significant difference method (HSD). John Tukey's honest significant difference method is to reject the equality of a pair of means based, not on the t -distribution, but the studentized range distribution. To implement Tukey's method with a FER of α , reject $H_0 : \mu_i = \mu_j$ when

$$|\bar{Y}_i - \bar{Y}_j| \geq \frac{q_{crit}}{\sqrt{2}} s_{\text{pooled}} \sqrt{\frac{1}{n_i} + \frac{1}{n_j}},$$

where q_{crit} is the α level critical value of the studentized range distribution. For the doughnut fats, the groupings based on Tukey and Bonferroni comparisons are identical.

```
##### Tukey's honest significant difference method (HSD)
## Fat
# Tukey 95% Individual p-values
# All Pairwise Comparisons among Levels of fat
TukeyHSD(fit.f)

## Tukey multiple comparisons of means
## 95% family-wise confidence level
##
## Fit: aov(formula = amount ~ type, data = fat.long)
##
```

```

## $type
##          diff      lwr      upr   p adj
## fat2-fat1 10.5000 -2.719 23.7190 0.1511
## fat3-fat1  0.3333 -12.886 13.5524 0.9999
## fat4-fat1 -12.5000 -25.719  0.7190 0.0679
## fat3-fat2 -10.1667 -23.386  3.0524 0.1710
## fat4-fat2 -23.0000 -36.219 -9.7810 0.0005
## fat4-fat3 -12.8333 -26.052  0.3857 0.0590

## Glabella
# Tukey 95% Individual p-values
# All Pairwise Comparisons among Levels of pop
TukeyHSD(fit.g)

## Tukey multiple comparisons of means
## 95% family-wise confidence level
##
## Fit: aov(formula = thickness ~ pop, data = glabella.long)
##
## $pop
##          diff      lwr      upr   p adj
## afam-cauc  0.64904 -0.2943 1.5924 0.2260
## naaa-cauc  0.04464 -0.8824 0.9717 0.9924
## naaa-afam -0.60440 -1.5120 0.3033 0.2473

```

Another popular method controls the **false discovery rate** (FDR) instead of the FER. The FDR is the expected proportion of false discoveries amongst the rejected hypotheses. The false discovery rate is a less stringent condition than the family-wise error rate, so these methods are more powerful than the others, though with a higher FER. I encourage you to learn more about the methods by Benjamini, Hochberg, and Yekutieli.

```

##### false discovery rate (FDR)
## Fat
# FDR
pairwise.t.test(fat.long$amount, fat.long$type,
                 pool.sd = TRUE, p.adjust.method = "BH")

##
## Pairwise comparisons using t tests with pooled SD
##
## data: fat.long$amount and fat.long$type
##
##      fat1     fat2     fat3
## fat2 0.05248 -       -
## fat3 0.94443 0.05248 -
## fat4 0.03095 0.00056 0.03095
##
## P value adjustment method: BH

```

```
## Glabella
# FDR
pairwise.t.test(glabella.long
```

■ CLICKER Qs — ANOVA, multiple comparisons ■

5.4 Checking Assumptions in ANOVA Problems

The classical ANOVA assumes that the populations have normal frequency curves and the populations have equal variances (or spreads). You can test the normality assumption using multiple normal QQ-plots and normal scores tests, which we discussed in Chapter 4. An alternative approach that is useful with three or more samples is to make a single normal scores plot for the entire data set. The samples must be *centered* at the same location for this to be meaningful. (WHY?) This is done by subtracting the sample mean from each observation in the sample, giving the so-called **residuals**. A normal scores plot or histogram of the residuals should resemble a sample from a normal population. These two plots can be generated with output in `$residuals` from the `anova()` procedure.

For the glabella residuals, there are a few observations outside the confidence bands, but the formal normality tests each have p-values > 0.2 , so there's weak but unconvincing evidence of nonnormality.

```
#### Checking Assumptions in ANOVA Problems
# plot of data
par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
```

```

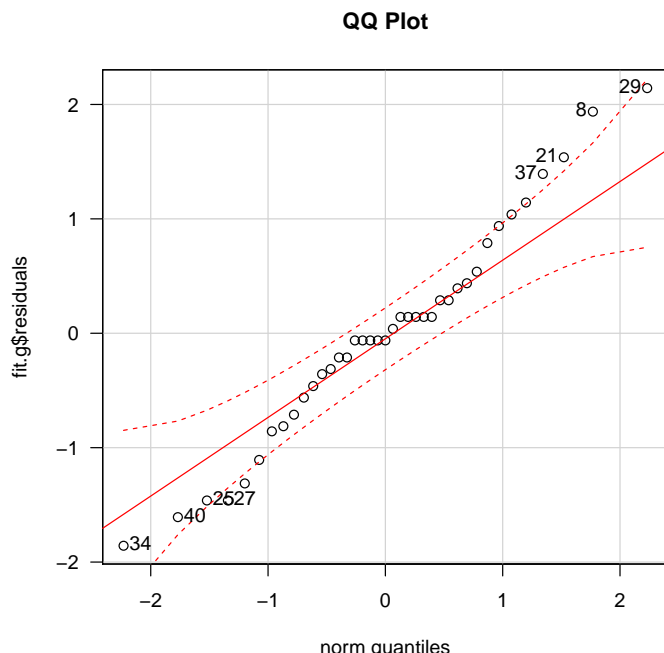
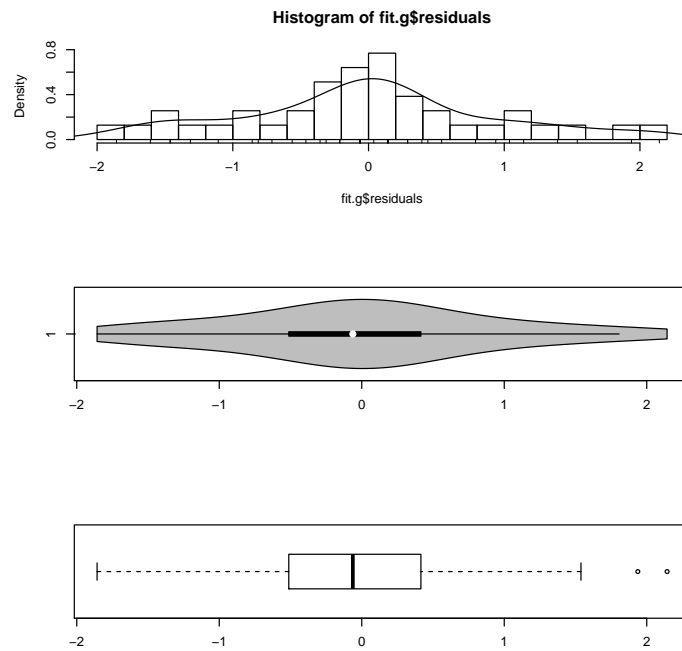
hist(fit.g$residuals, freq = FALSE, breaks = 20)
points(density(fit.g$residuals), type = "l")
rug(fit.g$residuals)

# violin plot
library(vioplot)
vioplot(fit.g$residuals, horizontal=TRUE, col="gray")

# boxplot
boxplot(fit.g$residuals, horizontal=TRUE)

# QQ plot
par(mfrow=c(1,1))
library(car)
qqPlot(fit.g$residuals, las = 1, id.n = 8, id.cex = 1, lwd = 1, main="QQ Plot")
## 29 8 34 40 21 25 27 37
## 39 38 1 2 37 3 4 36

```



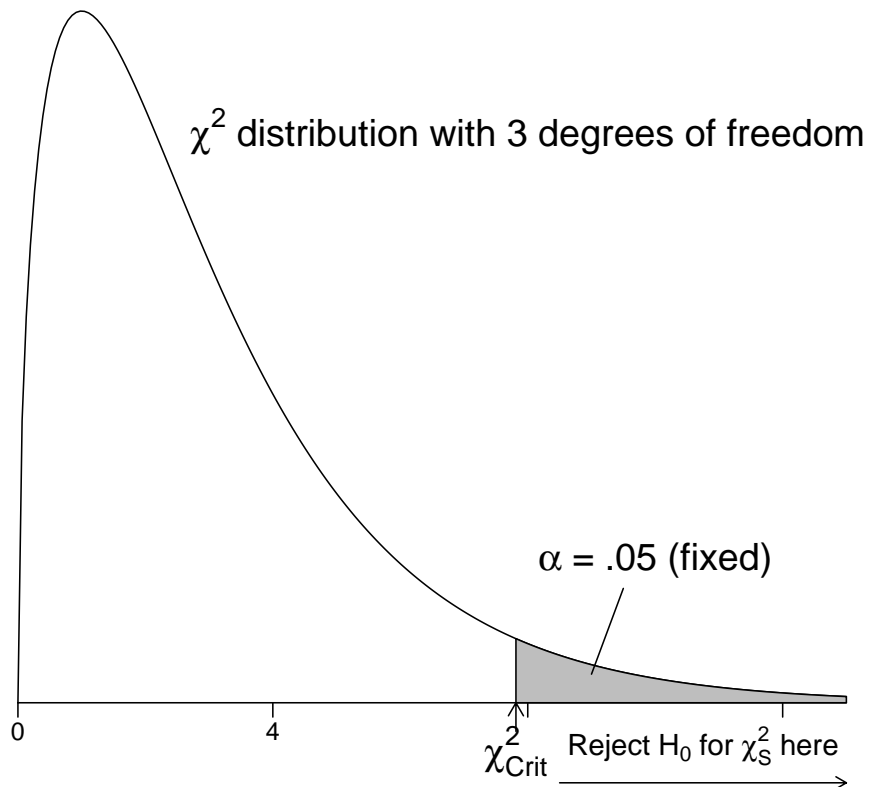
```

shapiro.test(fit.g$residuals)
##
##  Shapiro-Wilk normality test
##
## data: fit.g$residuals
## W = 0.9769, p-value = 0.5927
library(nortest)
ad.test(fit.g$residuals)
##
## Anderson-Darling normality test
##
## data: fit.g$residuals
## A = 0.3773, p-value = 0.3926

```

```
# lillie.test(fit.g$residuals)
cvm.test(fit.g$residuals)
##
##  Cramer-von Mises normality test
##
## data: fit.g$residuals
## W = 0.0709, p-value = 0.2648
```

In Chapter 4, I illustrated the use of Bartlett's test and Levene's test for equal population variances, and showed how to evaluate these tests in R.



R does the calculation for us, as illustrated below. Because the p-value > 0.5 , we fail to reject the null hypothesis that the population variances are equal. This result is not surprising given how close the sample variances are to each other.

```
## Test equal variance
# Barlett assumes populations are normal
bartlett.test(thickness ~ pop, data = glabella.long)
##
## Bartlett test of homogeneity of variances
##
```

```
## data: thickness by pop
## Bartlett's K-squared = 1.131, df = 2, p-value = 0.568
```

Levene's and Fligner's tests are consistent with Bartlett's.

```
# Levene does not assume normality, requires car package
library(car)
leveneTest(thickness ~ pop, data = glabella.long)

## Levene's Test for Homogeneity of Variance (center = median)
##          Df F value Pr(>F)
## group    2   0.53   0.59
##          36

# Fligner is a nonparametric test
fligner.test(thickness ~ pop, data = glabella.long)

##
## Fligner-Killeen test of homogeneity of variances
##
## data: thickness by pop
## Fligner-Killeen:med chi-squared = 1.031, df = 2, p-value =
## 0.5972
```

5.5 Example from the Child Health and Development Study (CHDS)

We consider data from the birth records of 680 live-born white male infants. The infants were born to mothers who reported for pre-natal care to three clinics of the Kaiser hospitals in northern California. As an initial analysis, we will examine whether maternal smoking has an effect on the birth weights of these children. To answer this question, we define 3 groups based on mother's smoking history: (1) mother does not currently smoke or never smoked, (2) mother smoked less than one pack of cigarettes a day during pregnancy, and (3) mother smoked at least one pack of cigarettes a day during pregnancy.

Let μ_i = pop mean birth weight (lb) for children in group i , ($i = 1, 2, 3$). We wish to test $H_0 : \mu_1 = \mu_2 = \mu_3$ against $H_A : \text{not } H_0$.

We read in the data, create a `smoke` factor variable, and plot the data by smoking group.

```
#### Example from the Child Health and Development Study (CHDS)
# description at http://statacumon.com/teach/ADA1/ADA1_notes_05-CHDS_desc.txt
# read data from website
# chds <- read.csv("http://statacumon.com/teach/ADA1/ADA1_notes_05-CHDS.csv")
# (for some reason this wasn't reading from website, so I'm reading it from my harddrive)
```

```

chds <- read.csv("ADA1_notes_05-CHDS.csv")

chds$smoke <- rep(NA, nrow(chds));
# no cigs
chds[(chds$m_smok == 0), "smoke"] <- "0 cigs";
# less than 1 pack (20 cigs = 1 pack)
chds[(chds$m_smok > 0) & (chds$m_smok < 20), "smoke"] <- "1-19 cigs";
# at least 1 pack (20 cigs = 1 pack)
chds[(chds$m_smok >= 20), "smoke"] <- "20+ cigs";
chds$smoke <- factor(chds$smoke)

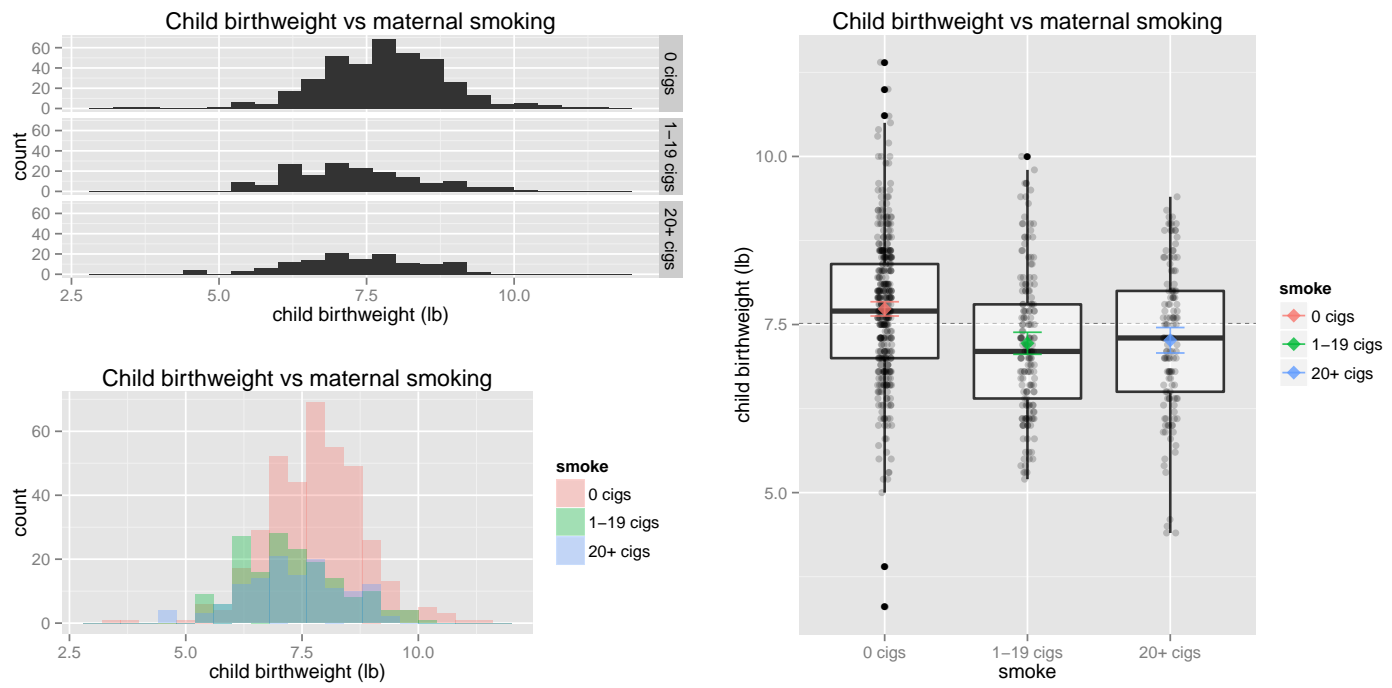
# histogram using ggplot
p1 <- ggplot(chds, aes(x = c_bwt))
p1 <- p1 + geom_histogram(binwidth = .4)
p1 <- p1 + facet_grid(smoke ~ .)
p1 <- p1 + labs(title = "Child birthweight vs maternal smoking") +
  xlab("child birthweight (lb)")
#print(p1)

p2 <- ggplot(chds, aes(x = c_bwt, fill=smoke))
p2 <- p2 + geom_histogram(binwidth = .4, alpha = 1/3, position="identity")
p2 <- p2 + labs(title = "Child birthweight vs maternal smoking") +
  xlab("child birthweight (lb)")
#print(p2)

library(gridExtra)
grid.arrange(p1, p2, ncol=1)

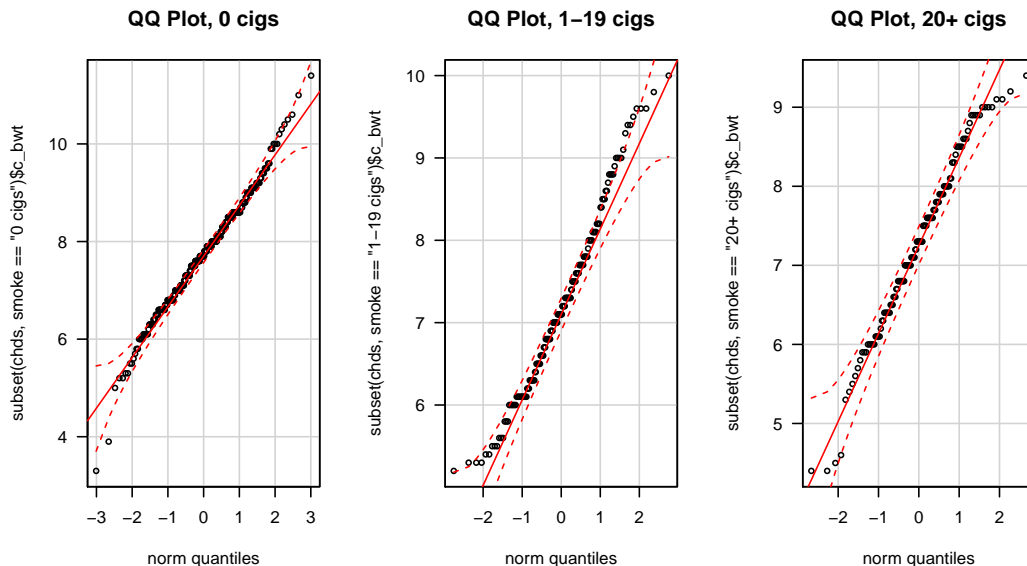
# Plot the data using ggplot
library(ggplot2)
p <- ggplot(chds, aes(x = smoke, y = c_bwt))
# plot a reference line for the global mean (assuming no groups)
p <- p + geom_hline(yintercept = mean(chds$c_bwt),
  colour = "black", linetype = "dashed", size = 0.3, alpha = 0.5)
# boxplot, size=.75 to stand out behind CI
p <- p + geom_boxplot(size = 0.75, alpha = 0.5)
# points for observed data
p <- p + geom_point(position = position_jitter(w = 0.05, h = 0), alpha = 0.2)
# diamond at mean for each group
p <- p + stat_summary(fun.y = mean, geom = "point", shape = 18, size = 4,
  aes(colour=smoke), alpha = 0.8)
# confidence limits based on normal distribution
p <- p + stat_summary(fun.data = "mean_cl_normal", geom = "errorbar",
  width = .2, aes(colour=smoke), alpha = 0.8)
p <- p + labs(title = "Child birthweight vs maternal smoking") +
  ylab("child birthweight (lb)")
print(p)

```



Looking at the boxplots, there is some evidence of non-normality here. Although there are outliers in the no smoking group, we need to recognize that the sample size for this group is fairly large (381). Given that boxplots are calibrated in such a way that 7 outliers per 1000 observations are expected when sampling from a normal population, 5 outliers (only 4 are visible!) out of 381 seems a bit excessive. A formal test rejects the hypothesis of normality in the no and low smoker groups. The normal probability plot and the histogram of the residuals also suggests that the population distributions are heavy tailed.

```
library(car)
par(mfrow=c(1,3))
qqPlot(subset(chds, smoke == "0 cigs")$c_bwt, las = 1, id.n = 0,
       id.cex = 1, lwd = 1, main="QQ Plot, 0 cigs")
qqPlot(subset(chds, smoke == "1-19 cigs")$c_bwt, las = 1, id.n = 0,
       id.cex = 1, lwd = 1, main="QQ Plot, 1-19 cigs")
qqPlot(subset(chds, smoke == "20+ cigs")$c_bwt, las = 1, id.n = 0,
       id.cex = 1, lwd = 1, main="QQ Plot, 20+ cigs")
```



```

library(nortest)
# 0 cigs -----
shapiro.test(subset(chds, smoke == "0 cigs" )$c_bwt)

##
## Shapiro-Wilk normality test
##
## data: subset(chds, smoke == "0 cigs")$c_bwt
## W = 0.9872, p-value = 0.00199
ad.test(   subset(chds, smoke == "0 cigs" )$c_bwt)

##
## Anderson-Darling normality test
##
## data: subset(chds, smoke == "0 cigs")$c_bwt
## A = 0.9283, p-value = 0.01831
cvm.test(   subset(chds, smoke == "0 cigs" )$c_bwt)

##
## Cramer-von Mises normality test
##
## data: subset(chds, smoke == "0 cigs")$c_bwt
## W = 0.1384, p-value = 0.03374
# 1-19 cigs -----
shapiro.test(subset(chds, smoke == "1-19 cigs")$c_bwt)

##
## Shapiro-Wilk normality test
##
## data: subset(chds, smoke == "1-19 cigs")$c_bwt
## W = 0.9785, p-value = 0.009926
ad.test(   subset(chds, smoke == "1-19 cigs")$c_bwt)

##
## Anderson-Darling normality test
##
## data: subset(chds, smoke == "1-19 cigs")$c_bwt
## A = 0.8308, p-value = 0.03149

```

```
cvm.test( subset(chds, smoke == "1-19 cigs")$c_bwt)
##
## Cramer-von Mises normality test
##
## data: subset(chds, smoke == "1-19 cigs")$c_bwt
## W = 0.1133, p-value = 0.07317

# 20+ cigs -----
shapiro.test(subset(chds, smoke == "20+ cigs" )$c_bwt)

##
## Shapiro-Wilk normality test
##
## data: subset(chds, smoke == "20+ cigs")$c_bwt
## W = 0.9813, p-value = 0.06962

ad.test( subset(chds, smoke == "20+ cigs" )$c_bwt)

##
## Anderson-Darling normality test
##
## data: subset(chds, smoke == "20+ cigs")$c_bwt
## A = 0.4001, p-value = 0.3578

cvm.test( subset(chds, smoke == "20+ cigs" )$c_bwt)

##
## Cramer-von Mises normality test
##
## data: subset(chds, smoke == "20+ cigs")$c_bwt
## W = 0.0405, p-value = 0.6694
```

Fit the ANOVA (we'll look at the table below).

```
fit.c <- aov(c_bwt ~ smoke, data = chds)
```

A formal test of normality on the residuals of the combined sample is marginally significant (SW p-value= 0.047, others > 0.10). However, I am not overly concerned about this for the following reasons: in large samples, small deviations from normality are often statistically significant and in my experience, the small deviations we are seeing here are not likely to impact our conclusions, in the sense that non-parametric methods that do not require normality will lead to the same conclusions.

```
# plot of data
par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(fit.c$residuals, freq = FALSE, breaks = 20)
points(density(fit.c$residuals), type = "l")
rug(fit.c$residuals)

# violin plot
library(vioplot)
```

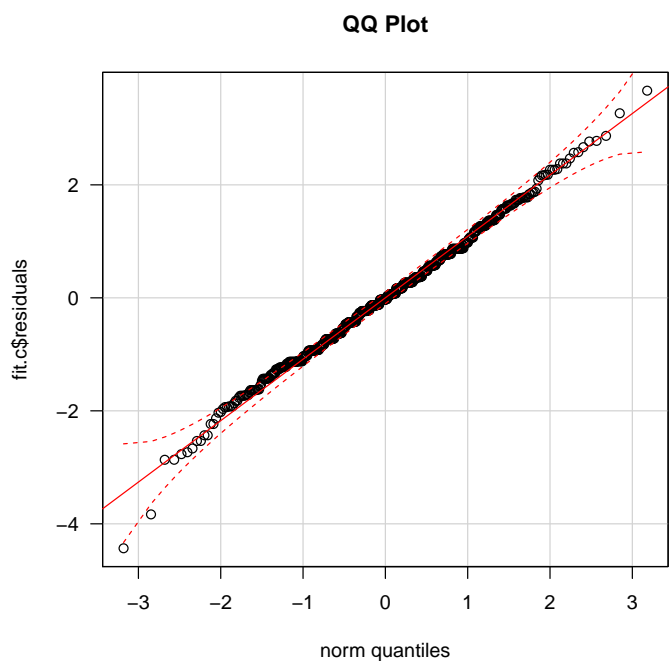
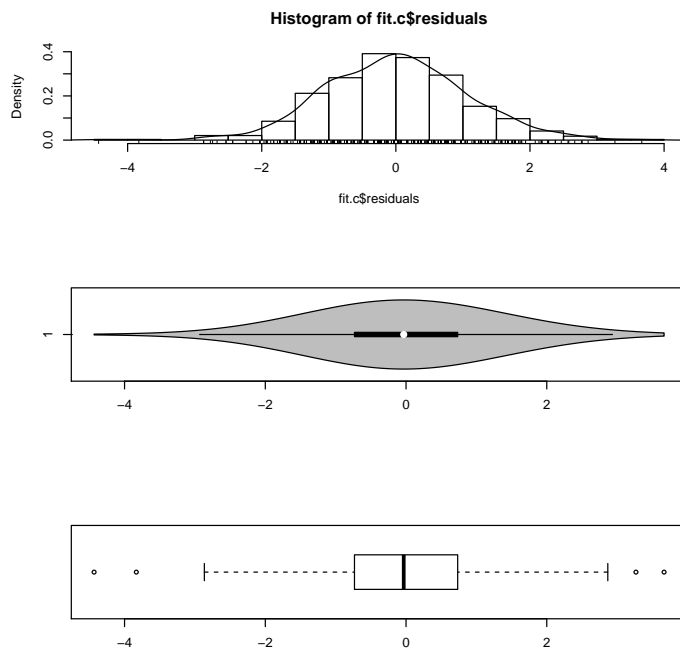
```

vioplot(fit.c$residuals, horizontal=TRUE, col="gray")

# boxplot
boxplot(fit.c$residuals, horizontal=TRUE)

# QQ plot
par(mfrow=c(1,1))
library(car)
qqPlot(fit.c$residuals, las = 1, id.n = 0, id.cex = 1, lwd = 1, main="QQ Plot")

```



```

shapiro.test(fit.c$residuals)

##
##  Shapiro-Wilk normality test
##
## data: fit.c$residuals
## W = 0.9955, p-value = 0.04758

library(nortest)
ad.test(fit.c$residuals)

##
##  Anderson-Darling normality test
##
## data: fit.c$residuals
## A = 0.6218, p-value = 0.1051

cvm.test(fit.c$residuals)

##
##  Cramer-von Mises normality test
##
## data: fit.c$residuals
## W = 0.092, p-value = 0.1449

```

Looking at the summaries, we see that the sample standard deviations are close. Formal tests of equal population variances are far from significant. The p-values for Bartlett's test and Levene's test are greater than 0.4. Thus, the standard ANOVA appears to be appropriate here.

```
# calculate summaries
chds.summary <- ddply(chds, "smoke",
  function(X) { data.frame( m = mean(X$c_bwt),
    s = sd(X$c_bwt),
    n = length(X$c_bwt) ) } )
chds.summary
##          smoke      m      s      n
## 1      0 cigs 7.733 1.052 381
## 2 1-19 cigs 7.221 1.078 169
## 3 20+ cigs 7.266 1.091 130

## Test equal variance
# assumes populations are normal
bartlett.test(c_bwt ~ smoke, data = chds)

##
##  Bartlett test of homogeneity of variances
##
## data: c_bwt by smoke
## Bartlett's K-squared = 0.3055, df = 2, p-value = 0.8583

# does not assume normality, requires car package
library(car)
leveneTest(c_bwt ~ smoke, data = chds)
## Levene's Test for Homogeneity of Variance (center = median)
##          Df F value Pr(>F)
## group     2     0.76   0.47
##          677

# nonparametric test
fligner.test(c_bwt ~ smoke, data = chds)

##
## Fligner-Killeen test of homogeneity of variances
##
## data: c_bwt by smoke
## Fligner-Killeen:med chi-squared = 2.093, df = 2, p-value =
## 0.3512
```

The p-value for the F -test is less than 0.0001. We would reject H_0 at any of the usual test levels (such as 0.05 or 0.01). The data suggest that the population mean birth weights differ across smoking status groups.

```
summary(fit.c)
##           Df Sum Sq Mean Sq F value  Pr(>F)
## smoke       2     41   20.35    17.9 2.6e-08 ***
## Residuals  677   769    1.14
```

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
fit.c
## Call:
##   aov(formula = c_bwt ~ smoke, data = chds)
##
## Terms:
##           smoke Residuals
## Sum of Squares    40.7     769.5
## Deg. of Freedom      2       677
##
## Residual standard error: 1.066
## Estimated effects may be unbalanced
```

The Tukey multiple comparisons suggest that the mean birth weights are different (higher) for children born to mothers that did not smoke during pregnancy.

```
## CHDS
# Tukey 95% Individual p-values
TukeyHSD(fit.c)

## Tukey multiple comparisons of means
## 95% family-wise confidence level
##
## Fit: aov(formula = c_bwt ~ smoke, data = chds)
##
## $smoke
##          diff      lwr      upr   p adj
## 1-19 cigs-0 cigs -0.51151 -0.7429 -0.2801 0.0000
## 20+ cigs-0 cigs -0.46665 -0.7210 -0.2123 0.0001
## 20+ cigs-1-19 cigs  0.04485 -0.2473  0.3370 0.9308
```

Chapter 6

Nonparametric Methods

Learning objectives

After completing this topic, you should be able to:

select the appropriate procedure based on assumptions.

explain reason for using one procedure over another.

decide whether the medians between multiple populations are different.

Achieving these goals contributes to mastery in these course learning outcomes:

3. select correct statistical procedure.
5. define parameters of interest and hypotheses in words and notation.
6. summarize data visually, numerically, and descriptively.
8. use statistical software.
10. identify and explain statistical methods, assumptions, and limitations.
12. make evidence-based decisions.

6.1 Introduction

Nonparametric methods do not require the normality assumption of classical techniques. When the normality assumption is met, the ANOVA and *t*-test are most powerful, in that if the alternative is true these methods will make the correct decision with highest probability. However, if the normality assumption is not met, results from the ANOVA and *t*-test can be misleading and too liberal. I will describe and illustrate selected **non-parametric methods**, and compare them with classical methods. Some motivation and discussion of

the strengths and weaknesses of non-parametric methods is given.

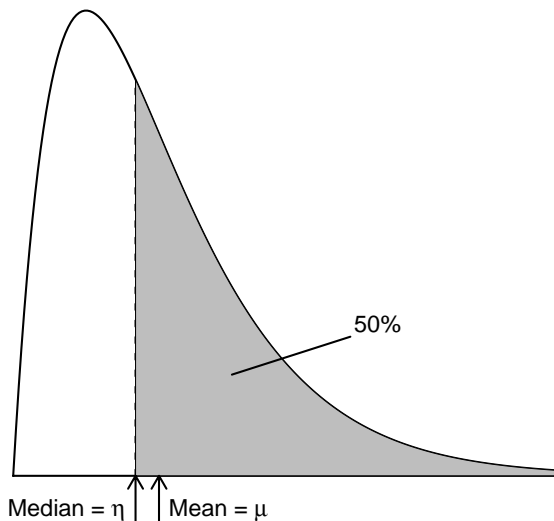
6.2 The Sign Test and CI for a Population Median

The **sign test** assumes that you have a random sample from a population, but makes **no assumption** about the population shape. The standard t -test provides inferences on a population mean. The sign test, in contrast, provides inferences about a **population median**.

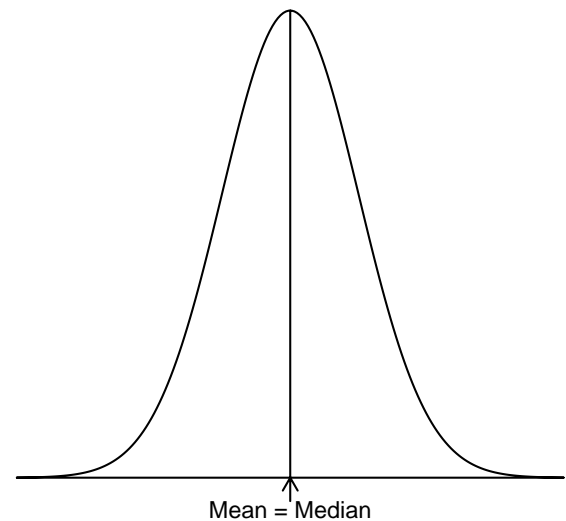
If the population frequency curve is symmetric (see below), then the population median, identified by η , and the population mean μ are identical. In this case the sign procedures provide inferences for the population mean, though less powerfully than the t -test.

The idea behind the sign test is straightforward. Suppose you have a sample of size m from the population, and you wish to test $H_0 : \eta = \eta_0$ (a given value). Let S be the number of sampled observations *above* η_0 . If H_0 is true, you expect S to be approximately one-half the sample size, $0.5m$. If S is much greater than $0.5m$, the data suggests that $\eta > \eta_0$. If S is much less than $0.5m$, the data suggests that $\eta < \eta_0$.

Mean and Median differ with skewed distributions



Mean and Median are the same with symmetric distributions



S has a **Binomial distribution** when H_0 is true. The Binomial distri-

bution is used to construct a test with size α (approximately). For a two-sided alternative $H_A : \eta \neq \eta_0$, the test rejects H_0 when S is significantly different from $0.5m$, as determined from the reference Binomial distribution. One-sided tests use the corresponding lower or upper tail of the distribution. To generate a CI for η , you can exploit the duality between CI and tests. A $100(1 - \alpha)\%$ CI for η consists of all values η_0 not rejected by a two-sided size α test of $H_0 : \eta = \eta_0$.

Not all test sizes and confidence levels are possible because the test statistic S is discrete valued. R's `SIGN.test()` in the `BSDA` package gives an exact p-value for the test, and approximates the desired confidence level using a linear interpolation algorithm.

Example: Income Data Recall that the income distribution is extremely skewed, with two extreme outliers at 46 and 1110.

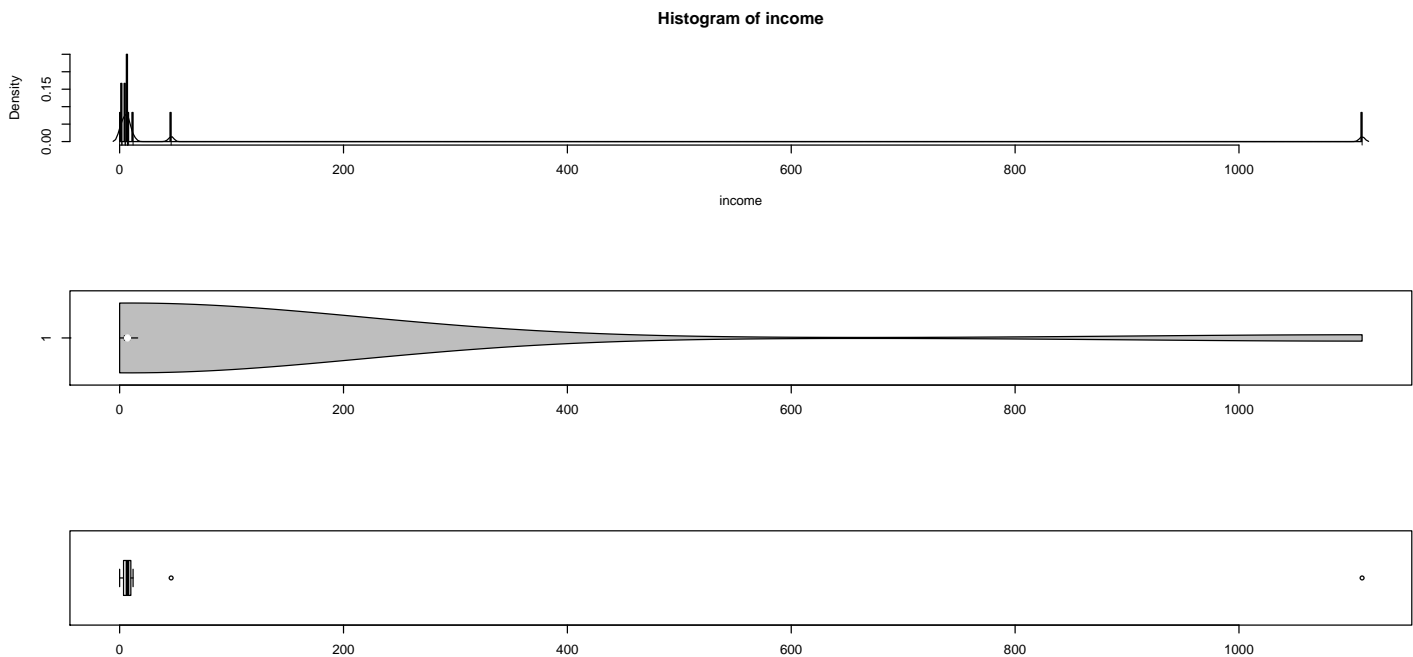
```
#### Example: Income Data
income <- c(7, 1110, 7, 5, 8, 12, 0, 5, 2, 2, 46, 7)
# sort in decreasing order
income <- sort(income, decreasing = TRUE)
income
## [1] 1110   46   12    8    7    7    7    5    5    2    2    0
summary(income)
##   Min. 1st Qu. Median     Mean 3rd Qu.    Max.
##   0.0    4.2    7.0   101.0    9.0  1110.0
sd(income)
## [1] 318
```

The income data is unimodal, skewed right, with two extreme outliers.

```
par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(income, freq = FALSE, breaks = 1000)
points(density(income), type = "l")
rug(income)

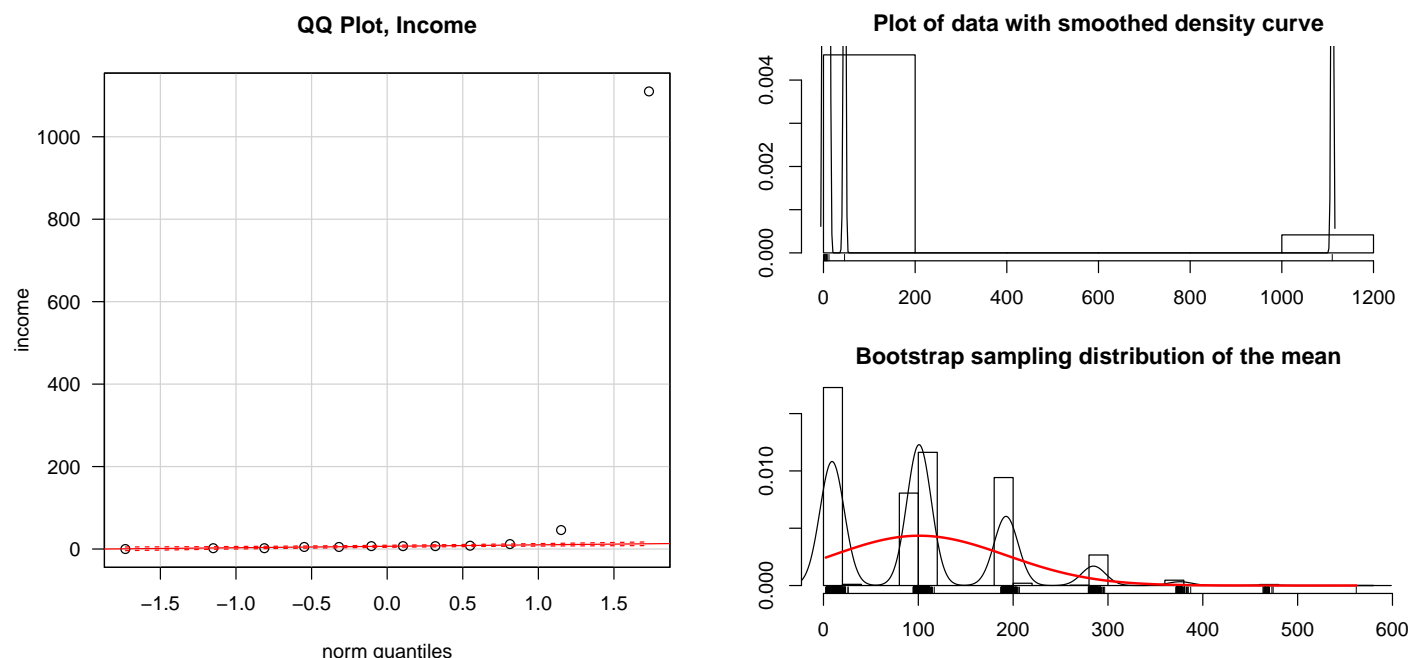
# violin plot
library(vioplot)
vioplot(income, horizontal=TRUE, col="gray")

# boxplot
boxplot(income, horizontal=TRUE)
```



The normal Q-Q-plot of the sample data indicates strong deviation from normality, and the CLT can't save us: even the bootstrap sampling distribution of the mean indicates strong deviation from normality.

```
library(car)
qqPlot(income, las = 1, id.n = 0, id.cex = 1, lwd = 1, main="QQ Plot, Income")
bs.one.samp.dist(income)
```



The presence of the outliers has a dramatic effect on the 95% CI for the population mean income μ , which goes from -101 to 303 (in 1000 dollar units). This t -CI is suspect because the normality assumption is unreasonable. A CI for the population median income η is more sensible because the median is likely to be a more reasonable measure of typical value. Using the sign procedure,

you are 95% confident that the population median income is between 2.32 and 11.57 (times \$1000).

```
library(BSDA)
t.test(income)

##
## One Sample t-test
##
## data: income
## t = 1.099, df = 11, p-value = 0.2951
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
## -101.1 303.0
## sample estimates:
## mean of x
##      100.9

SIGN.test(income)

##
## One-sample Sign-Test
##
## data: income
## s = 11, p-value = 0.0009766
## alternative hypothesis: true median is not equal to 0
## 95 percent confidence interval:
##    2.319 11.575
## sample estimates:
## median of x
##      7
##          Conf.Level L.E.pt U.E.pt
## Lower Achieved CI      0.8540  5.000   8.00
## Interpolated CI        0.9500  2.319  11.57
## Upper Achieved CI      0.9614  2.000  12.00
```

Example: Age at First Heart Transplant Recall that the distribution of ages is skewed to the left with a lower outlier. A question of interest is whether the “typical age” at first transplant is 50. This can be formulated as a test about the population median η or as a test about the population mean μ , depending on the interpretation.

```
##### Example: Age at First Heart Transplant
age <- c(54, 42, 51, 54, 49, 56, 33, 58, 54, 64, 49)
# sort in decreasing order
age <- sort(age, decreasing = TRUE)
age
## [1] 64 58 56 54 54 54 51 49 49 42 33
summary(age)
##    Min.  1st Qu.  Median      Mean  3rd Qu.  Max.
##    33.0    49.0    54.0    51.3    55.0    64.0
sd(age)
```

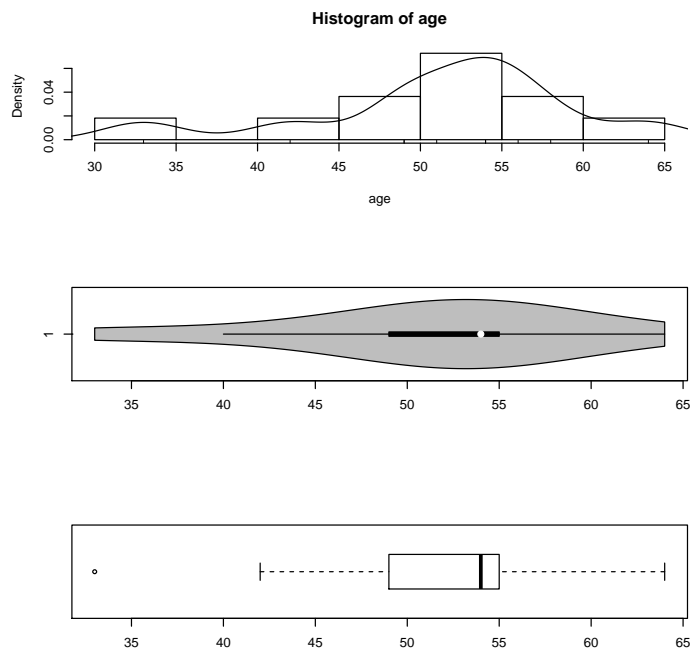
```
## [1] 8.259
```

The age data is unimodal, skewed left, no extreme outliers.

```
par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(age, freq = FALSE, breaks = 10)
points(density(age), type = "l")
rug(age)

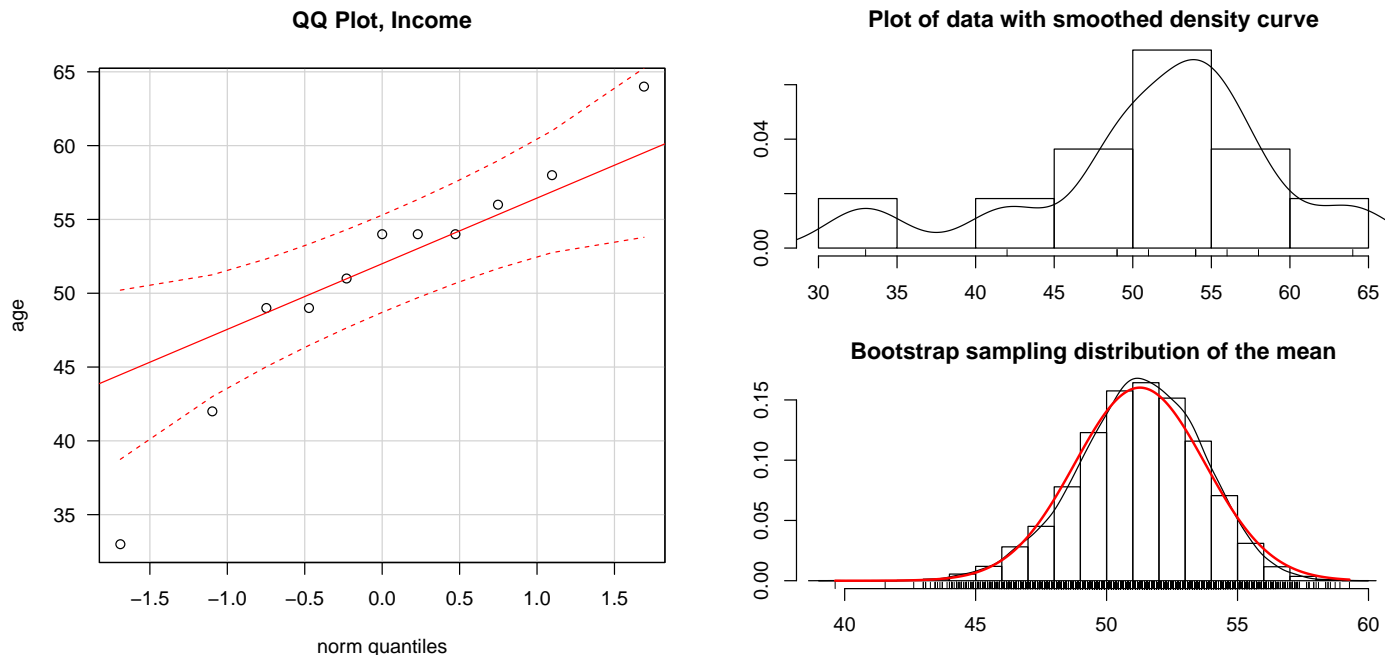
# violin plot
library(vioplot)
vioplot(age, horizontal=TRUE, col="gray")

# boxplot
boxplot(age, horizontal=TRUE)
```



The normal QQ-plot of the sample data indicates mild deviation from normality in the left tail (2 points of 11 outside the bands), and the bootstrap sampling distribution of the mean indicates weak deviation from normality. It is good practice in this case to use the nonparametric test as a double-check of the *t*-test, with the nonparametric test being the more conservative test.

```
library(car)
qqPlot(age, las = 1, id.n = 0, id.cex = 1, lwd = 1, main="QQ Plot, Income")
bs.one.samp.dist(age)
```



The sign test for $H_0 : \eta = 50$ against $H_A : \eta \neq 50$ has a p-value of 0.549, which is not sufficient to reject H_0 . A 95% CI for η is 47.0 to 56.6 years, which includes the hypothesized median age of 50. Similar conclusions are reached with the t -CI and the test on μ , but you should have less confidence in these results because the normality assumption is tenuous.

```
library(BSDA)
t.test(age, mu=50)

##
##  One Sample t-test
##
## data: age
## t = 0.5111, df = 10, p-value = 0.6204
## alternative hypothesis: true mean is not equal to 50
## 95 percent confidence interval:
## 45.72 56.82
## sample estimates:
## mean of x
##      51.27

SIGN.test(age, md=50)

##
##  One-sample Sign-Test
##
## data: age
## s = 7, p-value = 0.5488
## alternative hypothesis: true median is not equal to 50
## 95 percent confidence interval:
## 46.99 56.57
## sample estimates:
## median of x
##      54
## Conf.Level L.E.pt U.E.pt
```

```
## Lower Achieved CI      0.9346 49.00 56.00
## Interpolated CI        0.9500 46.99 56.57
## Upper Achieved CI       0.9883 42.00 58.00
```

6.3 Wilcoxon Signed-Rank Procedures

The **Wilcoxon** procedure assumes you have a random sample from a population with a **symmetric** frequency curve. The curve need not be normal. The test and CI can be viewed as procedures for either the population median or mean.

To illustrate the computation of the Wilcoxon statistic W , suppose you wish to test $H_0 : \mu = \mu_0 = 10$ on the made-up data below. The test statistic requires us to compute the **signs** of $X_i - \mu_0$ and the **ranks** of $|X_i - \mu_0|$. Ties in $|X_i - \mu_0|$ get the average rank and observations at μ_0 (here 10) are always discarded. The Wilcoxon statistic is the **sum of the signed ranks for observations above $\mu_0 = 10$** . For us

$$W = 6 + 4.5 + 8 + 2 + 4.5 + 7 = 32.$$

X_i	$X_i - 10$	sign	$ X_i - 10 $	rank	sign \times rank
20	10	+	10	6	6
18	8	+	8	4.5	4.5
23	13	+	13	8	8
5	-5	-	5	3	-3
14	4	+	4	2	2
8	-2	-	2	1	-1
18	8	+	8	4.5	4.5
22	12	+	12	7	7

The sum of all ranks is always $0.5m(m+1)$, where m is the sample size. If H_0 is true, you expect W to be approximately $0.5 \times 0.5m(m+1) = 0.25m(m+1)$. Why? Recall that W adds up the ranks for observations above μ_0 . If H_0 is true, you expect $1/2$ of all observations to be above μ_0 , assuming the population distribution is symmetric. The ranks of observations above μ_0 should add to approximately $1/2$ times the sum of all ranks. You reject H_0 in favor of

$H_A : \mu \neq \mu_0$ if W is much larger than, or much smaller than $0.25m(m + 1)$. One sided tests can also be constructed. The Wilcoxon CI for μ is computed in a manner analogous to that described for the sign CI.

Here, $m = 8$ so the sum of all ranks is $0.5 \times 8 \times 9 = 36$ (check yourself). The expected value of W is $0.5 \times 0.5 \times 8 \times 9 = 18$. Is the observed value of $W = 32$ **far from** the expected value of 18? To formally answer this question, we need to use the Wilcoxon procedures, which are implemented in R with `wilcox.test()`.

Example: Made-up Data The boxplot indicates that the distribution is fairly symmetric, so the Wilcoxon method is reasonable (so is a t -CI and test).

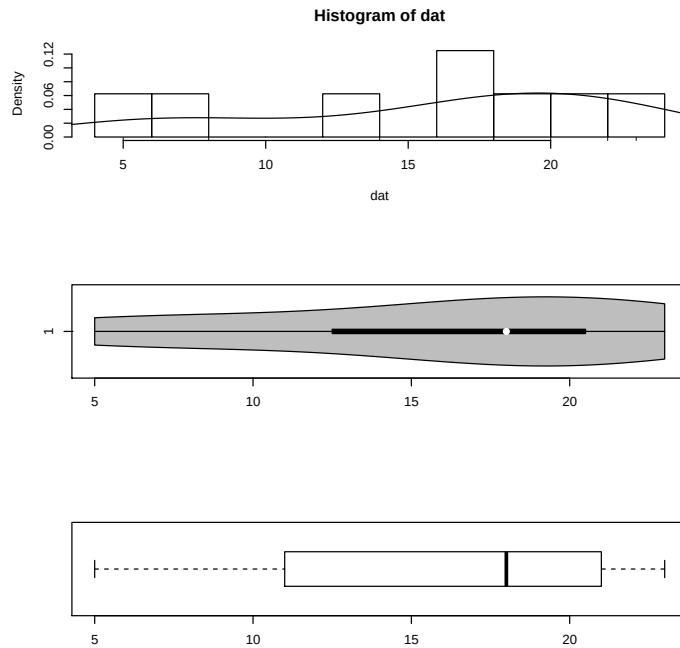
```
#### Example: Made-up Data
dat <- c(20, 18, 23, 5, 14, 8, 18, 22)
# sort in decreasing order
dat <- sort(dat, decreasing = TRUE)
dat
## [1] 23 22 20 18 18 14 8 5
summary(dat)
##    Min. 1st Qu. Median     Mean 3rd Qu.    Max.
##    5.0    12.5   18.0    16.0   20.5    23.0
sd(dat)
## [1] 6.525
```

The `dat` data is unimodal, skewed left, no extreme outliers.

```
par(mfrow=c(3,1))
# Histogram overlaid with kernel density curve
hist(dat, freq = FALSE, breaks = 10)
points(density(dat), type = "l")
rug(dat)

# violin plot
library(vioplot)
vioplot(dat, horizontal=TRUE, col="gray")

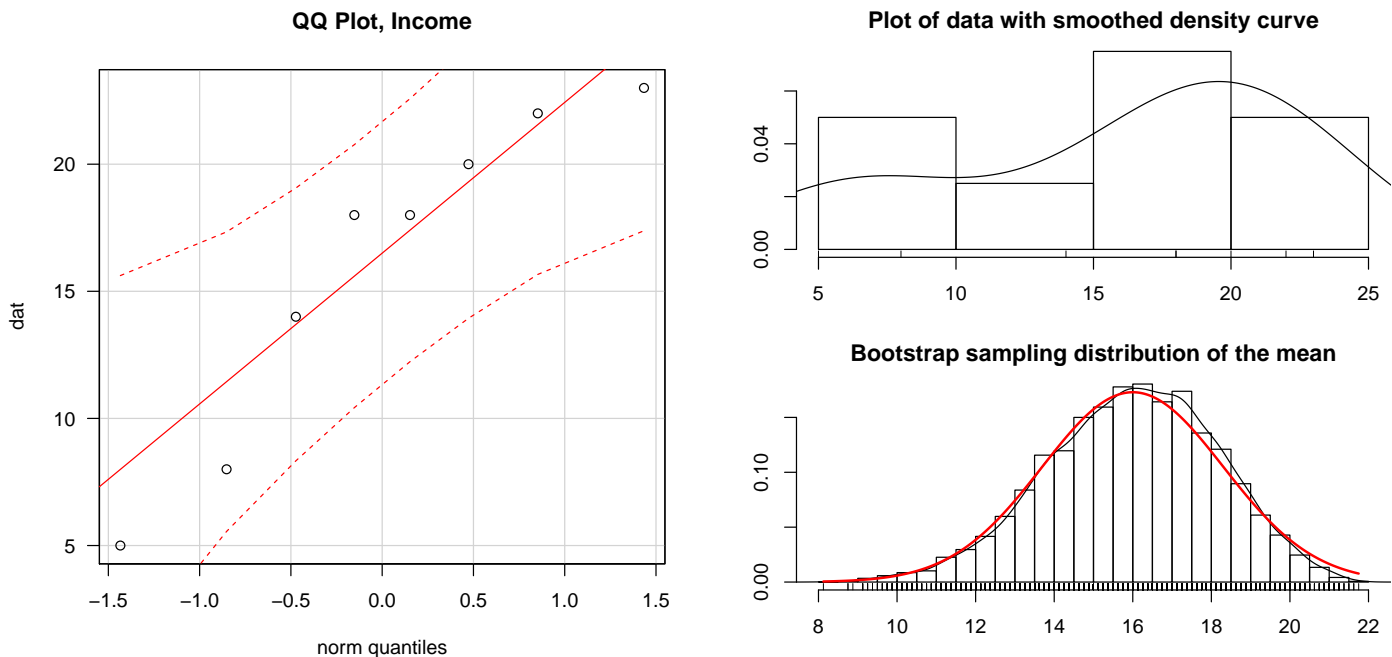
# boxplot
boxplot(dat, horizontal=TRUE)
```



The normal QQ-plot of the sample data indicates insufficient evidence of deviation from normality though both the QQ-plot and the bootstrap sampling distribution of the mean indicates weak left-skewness. Either the Wilcoxon or t -test are appropriate.

```
par(mfrow=c(1,1))
library(car)
qqPlot(dat, las = 1, id.n = 0, id.cex = 1, lwd = 1, main="QQ Plot, Income")

bs.one.samp.dist(dat)
```



The Wilcoxon p-value with continuity correction for testing $H_0 : \mu = 10$ against a two-sided alternative is 0.058. This would not lead to rejecting H_0 at

the 5% level.

```
t.test(dat, mu=10)
##
##  One Sample t-test
##
## data:  dat
## t = 2.601, df = 7, p-value = 0.03537
## alternative hypothesis: true mean is not equal to 10
## 95 percent confidence interval:
## 10.55 21.45
## sample estimates:
## mean of x
##      16

# with continuity correction in the normal approximation for the p-value
wilcox.test(dat, mu=10, conf.int=TRUE)

## Warning: cannot compute exact p-value with ties
## Warning: cannot compute exact confidence interval with ties
##
## Wilcoxon signed rank test with continuity correction
##
## data:  dat
## V = 32, p-value = 0.0584
## alternative hypothesis: true location is not equal to 10
## 95 percent confidence interval:
## 9.5 21.5
## sample estimates:
## (pseudo)median
##      16.01

# without continuity correction
wilcox.test(dat, mu=10, conf.int=TRUE, correct=FALSE)

## Warning: cannot compute exact p-value with ties
## Warning: cannot compute exact confidence interval with ties
##
## Wilcoxon signed rank test
##
## data:  dat
## V = 32, p-value = 0.04967
## alternative hypothesis: true location is not equal to 10
## 95 percent confidence interval:
## 11 21
## sample estimates:
## (pseudo)median
##      16.01
```

6.3.1 Nonparametric Analyses of Paired Data

Nonparametric methods for single samples can be used to analyze paired data because the difference between responses within pairs is the unit of analysis.

Example: Sleep Remedies I will illustrate Wilcoxon methods on the paired comparison of two remedies A and B for insomnia. The number of hours of sleep gained on each method was recorded.

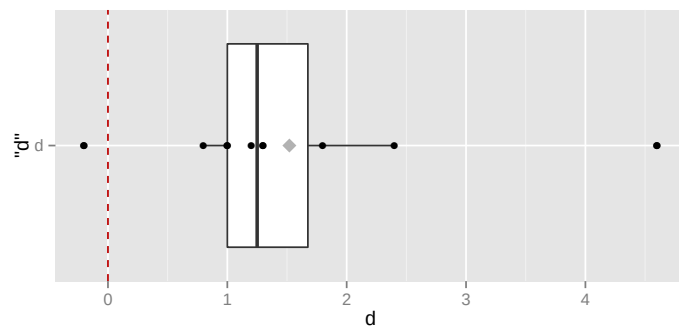
```
#### Example: Sleep Remedies
# Data and numerical summaries
a <- c( 0.7, -1.6, -0.2, -1.2,  0.1,  3.4,  3.7,  0.8,  0.0,  2.0)
b <- c( 1.9,  0.8,  1.1,  0.1, -0.1,  4.4,  5.5,  1.6,  4.6,  3.0)
d <- b - a;
sleep <- data.frame(a, b, d)
summary(sleep$d)

##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.
##    -0.20    1.00   1.25    1.52    1.67    4.60

shapiro.test(sleep$d)

##
##  Shapiro-Wilk normality test
##
## data: sleep$d
## W = 0.838, p-value = 0.04173

# boxplot
library(ggplot2)
p3 <- ggplot(sleep, aes(x = "d", y = d))
p3 <- p3 + geom_hline(yintercept=0, colour="#BB0000", linetype="dashed")
p3 <- p3 + geom_boxplot()
p3 <- p3 + geom_point()
p3 <- p3 + stat_summary(fun.y = mean, geom = "point", shape = 18,
                        size = 4, alpha = 0.3)
p3 <- p3 + coord_flip()
print(p3)
```



The boxplot shows that distribution of differences is reasonably symmetric but not normal. Recall that the Shapiro-Wilk test of normality was significant

at the 5% level ($p\text{-value}=0.042$). It is sensible to use the Wilcoxon procedure on the differences. Let μ_B be the population mean sleep gain on remedy B, and μ_A be the population mean sleep gain on remedy A. You are 95% confident that $\mu_B - \mu_A$ is between 0.8 and 2.8 hours. Putting this another way, you are 95% confident that μ_B exceeds μ_A by between 0.8 and 2.8 hours. The p -value for testing $H_0 : \mu_B - \mu_A = 0$ against a two-sided alternative is 0.008, which strongly suggests that $\mu_B \neq \mu_A$. This agrees with the CI. Note that the t -CI and test give qualitatively similar conclusions as the Wilcoxon methods, but the t -test p -value is about half as large.

If you are uncomfortable with the symmetry assumption, you could use the sign CI for the population median difference between B and A. I will note that a 95% CI for the median difference goes from 0.86 to 2.2 hours.

```
t.test(sleep$d, mu=0)
##
##  One Sample t-test
##
## data: sleep$d
## t = 3.78, df = 9, p-value = 0.004352
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  0.6102 2.4298
## sample estimates:
## mean of x
##      1.52

# with continuity correction in the normal approximation for the p-value
wilcox.test(sleep$d, mu=0, conf.int=TRUE)

## Warning: cannot compute exact p-value with ties
## Warning: cannot compute exact confidence interval with ties
##
## Wilcoxon signed rank test with continuity correction
##
## data: sleep$d
## V = 54, p-value = 0.008004
## alternative hypothesis: true location is not equal to 0
## 95 percent confidence interval:
##  0.7999 2.8000
## sample estimates:
## (pseudo)median
##      1.3

# can use the paired= option
#wilcox.test(sleep$b, sleep$a, paired=TRUE, mu=0, conf.int=TRUE)
# if don't assume symmetry, can use sign test
#SIGN.test(sleep$d)
```

6.3.2 Comments on One-Sample Nonparametric Methods

For this discussion, I will assume that the underlying population distribution is (approximately) symmetric, which implies that population means and medians are equal (approximately). For symmetric distributions the t , sign, and Wilcoxon procedures are all appropriate.

If the underlying population distribution is extremely skewed, you can use the sign procedure to get a CI for the population median. Alternatively, as illustrated on HW 2, you can transform the data to a scale where the underlying distribution is nearly normal, and then use the classical t -methods. Moderate degrees of skewness will not likely have a big impact on the standard t -test and CI.

The one-sample t -test and CI are optimal when the underlying population frequency curve is normal. Essentially this means that the t -CI is, on average, narrowest among all CI procedures with given level, or that the t -test has the highest power among all tests with a given size. The width of a CI provides a measure of the sensitivity of the estimation method. For a given level CI, the narrower CI better pinpoints the unknown population mean.

With heavy-tailed symmetric distributions, the t -test and CI tend to be conservative. Thus, for example, a nominal 95% t -CI has actual coverage rates higher than 95%, and the nominal 5% t -test has an actual size smaller than 5%. The t -test and CI possess a property that is commonly called **robustness of validity**. However, data from heavy-tailed distributions can have a profound effect on the **sensitivity** of the t -test and CI. Outliers can dramatically inflate the standard error of the mean, causing the CI to be needlessly wide, and tests to have diminished power (outliers typically inflate p-values for the t -test). The sign and Wilcoxon procedures downweight the influence of outliers by looking at sign or signed-ranks instead of the actual data values. These two nonparametric methods are somewhat less efficient than the t -methods when the population is normal (efficiency is about 0.64 and 0.96 for the sign and Wilcoxon methods relative to the normal t -methods, where efficiency is the ratio of sample sizes needed for equal power), but can be infinitely more efficient with heavier than normal tailed distributions. In essence, the t -methods do not

have a **robustness of sensitivity**.

Nonparametric methods have gained widespread acceptance in many scientific disciplines, but not all. Scientists in some disciplines continue to use classical *t*-methods because they believe that the methods are robust to non-normality. As noted above, this is a robustness of validity, not sensitivity. This misconception is unfortunate, and results in the routine use of methods that are less powerful than the non-parametric techniques. **Scientists need to be flexible and adapt their tools to the problem at hand, rather than use the same tool indiscriminately!** I have run into suspicion that use of nonparametric methods was an attempt to “cheat” in some way — properly applied, they are excellent tools that *should* be used.

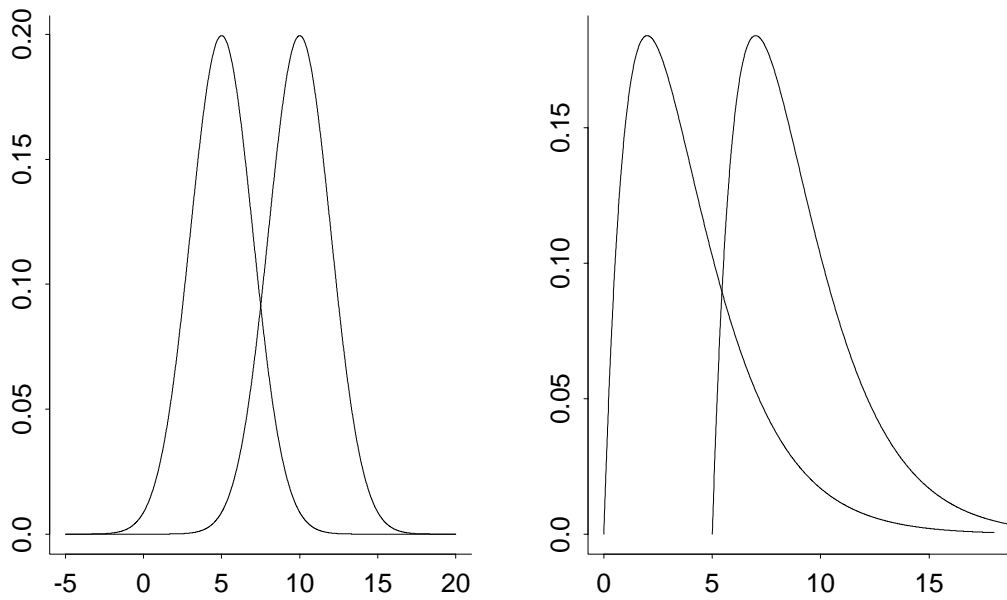
A minor weakness of nonparametric methods is that they do not easily generalize to complex modelling problems. A great deal of progress has been made in this area, but most software packages have not included the more advanced techniques (R is among the forerunners).

Nonparametric statistics used to refer almost exclusively to the set of methods such as we have been discussing that provided analogs like tests and CIs to the normal theory methods without requiring the assumption of sampling from normal distributions. There is now a large area of statistics also called nonparametric methods not focused on these goals at all. In our department we (used to) have a course titled “Nonparametric Curve Estimation & Image Reconstruction”, where the focus is much more general than relaxing an assumption of normality. In that sense, what we are covering in this course could be considered “classical” nonparametrics.

6.4 (Wilcoxon-)Mann-Whitney Two-Sample Procedure

The WMW procedure assumes you have independent random samples from the two populations, and assumes that the populations have the **same shapes and spreads** (the frequency curves for the two populations are “shifted” versions of each other — see below). The frequency curves are not required to be symmetric. The WMW procedures give a CI and tests on the difference $\eta_1 - \eta_2$

between the two population medians. If the populations are symmetric, then the methods apply to $\mu_1 - \mu_2$.



The R help on `?wilcox.test` gives references to how the exact WMW procedure is actually calculated; here is a good approximation to the exact method that is easier to understand. The WMW procedure is based on ranks. The two samples are combined, ranked from smallest to largest (1=smallest) and separated back into the original samples. If the two populations have equal medians, you expect the average rank in the two samples to be roughly equal. The WMW test computes a classical two sample t -test using the pooled variance on the ranks to assess whether the sample mean ranks are significantly different.

Example: Comparison of Cooling Rates of Uwet and Walker Co. Meteorites The Uwet¹ (Cross River, Nigeria) and Walker² County (Alabama, US) meteorite cooling rate data are below. A primary interest is comparing the population “typical” cooling rate measurements.

¹<http://www.lpi.usra.edu/meteor/metbull.php?code=24138>

²<http://www.lpi.usra.edu/meteor/metbull.php?code=24204>

```
#### Example: Comparison of Cooling Rates of Uwet and Walker Co. Meteorites
```

```
Uwet <- c(0.21, 0.25, 0.16, 0.23, 0.47, 1.20, 0.29, 1.10, 0.16)
```

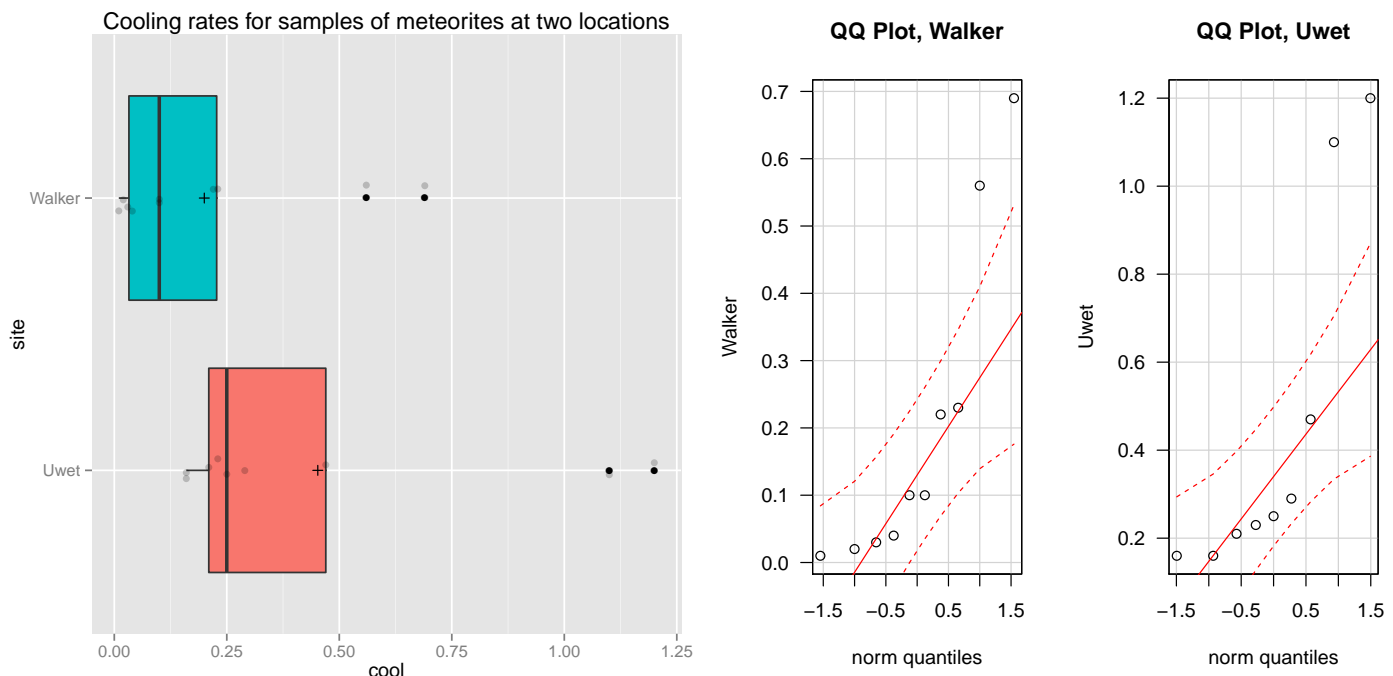
```
Walker <- c(0.69, 0.23, 0.10, 0.03, 0.56, 0.10, 0.01, 0.02, 0.04, 0.22)
```

The boxplots and normal QQ-plots show that the distributions are rather skewed to the right. The AD test of normality indicate that a normality assumption is unreasonable for each population.

```
met <- data.frame(Uwet=c(Uwet,NA), Walker)
library(reshape2)
met.long <- melt(met, variable.name = "site", value.name = "cool", na.rm=TRUE)

## Using as id variables
# naming variables manually, the variable.name and value.name not working 11/2012
names(met.long) <- c("site", "cool")
library(ggplot2)
p <- ggplot(met.long, aes(x = site, y = cool, fill=site))
p <- p + geom_boxplot()
p <- p + geom_point(position = position_jitter(w = 0.05, h = 0), alpha = 0.2)
p <- p + stat_summary(fun.y = mean, geom = "point", shape = 3, size = 2)
p <- p + coord_flip()
p <- p + labs(title = "Cooling rates for samples of meteorites at two locations")
p <- p + theme(legend.position="none")
print(p)

par(mfrow=c(1,2))
library(car)
qqPlot(Walker, las = 1, id.n = 0, id.cex = 1, lwd = 1, main="QQ Plot, Walker")
qqPlot(Uwet, las = 1, id.n = 0, id.cex = 1, lwd = 1, main="QQ Plot, Uwet")
```



I carried out the standard two-sample procedures to see what happens. The pooled-variance and Satterthwaite results are comparable, which is expected because the sample standard deviations and sample sizes are roughly equal.

Both tests indicate that the mean cooling rates for Uwet and Walker Co. meteorites are not significantly different at the 10% level. You are 95% confident that the mean cooling rate for Uwet is at most 0.1 less, and no more than 0.6 greater than that for Walker Co. (in degrees per million years).

```
# numerical summaries
summary(Uwet)

##      Min. 1st Qu. Median     Mean 3rd Qu.    Max.
## 0.160   0.210   0.250   0.452   0.470   1.200

c(sd(Uwet), IQR(Uwet), length(Uwet))
## [1] 0.407 0.260 9.000

summary(Walker)

##      Min. 1st Qu. Median     Mean 3rd Qu.    Max.
## 0.0100  0.0325  0.1000  0.2000  0.2280  0.6900

c(sd(Walker), IQR(Walker), length(Walker))
## [1] 0.239 0.195 10.000

t.test(Uwet, Walker, var.equal = TRUE)

##
## Two Sample t-test
##
## data: Uwet and Walker
## t = 1.669, df = 17, p-value = 0.1134
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.06663 0.57107
## sample estimates:
## mean of x mean of y
## 0.4522    0.2000

t.test(Uwet, Walker)

##
## Welch Two Sample t-test
##
## data: Uwet and Walker
## t = 1.624, df = 12.65, p-value = 0.129
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.08421 0.58865
## sample estimates:
## mean of x mean of y
## 0.4522    0.2000
```

Given the marked skewness, a nonparametric procedure is more appropriate. The Wilcoxon-Mann-Whitney comparison of population medians is reasonable. Why? The WMW test of equal population medians is significant (barely) at the 5% level. You are 95% confident that median cooling rate for Uwet exceeds

that for Walker by between 0+ and 0.45 degrees per million years.

```
wilcox.test(Uwet, Walker, conf.int = TRUE)
## Warning: cannot compute exact p-value with ties
## Warning: cannot compute exact confidence intervals with ties
##
## Wilcoxon rank sum test with continuity correction
##
## data: Uwet and Walker
## W = 69.5, p-value = 0.04974
## alternative hypothesis: true location shift is not equal to 0
## 95 percent confidence interval:
## 4.497e-05 4.500e-01
## sample estimates:
## difference in location
## 0.1703
```

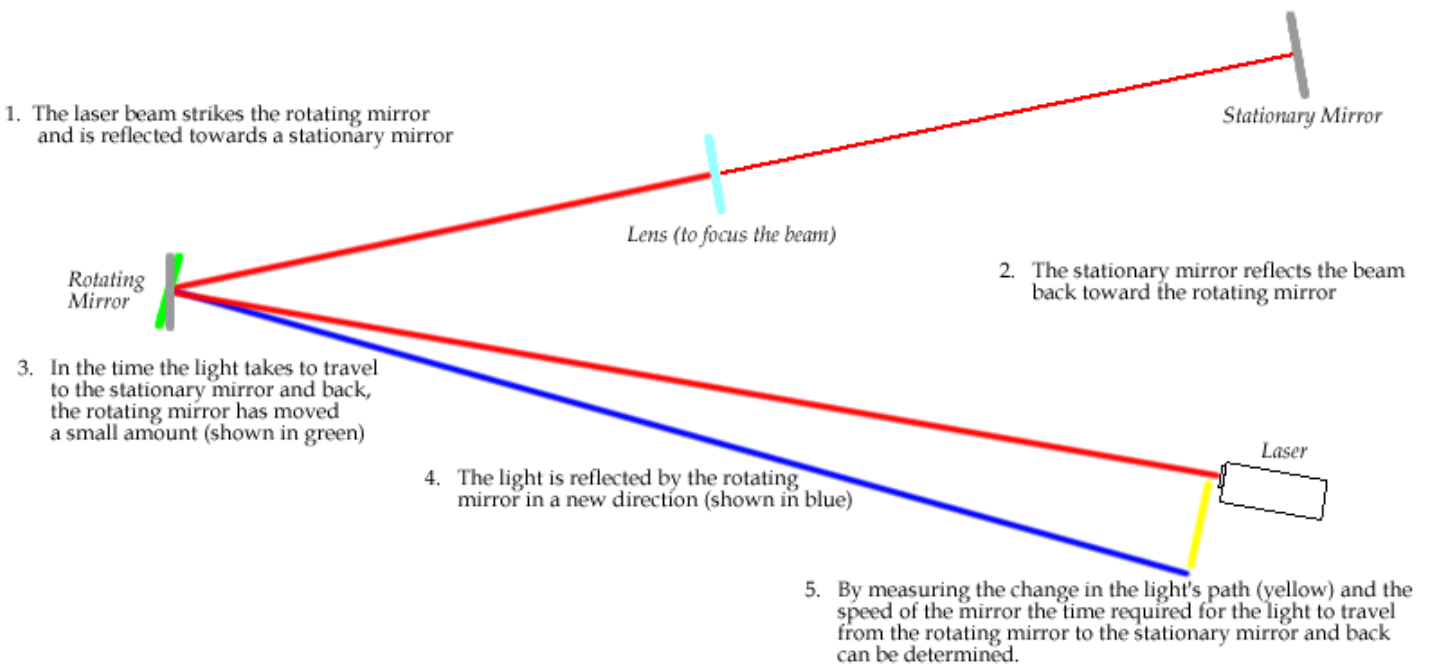
The difference between the WMW and t -test p-values and CI lengths (i.e. the WMW CI is narrower and the p-value smaller) reflects the effect of the outliers on the sensitivity of the standard tests and CI.

I conducted a pooled-variance two-sample t -test on ranks to show you that the p-value is close to the WMW p-value, as expected.

```
rank(met.long$cool)
## [1] 9.0 13.0 7.5 11.5 15.0 19.0 14.0 18.0 7.5 17.0 11.5 5.5 3.0
## [14] 16.0 5.5 1.0 2.0 4.0 10.0
by(rank(met.long$cool), met.long$site, summary)
## met.long$site: Uwet
##   Min. 1st Qu. Median   Mean 3rd Qu.   Max.
##   7.5     9.0    13.0   12.7    15.0   19.0
## -----
## met.long$site: Walker
##   Min. 1st Qu. Median   Mean 3rd Qu.   Max.
##   1.00    3.25    5.50    7.55   11.10   17.00
# note: the CI for ranks is not interpretable
t.test(rank(met.long$cool) ~ met.long$site, var.equal = TRUE)
##
## Two Sample t-test
##
## data: rank(met.long$cool) by met.long$site
## t = 2.208, df = 17, p-value = 0.04125
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 0.2305 10.1140
## sample estimates:
## mean in group Uwet mean in group Walker
## 12.72                7.55
```

Example: Newcombe's Data Experiments of historical importance were performed beginning in the eighteenth century to determine physical constants, such as the mean density of the earth, the distance from the earth to the sun, and the velocity of light. An interesting series of experiments to determine the velocity of light was begun in 1875. The first method used, and reused with refinements several times thereafter, was the rotating mirror method³. In this method a beam of light is reflected off a rapidly rotating mirror to a fixed mirror at a carefully measured distance from the source. The returning light is re-reflected from the rotating mirror at a different angle, because the mirror has turned slightly during the passage of the corresponding light pulses. From the speed of rotation of the mirror and from careful measurements of the angular difference between the outward-bound and returning light beams, the passage time of light can be calculated for the given distance. After averaging several calculations and applying various corrections, the experimenter can combine mean passage time and distance for a determination of the velocity of light. Simon Newcombe, a distinguished American scientist, used this method during the year 1882 to generate the passage time measurements given below, in microseconds. The travel path for this experiment was 3721 meters in length, extending from Ft. Meyer, on the west bank of the Potomac River in Washington, D.C., to a fixed mirror at the base of the Washington Monument.

³[http://en.wikipedia.org/wiki/File:Speed_of_light_\(foucault\).PNG](http://en.wikipedia.org/wiki/File:Speed_of_light_(foucault).PNG)



The problem is to determine a 95% CI for the “true” passage time, which is taken to be the typical time (mean or median) of the population of measurements that were or could have been taken by this experiment.

Example: Newcombe's Data

```
time <- c(24.828, 24.833, 24.834, 24.826, 24.824, 24.756  
       , 24.827, 24.840, 24.829, 24.816, 24.798, 24.822  
       , 24.824, 24.825, 24.823, 24.821, 24.830, 24.829  
       , 24.831, 24.824, 24.836, 24.819, 24.820, 24.832  
       , 24.836, 24.825, 24.828, 24.828, 24.821, 24.829  
       , 24.837, 24.828, 24.830, 24.825, 24.826, 24.832  
       , 24.836, 24.830, 24.836, 24.826, 24.822, 24.823  
       , 24.827, 24.828, 24.831, 24.827, 24.827, 24.827  
       , 24.826, 24.826, 24.832, 24.833, 24.832, 24.824  
       , 24.839, 24.824, 24.832, 24.828, 24.825, 24.825  
       , 24.829, 24.828, 24.816, 24.827, 24.829, 24.823)
```

```
library(nortest)
```

```
ad.test(time)
```

##

```
## Anderson-Darling normality test
```

##

```
## data: time
```

```
## A = 5.884, p-value = 1.217e-14
```

```
# Histogram overlaid with kernel density curve
```

```
Passage_df <- data.frame(time)
```

```
p1 <- ggplot(Passage_df, aes(x = time))
```

```
# Histogram with density instead of count on y-axis
```

```
p1 <- p1 + geom_histogram(aes(y=..density..))
```

, binwidth=0.001

, colour="black", fill="white")

```

# Overlay with transparent density plot
p1 <- p1 + geom_density(alpha=0.1, fill="#FF6666")
p1 <- p1 + geom_point(aes(y = -1)
                      , position = position_jitter(height = .5)
                      , alpha = 1/5)

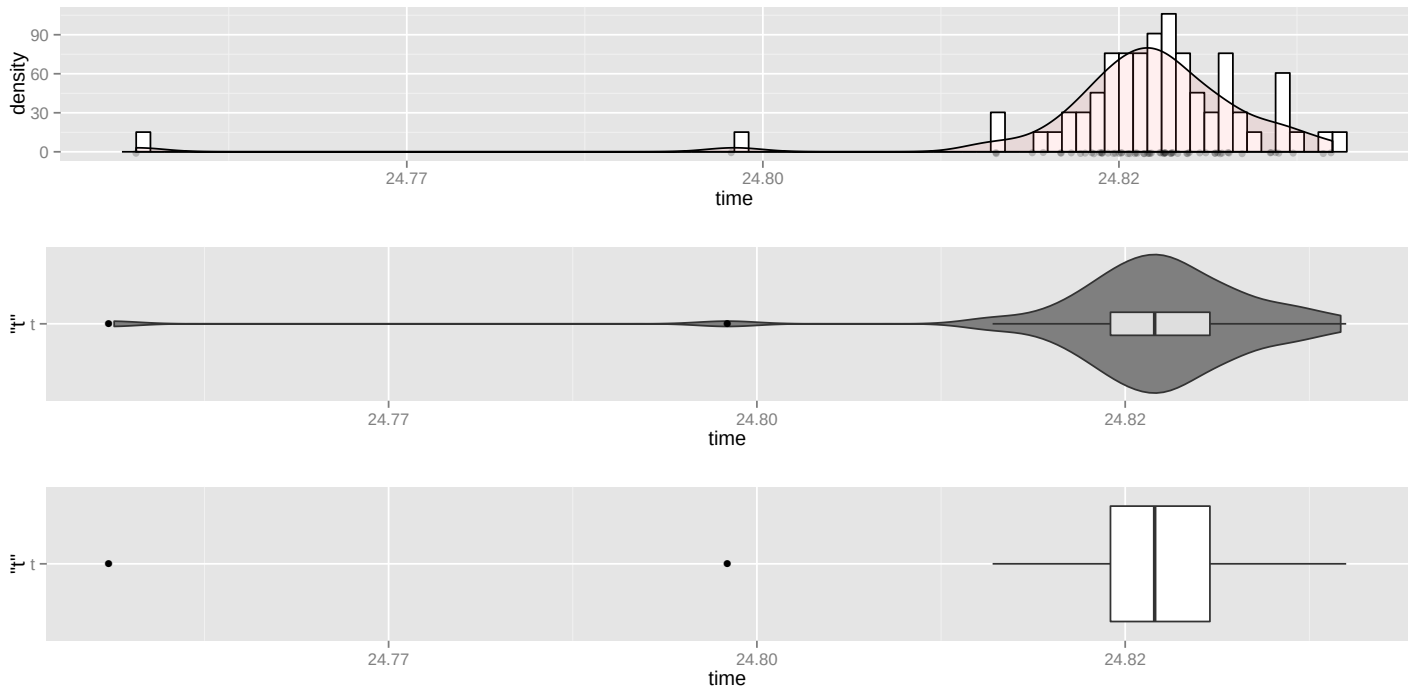
# violin plot
p2 <- ggplot(Passage_df, aes(x = "t", y = time))
p2 <- p2 + geom_violin(fill = "gray50")
p2 <- p2 + geom_boxplot(width = 0.2, alpha = 3/4)
p2 <- p2 + coord_flip()

# boxplot
p3 <- ggplot(Passage_df, aes(x = "t", y = time))
p3 <- p3 + geom_boxplot()
p3 <- p3 + coord_flip()

library(gridExtra)
grid.arrange(p1, p2, p3, ncol=1)

## Warning: position_stack requires constant width: output may be incorrect

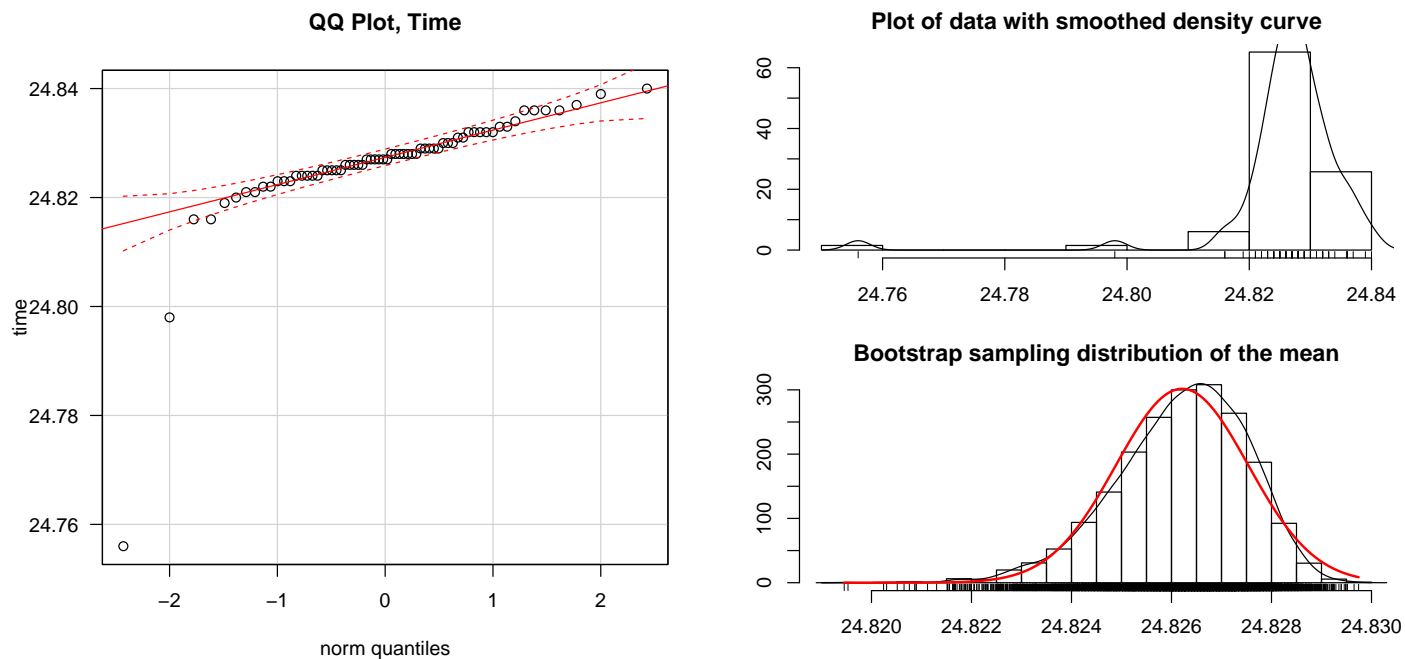
```



```

par(mfrow=c(1,1))
library(car)
qqPlot(time, las = 1, id.n = 0, id.cex = 1, lwd = 1, main="QQ Plot, Time")
bs.one.samp.dist(time)

```



The data set is skewed to the left, due to the presence of two extreme outliers that could potentially be misrecorded observations. Without additional information I would be hesitant to apply normal theory methods (the t -test), even though the sample size is “large” (bootstrap sampling distribution is still left-skewed). Furthermore, the t -test still suffers from a lack of robustness of sensitivity, even in large samples. A formal QQ-plot and normal test rejects, at the 0.01 level, the normality assumption needed for the standard methods.

The table below gives 95% t , sign, and Wilcoxon CIs. I am more comfortable with the sign CI for the population median than the Wilcoxon method, which assumes symmetry.

```
t.sum <- t.test(time)
t.sum$conf.int
## [1] 24.82 24.83
## attr(,"conf.level")
## [1] 0.95
diff(t.test(time)$conf.int)
## [1] 0.005283

s.sum <- SIGN.test(time)
##
## One-sample Sign-Test
##
## data: time
## s = 66, p-value < 2.2e-16
## alternative hypothesis: true median is not equal to 0
## 95 percent confidence interval:
## 24.83 24.83
```

```

## sample estimates:
## median of x
##      24.83
s.sum[2,c(2,3)]
## L.E.pt U.E.pt
## 24.83 24.83
diff(s.sum[2,c(2,3)])
## U.E.pt
## 0.0025

w.sum <- wilcox.test(time, conf.int=TRUE)
w.sum$conf.int
## [1] 24.83 24.83
## attr(,"conf.level")
## [1] 0.95
diff(w.sum$conf.int)
## [1] 0.002488

```

parameter	Method	CI Limits	Width
mean	t	(24.8236, 24.8289)	0.0053
median	sign	(24.8260, 24.8285)	0.0025
median	Wilcoxon	(24.8260, 24.8285)	0.0025

Note the big difference between the nonparametric and the t -CI. The nonparametric CIs are about 1/2 as wide as the t -CI. This reflects the impact that outliers have on the standard deviation, which directly influences the CI width.

6.5 Alternatives for ANOVA and Planned Comparisons

The classical ANOVA assumes that the populations have normal frequency curves and the populations have equal variances (or spreads). You learned formal tests for these assumptions in Chapter 5. When the assumptions do not hold, you can try one of the following two approaches. Before describing alternative methods, I will note that deviations from normality in one or more samples might be expected in a comparison involving many samples. You should downplay small deviations from normality in problems involving many samples.

6.5.1 Kruskal-Wallis ANOVA

The **Kruskal-Wallis** (KW) test is a non-parametric method for testing the hypothesis of equal population medians against the alternative that not all population medians are equal. The procedure assumes you have independent random samples from populations with frequency curves having **identical shapes and spreads**. The KW ANOVA is essentially the standard ANOVA based on ranked data. That is, we combine the samples, rank the observations from smallest to largest, and then return the ranks to the original samples and do the standard ANOVA using the ranks. The KW ANOVA is a multiple sample analog of the Wilcoxon-Mann-Whitney two sample procedure. Hence, multiple comparisons for a KW analysis, be they FSD or Bonferroni comparisons, are based on the two sample WMW procedure.

6.5.2 Transforming Data

The distributions in many data sets are skewed to the right with outliers. If the sample spreads, say s and IQR, increase with an increasing mean or median, you can often **transform data** to a scale where the normality and the constant spread assumption are more nearly satisfied. The transformed data are analyzed using the standard ANOVA. The two most commonly used transforms for this problem are the square root and natural logarithm, provided the data are non-negative⁴.

If the original distributions are nearly symmetric, but heavy-tailed, non-linear transformations will tend to destroy the symmetry. Many statisticians recommend methods based on trimmed means for such data. These methods are not commonly used by other researchers.

⁴The aim behind the choice of a **variance-stabilizing transformation** is to find a simple function f to apply to values y in a data set to create new values $y' = f(y)$ such that the variability of the values y' is not related to their mean value. For example, suppose that the values y are realizations from a Poisson distribution. Because for the Poisson distribution the variance is identical to the mean, the variance varies with the mean. However, if the simple variance-stabilizing transformation $y' = \sqrt{y}$ is applied, the sampling variance will be independent of the mean. A few distributional examples are provided in the table below.

Distribution	Variance= $g(\text{mean})$	Transformation $y' = f(y)$
Poisson	$\sigma^2 = \mu$	$y' = \sqrt{y}$
binomial	$\sigma^2 = \mu(1 - \mu)$	$y' = \arcsin(\sqrt{y})$
lognormal	$\sigma^2 = \mu^2$	$y' = \log(y)$

Example: Hydrocarbon (HC) Emissions Data These data are the HC emissions at idling speed, in ppm, for automobiles of different years of manufacture. The data are a random sample of all automobiles tested at an Albuquerque shopping center. (It looks like we need to find some newer cars!)

```
#### Example: Hydrocarbon (HC) Emissions Data
```

```
emis <- read.table(text="
```

	Pre-y63	y63-7	y68-9	y70-1	y72-4
2351	620	1088	141	140	
1293	940	388	359	160	
541	350	111	247	20	
1058	700	558	940	20	
411	1150	294	882	223	
570	2000	211	494	60	
800	823	460	306	20	
630	1058	470	200	95	
905	423	353	100	360	
347	900	71	300	70	
NA	405	241	223	220	
NA	780	2999	190	400	
NA	270	199	140	217	
NA	NA	188	880	58	
NA	NA	353	200	235	
NA	NA	117	223	1880	
NA	NA	NA	188	200	
NA	NA	NA	435	175	
NA	NA	NA	940	85	
NA	NA	NA	241	NA	

```
", header=TRUE)
```

```
#emis
```

```
# convert to long format
```

```
emis.long <- melt(emis,
                    variable.name = "year",
                    value.name = "hc",
                    na.rm = TRUE
      )
```

```
## Using as id variables
```

```
# naming variables manually, the variable.name and value.name not working 11/2012
```

```
names(emis.long) <- c("year", "hc")
```

```
# summary of each year
```

```
by(emis.long$hc, emis.long$year, summary)
```

```
## emis.long$year: Pre.y63
```

	Min.	1st Qu.	Median	Mean	3rd Qu.	Max.
##	347	548	715	891	1020	2350

```
## -----
```

```
## emis.long$year: y63.7
```

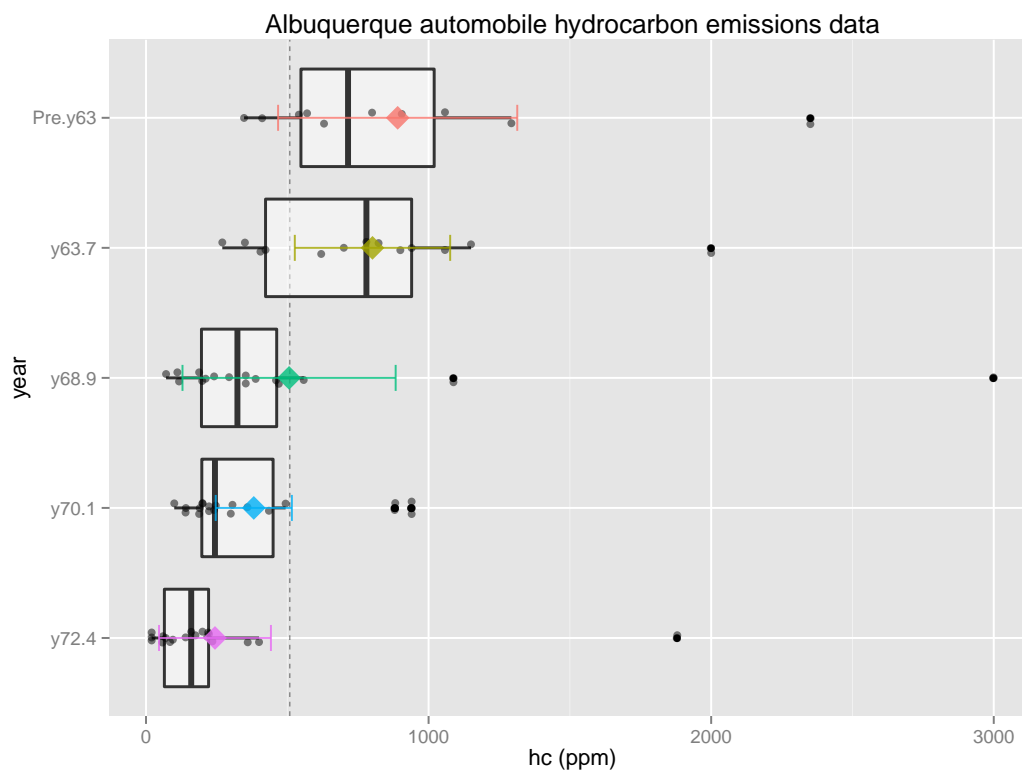
	Min.	1st Qu.	Median	Mean	3rd Qu.	Max.
##	270	423	780	801	940	2000

```

## -----
## emis.long$year: y68.9
##   Min. 1st Qu. Median     Mean 3rd Qu.    Max.
##    71      196    324      506     462    3000
## -----
## emis.long$year: y70.1
##   Min. 1st Qu. Median     Mean 3rd Qu.    Max.
##   100      198    244      381     450    940
## -----
## emis.long$year: y72.4
##   Min. 1st Qu. Median     Mean 3rd Qu.    Max.
##    20       65    160      244     222   1880
# IQR and sd of each year
by(emis.long$hc, emis.long$year, function(X) { c(IQR(X), sd(X), length(X)) })
## emis.long$year: Pre.y63
## [1] 471.5 591.6 10.0
## -----
## emis.long$year: y63.7
## [1] 517.0 454.9 13.0
## -----
## emis.long$year: y68.9
## [1] 266.2 707.8 16.0
## -----
## emis.long$year: y70.1
## [1] 252.2 287.9 20.0
## -----
## emis.long$year: y72.4
## [1] 156.5 410.8 19.0

# Plot the data using ggplot
library(ggplot2)
p <- ggplot(emis.long, aes(x = year, y = hc))
# plot a reference line for the global mean (assuming no groups)
p <- p + geom_hline(yintercept = mean(emis.long$hc),
                      colour = "black", linetype = "dashed", size = 0.3, alpha = 0.5)
# boxplot, size=.75 to stand out behind CI
p <- p + geom_boxplot(size = 0.75, alpha = 0.5)
# points for observed data
p <- p + geom_point(position = position_jitter(w = 0.05, h = 0), alpha = 0.5)
# diamond at mean for each group
p <- p + stat_summary(fun.y = mean, geom = "point", shape = 18, size = 6,
                      aes(colour=year), alpha = 0.8)
# confidence limits based on normal distribution
p <- p + stat_summary(fun.data = "mean_cl_normal", geom = "errorbar",
                      width = .2, aes(colour=year), alpha = 0.8)
p <- p + labs(title = "Albuquerque automobile hydrocarbon emissions data") + ylab("hc (ppm)")
# to reverse order that years print, so oldest is first on top
p <- p + scale_x_discrete(limits = rev(levels(emis.long$year)) )
p <- p + coord_flip()
p <- p + theme(legend.position="none")
print(p)

```



The standard ANOVA shows significant differences among the mean HC emissions. However, the standard ANOVA is inappropriate because the distributions are extremely skewed to the right due to presence of outliers in each sample.

```

fit.e <- aov(hc ~ year, data = emis.long)
summary(fit.e)

##           Df  Sum Sq Mean Sq F value Pr(>F)
## year        4 4226834 1056709   4.34 0.0033 **
## Residuals   73 17759968  243287
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
fit.e
## Call:
##   aov(formula = hc ~ year, data = emis.long)
##
## Terms:
##   year  Residuals
## Sum of Squares 4226834 17759968
## Deg. of Freedom     4       73
##
## Residual standard error: 493.2
## Estimated effects may be unbalanced

```

The boxplots show that the typical HC emissions appear to decrease as the age of car increases (the simplest description). Although the spread in the samples, as measured by the IQR, also decreases as age increases, I am more

comfortable with the KW ANOVA, in part because the KW analysis is not too sensitive to differences in spreads among samples. This point is elaborated upon later. As described earlier, the KW ANOVA is essentially an ANOVA based on the ranks. I give below the ANOVA based on ranks and the output from the KW procedure. They give similar p-values, and lead to the conclusion that there are significant differences among the population median HC emissions. A simple description is that the population median emission tends to decrease with the age of the car. You should follow up this analysis with Mann-Whitney multiple comparisons.

```
# ANOVA of rank, for illustration that this is similar to what KW is doing
fit.er <- aov(rank(hc) ~ year, data = emis.long)
summary(fit.er)

##          Df Sum Sq Mean Sq F value    Pr(>F)
## year        4 16329   4082     12.8 5.7e-08 ***
## Residuals   73 23200    318
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

fit.er
## Call:
##   aov(formula = rank(hc) ~ year, data = emis.long)
##
## Terms:
##           year Residuals
## Sum of Squares 16329      23200
## Deg. of Freedom   4         73
##
## Residual standard error: 17.83
## Estimated effects may be unbalanced

# KW ANOVA
fit.ek <- kruskal.test(hc ~ year, data = emis.long)
fit.ek

##
## Kruskal-Wallis rank sum test
##
## data: hc by year
## Kruskal-Wallis chi-squared = 31.81, df = 4, p-value =
## 2.093e-06
```

It is common to transform the data to a log scale when the spread increases as the median or mean increases.

```
# log scale
emis.long$loghc <- log(emis.long$hc)
# summary of each year
by(emis.long$loghc, emis.long$year, summary)
```

```

## emis.long$year: Pre.y63
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.
##      5.85    6.31   6.57     6.63    6.93    7.76
## -----
## emis.long$year: y63.7
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.
##      5.60    6.05   6.66     6.55    6.85    7.60
## -----
## emis.long$year: y68.9
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.
##      4.26    5.28   5.78     5.76    6.14    8.01
## -----
## emis.long$year: y70.1
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.
##      4.61    5.29   5.50     5.71    6.11    6.85
## -----
## emis.long$year: y72.4
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.
##      3.00    4.17   5.08     4.84    5.40    7.54

# IQR and sd of each year
by(emis.long$loghc, emis.long$year, function(X) { c(IQR(X), sd(X), length(X)) })

## emis.long$year: Pre.y63
## [1] 0.6186 0.5702 10.0000
## -----
## emis.long$year: y63.7
## [1] 0.7985 0.5525 13.0000
## -----
## emis.long$year: y68.9
## [1] 0.8575 0.9062 16.0000
## -----
## emis.long$year: y70.1
## [1] 0.8216 0.6776 20.0000
## -----
## emis.long$year: y72.4
## [1] 1.229 1.139 19.000

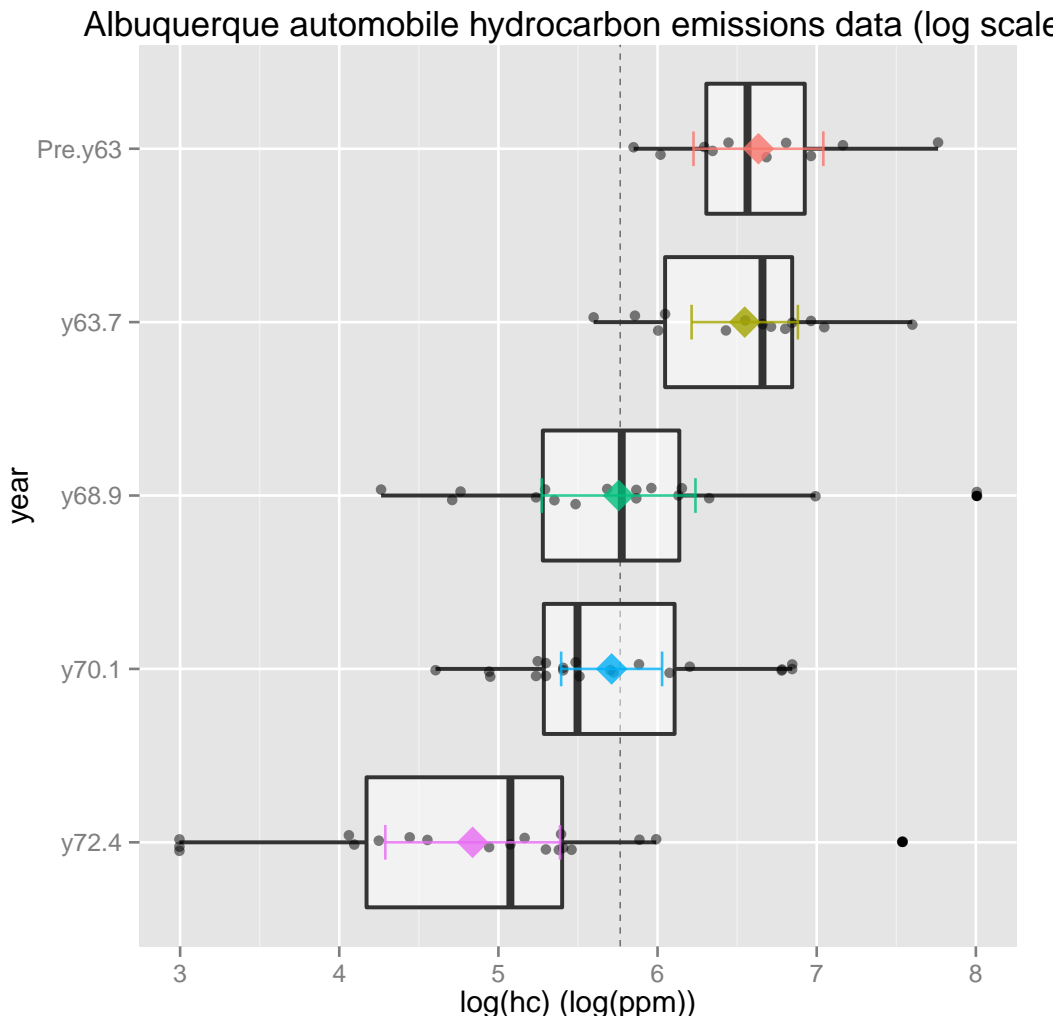
# Plot the data using ggplot
library(ggplot2)
p <- ggplot(emis.long, aes(x = year, y = loghc))
# plot a reference line for the global mean (assuming no groups)
p <- p + geom_hline(yintercept = mean(emis.long$loghc),
                      colour = "black", linetype = "dashed", size = 0.3, alpha = 0.5)
# boxplot, size=.75 to stand out behind CI
p <- p + geom_boxplot(size = 0.75, alpha = 0.5)
# points for observed data
p <- p + geom_point(position = position_jitter(w = 0.05, h = 0), alpha = 0.5)
# diamond at mean for each group
p <- p + stat_summary(fun.y = mean, geom = "point", shape = 18, size = 6,
                      aes(colour=year), alpha = 0.8)
# confidence limits based on normal distribution
p <- p + stat_summary(fun.data = "mean_cl_normal", geom = "errorbar",

```

```

width = .2, aes(colour=year), alpha = 0.8)
p <- p + labs(title = "Albuquerque automobile hydrocarbon emissions data (log scale)")
p <- p + ylab("log(hc) (log(ppm))")
# to reverse order that years print, so oldest is first on top
p <- p + scale_x_discrete(limits = rev(levels(emis.long$year)) )
p <- p + coord_flip()
p <- p + theme(legend.position="none")
print(p)

```



After transformation, the samples have roughly the same spread (IQR and s) and shape. The transformation does not completely eliminate the outliers. However, I am more comfortable with a standard ANOVA on this scale than with the original data. A difficulty here is that the ANOVA is comparing population mean log HC emission (so interpretations are on the log ppm scale, instead of the natural ppm scale). Summaries for the ANOVA on the log hydrocarbon emissions levels are given below.

```

# ANOVA of rank, for illustration that this is similar to what KW is doing
fit.le <- aov(loghc ~ year, data = emis.long)
summary(fit.le)

```

```

##          Df Sum Sq Mean Sq F value Pr(>F)
## year        4   31.9    7.97    11.4 3e-07 ***
## Residuals   73   51.0    0.70
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
fit.le
## Call:
##   aov(formula = loghc ~ year, data = emis.long)
##
## Terms:
##           year Residuals
## Sum of Squares 31.90      50.98
## Deg. of Freedom 4          73
##
## Residual standard error: 0.8357
## Estimated effects may be unbalanced

# KW ANOVA -- same conclusions as original scale, since based on ranks
fit.lek <- kruskal.test(loghc ~ year, data = emis.long)
fit.lek

##
## Kruskal-Wallis rank sum test
##
## data: loghc by year
## Kruskal-Wallis chi-squared = 31.81, df = 4, p-value =
## 2.093e-06

```

The boxplot of the log-transformed data reinforces the reasonableness of the original KW analysis. Why? The log-transformed distributions have fairly similar shapes and spreads, so a KW analysis on these data is sensible. The ranks for the original and log-transformed data are identical, so the KW analyses on the log-transformed data and the original data must lead to the same conclusions. This suggests that the KW ANOVA is not overly sensitive to differences in spreads among the samples.

There are two reasonable analyses here: the standard ANOVA using log HC emissions, and the KW analysis of the original data. The first analysis gives a comparison of mean log HC emissions. The second involves a comparison of median HC emissions. A statistician would present both analyses to the scientist who collected the data to make a decision on which was more meaningful (independently of the results⁵!). Multiple comparisons would be performed relative to the selected analysis (*t*-tests for ANOVA or WMW-tests for KW ANOVA).

⁵It is unethical to choose a method based on the results it gives.

Example: Hodgkin's Disease Study Plasma bradykininogen levels were measured in normal subjects, in patients with active Hodgkin's disease, and in patients with inactive Hodgkin's disease. The globulin bradykininogen is the precursor substance for bradykinin, which is thought to be a chemical mediator of inflammation. The data (in micrograms of bradykininogen per milliliter of plasma) are displayed below. The three samples are denoted by **nc** for normal controls, **ahd** for active Hodgkin's disease patients, and **ihd** for inactive Hodgkin's disease patients.

The medical investigators wanted to know if the three samples differed in their bradykininogen levels. Carry out the statistical analysis you consider to be most appropriate, and state your conclusions to this question.

Read in the data, look at summaries on the original scale, and create a plot. Also, look at summaries on the log scale and create a plot.

```
#### Example: Hodgkin's Disease Study
```

```
hd <- read.table(text="
```

nc	ahd	ihd
5.37	3.96	5.37
5.80	3.04	10.60
4.70	5.28	5.02
5.70	3.40	14.30
3.40	4.10	9.90
8.60	3.61	4.27
7.48	6.16	5.75
5.77	3.22	5.03
7.15	7.48	5.74
6.49	3.87	7.85
4.09	4.27	6.82
5.94	4.05	7.90
6.38	2.40	8.36
9.24	5.81	5.72
5.66	4.29	6.00
4.53	2.77	4.75
6.51	4.40	5.83
7.00	NA	7.30
6.20	NA	7.52
7.04	NA	5.32
4.82	NA	6.05
6.73	NA	5.68
5.26	NA	7.57
NA	NA	5.68
NA	NA	8.91
NA	NA	5.39
NA	NA	4.40
NA	NA	7.13

```
", header=TRUE)
```

```

#hd

# convert to long format
hd.long <- melt(hd,
                 variable.name = "patient",
                 value.name = "level",
                 na.rm = TRUE
               )
## Using as id variables

# naming variables manually, the variable.name and value.name not working 11/2012
names(hd.long) <- c("patient", "level")
# summary of each patient
by(hd.long$level, hd.long$patient, summary)
## hd.long$patient: nc
##   Min. 1st Qu. Median   Mean 3rd Qu.   Max.
##   3.40    5.32   5.94    6.08   6.86    9.24
## -----
## hd.long$patient: ahd
##   Min. 1st Qu. Median   Mean 3rd Qu.   Max.
##   2.40    3.40   4.05    4.24   4.40    7.48
## -----
## hd.long$patient: ihd
##   Min. 1st Qu. Median   Mean 3rd Qu.   Max.
##   4.27    5.38   5.92    6.79   7.64   14.30

# IQR and sd of each patient
by(hd.long$level, hd.long$patient, function(X) { c(IQR(X), sd(X), length(X)) })
## hd.long$patient: nc
## [1] 1.550 1.362 23.000
## -----
## hd.long$patient: ahd
## [1] 1.000 1.303 17.000
## -----
## hd.long$patient: ihd
## [1] 2.255 2.176 28.000

# log scale
hd.long$loglevel <- log(hd.long$level)
# summary of each patient
by(hd.long$loglevel, hd.long$patient, summary)
## hd.long$patient: nc
##   Min. 1st Qu. Median   Mean 3rd Qu.   Max.
##   1.22    1.67   1.78    1.78   1.93    2.22
## -----
## hd.long$patient: ahd
##   Min. 1st Qu. Median   Mean 3rd Qu.   Max.
##   0.875   1.220   1.400   1.400   1.480   2.010
## -----
## hd.long$patient: ihd
##   Min. 1st Qu. Median   Mean 3rd Qu.   Max.
##   1.45    1.68   1.78    1.87   2.03    2.66

# IQR and sd of each patient
by(hd.long$loglevel, hd.long$patient, function(X) { c(IQR(X), sd(X), length(X)) })
## hd.long$patient: nc

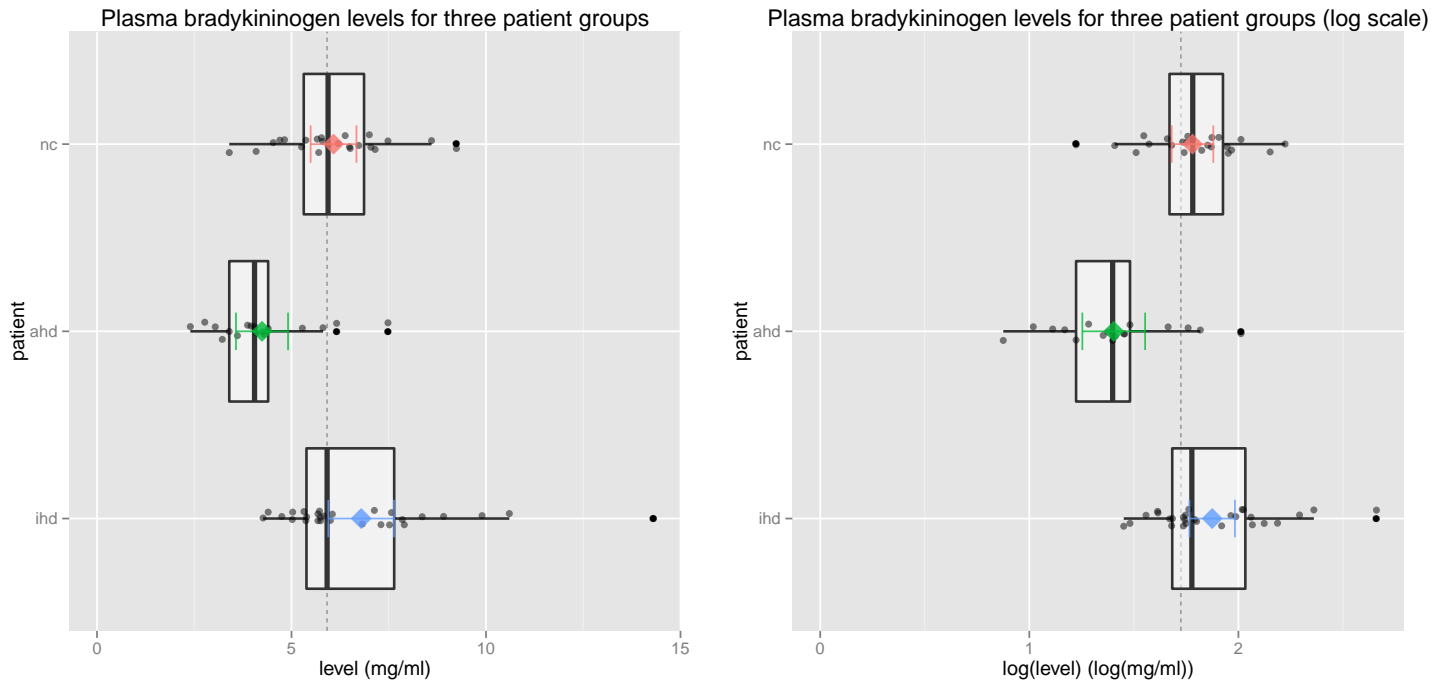
```

```
## [1] 0.2558 0.2303 23.0000
## -----
## hd.long$patient: ahd
## [1] 0.2578 0.2921 17.0000
## -----
## hd.long$patient: ihd
## [1] 0.3497 0.2803 28.0000

# Plot the data using ggplot
library(ggplot2)
p <- ggplot(hd.long, aes(x = patient, y = level))
# plot a reference line for the global mean (assuming no groups)
p <- p + geom_hline(yintercept = mean(hd.long$level),
                      colour = "black", linetype = "dashed", size = 0.3, alpha = 0.5)
# boxplot, size=.75 to stand out behind CI
p <- p + geom_boxplot(size = 0.75, alpha = 0.5)
# points for observed data
p <- p + geom_point(position = position_jitter(w = 0.05, h = 0), alpha = 0.5)
# diamond at mean for each group
p <- p + stat_summary(fun.y = mean, geom = "point", shape = 18, size = 6,
                      aes(colour=patient), alpha = 0.8)
# confidence limits based on normal distribution
p <- p + stat_summary(fun.data = "mean_cl_normal", geom = "errorbar",
                      width = .2, aes(colour=patient), alpha = 0.8)
p <- p + labs(title = "Plasma bradykininogen levels for three patient groups")
p <- p + ylab("level (mg/ml)")
# to reverse order that years print, so oldest is first on top
p <- p + scale_x_discrete(limits = rev(levels(hd.long$patient)) )
p <- p + ylim(c(0,max(hd.long$level)))
p <- p + coord_flip()
p <- p + theme(legend.position="none")
print(p)

## log scale
# Plot the data using ggplot
library(ggplot2)
p <- ggplot(hd.long, aes(x = patient, y = loglevel))
# plot a reference line for the global mean (assuming no groups)
p <- p + geom_hline(yintercept = mean(hd.long$loglevel),
                      colour = "black", linetype = "dashed", size = 0.3, alpha = 0.5)
# boxplot, size=.75 to stand out behind CI
p <- p + geom_boxplot(size = 0.75, alpha = 0.5)
# points for observed data
p <- p + geom_point(position = position_jitter(w = 0.05, h = 0), alpha = 0.5)
# diamond at mean for each group
p <- p + stat_summary(fun.y = mean, geom = "point", shape = 18, size = 6,
                      aes(colour=patient), alpha = 0.8)
# confidence limits based on normal distribution
p <- p + stat_summary(fun.data = "mean_cl_normal", geom = "errorbar",
                      width = .2, aes(colour=patient), alpha = 0.8)
p <- p + labs(title = "Plasma bradykininogen levels for three patient groups (log scale)")
p <- p + ylab("log(level) (log(mg/ml))")
# to reverse order that years print, so oldest is first on top
```

```
p <- p + scale_x_discrete(limits = rev(levels(hd.long$patient)) )
p <- p + ylim(c(0,max(hd.long$loglevel)))
p <- p + coord_flip()
p <- p + theme(legend.position="none")
print(p)
```



Although the spread (IQR, s) in the ihd sample is somewhat greater than the spread in the other samples, the presence of skewness and outliers in the boxplots is a greater concern regarding the use of the classical ANOVA. The shapes and spreads in the three samples are roughly identical, so a Kruskal-Wallis nonparametric ANOVA appears ideal. As a sidelight, I transformed plasma levels to a log scale to reduce the skewness and eliminate the outliers. The boxplots of the transformed data show reasonable symmetry across groups, but outliers are still present. I will stick with the Kruskal-Wallis ANOVA (although it would not be much of a problem to use the classical ANOVA on transformed data).

Let η_{nc} = population median plasma level for normal controls, η_{ahd} = population median plasma level for active Hodgkin's disease patients, and η_{ihd} = population median plasma level for inactive Hodgkin's disease patients. The KW test of $H_0 : \eta_{nc} = \eta_{ahd} = \eta_{ihd}$ versus $H_A : \text{not } H_0$ is highly significant ($p\text{-value}= 0.00003$), suggesting differences among the population median plasma levels. The Kruskal-Wallis ANOVA summary is given below.

```
# KW ANOVA
fit.h <- kruskal.test(level ~ patient, data = hd.long)
fit.h
##
## Kruskal-Wallis rank sum test
##
## data: level by patient
## Kruskal-Wallis chi-squared = 20.57, df = 2, p-value =
## 3.421e-05
```

I followed up the KW ANOVA with Bonferroni comparisons of the samples, using the Mann-Whitney two sample procedure. There are three comparisons, so an overall FER of 0.05 is achieved by doing the individual tests at the $0.05/3=0.0167$ level. Alternatively, you can use 98.33% CI for differences in population medians.

```
# with continuity correction in the normal approximation for the p-value
wilcox.test(hd$nc , hd$ahd, conf.int=TRUE, conf.level = 0.9833)
## Warning: cannot compute exact p-value with ties
## Warning: cannot compute exact confidence intervals with ties
##
## Wilcoxon rank sum test with continuity correction
##
## data: hd$nc and hd$ahd
## W = 329, p-value = 0.0002735
## alternative hypothesis: true location shift is not equal to 0
## 98.33 percent confidence interval:
##  0.8599 2.9001
## sample estimates:
## difference in location
##                      1.91
wilcox.test(hd$nc , hd$ihd, conf.int=TRUE, conf.level = 0.9833)
## Warning: cannot compute exact p-value with ties
## Warning: cannot compute exact confidence intervals with ties
##
## Wilcoxon rank sum test with continuity correction
##
## data: hd$nc and hd$ihd
## W = 276.5, p-value = 0.3943
## alternative hypothesis: true location shift is not equal to 0
## 98.33 percent confidence interval:
## -1.56  0.68
## sample estimates:
## difference in location
##                      -0.3414
wilcox.test(hd$ahd, hd$ihd, conf.int=TRUE, conf.level = 0.9833)
## Warning: cannot compute exact p-value with ties
## Warning: cannot compute exact confidence intervals with ties
```

```

## Wilcoxon rank sum test with continuity correction
##
## data: hd$ahd and hd$ihd
## W = 56, p-value = 2.143e-05
## alternative hypothesis: true location shift is not equal to 0
## 98.33 percent confidence interval:
## -3.50 -1.32
## sample estimates:
## difference in location
## -2.147

```

The only comparison with a p-value greater than 0.0167 involved the **nc** and **ihd** samples. The comparison leads to two groups, and is consistent with what we see in the boxplots.

ahd	nc	ihd
---	-----	

You have sufficient evidence to conclude that the plasma bradykininogen levels for active Hodgkin's disease patients (ahd) is lower than the population median levels for normal controls (nc) and for patients with inactive Hodgkin's disease (ihd). You do not have sufficient evidence to conclude that the population median levels for normal controls (nc) and for patients with inactive Hodgkin's disease (ihd) are different. The CIs give an indication of size of differences in the population medians.

6.5.3 Planned Comparisons

Bonferroni multiple comparisons are generally preferred to Fisher's least significant difference approach. Fisher's method does not control the familywise error rate and produces too many spurious significant differences (claims of significant differences that are due solely to chance variation and not to actual differences in population means). However, Bonferroni's method is usually very conservative when a large number of comparisons is performed — large differences in sample means are needed to claim significance. A way to reduce this conservatism is to avoid doing all possible comparisons. Instead, one should, when possible, decide *a priori* (before looking at the data) which comparisons are of primary interest, and then perform only those comparisons.

For example, suppose a medical study compares five new treatments with a control (a six group problem). The medical investigator may not be interested in all 15 possible comparisons, but only in which of the five treatments differ

on average from the control. Rather than performing the 15 comparisons, each at the say $0.05/15 = 0.0033$ level, she could examine the five comparisons of interest at the $0.05/5 = 0.01$ level. By deciding beforehand which comparisons are of interest, she can justify using a 0.01 level for the comparisons, instead of the more conservative 0.0033 level needed when doing all possible comparisons.

To illustrate this idea, consider the KW analysis of HC emissions. We saw that there are significant differences among the population median HC emissions. Given that the samples have a natural ordering

Sample	Year of manufacture
1	Pre-1963
2	63 – 67
3	68 – 69
4	70 – 71
5	72 – 74

you may primarily be interested in whether the population medians for cars manufactured in consecutive samples are identical. That is, you may be primarily interested in the following 4 comparisons:

Pre-1963	vs	63 – 67
63 – 67	vs	68 – 69
68 – 69	vs	70 – 71
70 – 71	vs	72 – 74

A Bonferroni analysis would carry out each comparison at the $0.05/4 = 0.0125$ level versus the $0.05/10 = 0.005$ level when all comparisons are done.

The following output was obtained for doing these four comparisons, based on Wilcoxon-Mann-Whitney two-sample tests (why?⁶). Two-year groups are claimed to be different if the p-value is 0.0125 or below, or equivalently, if a 98.75% CI for the difference in population medians does not contain zero.

```
#### Planned Comparisons
# with continuity correction in the normal approximation for the p-value
wilcox.test(emis$y63.7, emis$Pre.y63, conf.int=TRUE, conf.level = 0.9875)

## Warning: cannot compute exact p-value with ties
## Warning: cannot compute exact confidence intervals with ties
##
```

⁶The ANOVA is the multi-sample analog to the two-sample *t*-test for the mean, and the KW ANOVA is the multi-sample analog to the WMW two-sample test for the median. Thus, we follow up a KW ANOVA with WMW two-sample tests at the chosen multiple comparison adjusted error rate.

```
## Wilcoxon rank sum test with continuity correction
##
## data: emis$y63.7 and emis$Pre.y63
## W = 61.5, p-value = 0.8524
## alternative hypothesis: true location shift is not equal to 0
## 98.75 percent confidence interval:
## -530 428
## sample estimates:
## difference in location
## -15.48

wilcox.test(emis$y68.9, emis$y63.7 , conf.int=TRUE, conf.level = 0.9875)
## Warning: cannot compute exact p-value with ties
## Warning: cannot compute exact confidence intervals with ties
##
## Wilcoxon rank sum test with continuity correction
##
## data: emis$y68.9 and emis$y63.7
## W = 43, p-value = 0.007968
## alternative hypothesis: true location shift is not equal to 0
## 98.75 percent confidence interval:
## -709 -52
## sample estimates:
## difference in location
## -397.4

wilcox.test(emis$y70.1, emis$y68.9 , conf.int=TRUE, conf.level = 0.9875)
## Warning: cannot compute exact p-value with ties
## Warning: cannot compute exact confidence intervals with ties
##
## Wilcoxon rank sum test with continuity correction
##
## data: emis$y70.1 and emis$y68.9
## W = 156, p-value = 0.9112
## alternative hypothesis: true location shift is not equal to 0
## 98.75 percent confidence interval:
## -206 171
## sample estimates:
## difference in location
## -11

wilcox.test(emis$y72.4, emis$y70.1 , conf.int=TRUE, conf.level = 0.9875)
## Warning: cannot compute exact p-value with ties
## Warning: cannot compute exact confidence intervals with ties
##
## Wilcoxon rank sum test with continuity correction
##
## data: emis$y72.4 and emis$y70.1
## W = 92.5, p-value = 0.006384
## alternative hypothesis: true location shift is not equal to 0
## 98.75 percent confidence interval:
## -286 -6
## sample estimates:
```

```
## difference in location  
## -130
```

There are significant differences between the 1963-67 and 1968-69 samples, and between the 1970-71 and 1972-74 samples. You are 98.75% confident that the population median HC emissions for 1963-67 year cars is between 52 and 708.8 ppm greater than the population median for 1968-69 cars. Similarly, you are 98.75% confident that the population median HC emissions for 1970-71 year cars is between 6.1 and 285.9 ppm greater than the population median for 1972-74 cars. Overall, you are 95% confident among the four pairwise comparisons that you have not declared a difference significant when it isn't.

6.5.4 Two final ANOVA comments

It is not uncommon for researchers to combine data from groups not found to be significantly different. This is not, in general, a good practice. Just because you do not have sufficient evidence to show differences does not imply that you should treat the groups as if they are the same!

If the data distributions do not substantially deviate from normality, but the spreads are different across samples, you might consider the standard ANOVA followed with multiple comparisons using two-sample tests based on Satterthwaite's approximation.

6.6 Permutation tests

Permutation tests⁷ are a subset of non-parametric statistics. The basic premise is to use only the assumption that it is possible that all of the treatment groups are equivalent, and that every member of them is the same before sampling began (i.e., the position in the group to which they belong is not differentiable from other position before the positions are filled). From this, one can calculate a statistic and then see to what extent this statistic is special by seeing how likely it would be if the group assignments had been jumbled.

⁷[http://en.wikipedia.org/wiki/Resampling_\(statistics\)](http://en.wikipedia.org/wiki/Resampling_(statistics))

A permutation test (also called a randomization test, re-randomization test, or an exact test) is a type of statistical significance test in which the distribution of the test statistic under the null hypothesis is obtained by calculating all possible values of the test statistic under **rearrangements of the labels** on the observed data points. In other words, the method by which treatments are allocated to subjects in an experimental design is mirrored in the analysis of that design. If the labels are exchangeable under the null hypothesis, then the resulting tests yield exact significance levels. Confidence intervals can then be derived from the tests. The theory has evolved from the works of R.A. Fisher and E.J.G. Pitman in the 1930s.

Let's illustrate the basic idea of a permutation test using the Meteorites example. Suppose we have two groups Uwet and Walker whose sample means are \bar{Y}_U and \bar{Y}_W , and that we want to test, at 5% significance level, whether they come from the same distribution. Let $n_U = 9$ and $n_W = 10$ be the sample size corresponding to each group. The permutation test is designed to determine whether the observed difference between the sample means is large enough to reject the null hypothesis $H_0 : \mu_U = \mu_W$, that the two groups have identical means.

The test proceeds as follows. First, the difference in means between the two samples is calculated: this is the observed value of the test statistic, $T_{(obs)}$. Then the observations of groups Uwet and Walker are pooled.

```
#### Permutation tests
# Calculated the observed difference in means
  # met.long includes both Uwet and Walker groups
Tobs <- mean(met.long[(met.long$site == "Uwet"), 2]) -
  mean(met.long[(met.long$site == "Walker"), 2])
Tobs
## [1] 0.2522
```

Next, the difference in sample means is calculated and recorded for every possible way of dividing these pooled values into two groups of size $n_U = 9$ and $n_W = 10$ (i.e., for every permutation of the group labels Uwet and Walker). The set of these calculated differences is the exact distribution of possible differences under the null hypothesis that group label does not matter. This exact distribution can be approximated by drawing a large number of random permutations.

```
# Plan:
#   Initialize a vector in which to store the R number of difference of means.
#   Calculate R differences in means for R permutations, storing the results.
#       Note that there are prod(1:19) = 10^17 total permutations,
```

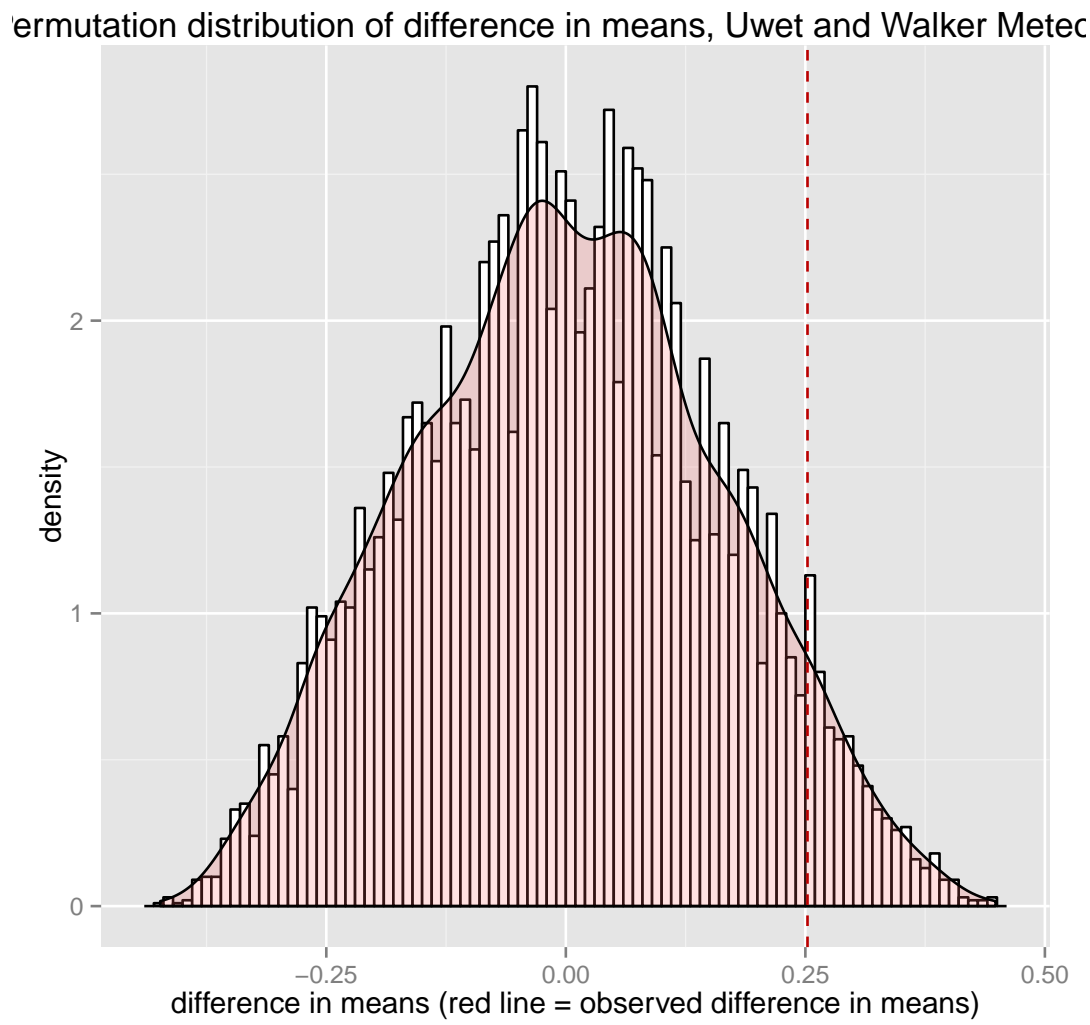
```
#       but the R repetitions will serve as a good approximation.
# Plot the permutation null distribution with an indication of the Tobs.

# R = a large number of repetitions
R <- 1e4
# initialize the vector of difference of means from the permutations
Tperm <- rep(NA, R)
# For each of R repetitions, permute the Uwet and Walker labels,
# calculate the difference of means with the permuted labels,
# and store the result in the i.R'th position of Tperm.
for (i.R in 1:R) {
  # permutation of 19 = 9+10 integers 1, 2, ..., 19
  ind.perm <- sample.int(nrow(met.long))
  # identify as "TRUE" numbers 1, ..., 9 (the number of Uwet labels)
  lab.U <- (ind.perm <= sum(met.long$site == "Uwet"))                #$
  # identify as "TRUE" numbers 10, ..., 19 (the number of Walker labels)
  # that is, all the non-Uwet labels
  lab.W <- !lab.U

  # calculate the difference in means and store in Tperm at index i.R
  Tperm[i.R] <- mean(met.long[lab.U, 2]) - mean(met.long[lab.W, 2])
}

# Plot the permutation null distribution with an indication of the Tobs.
dat <- data.frame(Tperm)

library(ggplot2)
p <- ggplot(dat, aes(x = Tperm))
#p <- p + scale_x_continuous(limits=c(-20,+20))
p <- p + geom_histogram(aes(y=..density..),
                        , binwidth=0.01
                        , colour="black", fill="white")
  # Overlay with transparent density plot
p <- p + geom_density(alpha=0.2, fill="#FF6666")
#p <- p + geom_point(aes(y = -0.05)
#                      , position = position_jitter(height = 0.01)
#                      , alpha = 1/5)
  # vertical line at Tobs
p <- p + geom_vline(aes(xintercept=Tobs), colour="#BB0000", linetype="dashed")
p <- p + labs(title = "Permutation distribution of difference in means, Uwet and Walker Meteor")
p <- p + xlab("difference in means (red line = observed difference in means)")
print(p)
```



Notice the contrast in this permutation distribution of the difference in means from a normal distribution.

The one-sided p-value of the test is calculated as the proportion of sampled permutations where the difference in means was at least as extreme as $T_{(obs)}$. The two-sided p-value of the test is calculated as the proportion of sampled permutations where the absolute difference was at least as extreme as $|T_{(obs)}|$.

```
# Calculate a two-sided p-value.
p.upper <- sum((Tperm >= abs(Tobs))) / R
p.upper
## [1] 0.0619

p.lower <- sum((Tperm <= -abs(Tobs))) / R
p.lower
## [1] 0.061

p.twosided <- p.lower + p.upper
p.twosided
## [1] 0.1229
```

Note that the two-sided p-value of 0.1229 is consistent, in this case, with the two-sample t -test p-values of 0.1134 (pooled) and 0.1290 (Satterthwaite), but different from 0.0497 (WMW). The permutation is a comparison of means **without** the normality assumption, though requires that the observations are exchangeable between populations under H_0 .

If the only purpose of the test is reject or not reject the null hypothesis, we can as an alternative sort the recorded differences, and then observe if $T_{(\text{obs})}$ is contained within the middle 95% of them. If it is not, we reject the hypothesis of equal means at the 5% significance level.

6.6.1 Automated in R

The `lmp` package provides permutation tests for linear models (including t -tests and ANOVA) and is particularly easy to implement. You can use it for all manner of ANOVA/ANCOVA designs, as well as simple, polynomial, and multiple regression. Simply use `lmp()` and `aovp()` where you would have used `lm()` and `aov()`. Note that the t -test can be performed with `lmp()`.

Below I calculate the standard t -test for the Meteorite data using `t.test()` and `lmp()`, then compare that with `lmp()` and what we calculated using our calculation of the permutation test.

```
# standard two-sample t-test with equal variances
t.summary <- t.test(cool ~ site, data = met.long, var.equal = TRUE)
t.summary

##
## Two Sample t-test
##
## data: cool by site
## t = 1.669, df = 17, p-value = 0.1134
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.06663 0.57107
## sample estimates:
## mean in group Uwet mean in group Walker
##           0.4522             0.2000

# linear model form of t-test, "siteWalker" has estimate, se, t-stat, and p-value
lm.summary <- lm(cool ~ site, data = met.long)
summary(lm.summary)

##
## Call:
## lm(formula = cool ~ site, data = met.long)
```

```

## 
## Residuals:
##   Min     1Q Median     3Q    Max
## -0.292 -0.196 -0.160  0.025  0.748
##
## 
## Coefficients:
##             Estimate Std. Error t value Pr(>|t|)
## (Intercept)  0.452     0.110    4.12  0.00071 ***
## siteWalker -0.252     0.151   -1.67  0.11344
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.329 on 17 degrees of freedom
## Multiple R-squared:  0.141, Adjusted R-squared:  0.0902
## F-statistic: 2.79 on 1 and 17 DF,  p-value: 0.113

# permutation test version
library(lmPerm)
lmp.summary <- lmp(cool ~ site, data = met.long)
## [1] "Settings: unique SS"
summary(lmp.summary)

##
## Call:
## lmp(formula = cool ~ site, data = met.long)
##
## Residuals:
##   Min     1Q Median     3Q    Max
## -0.292 -0.196 -0.160  0.025  0.748
##
## 
## Coefficients:
##             Estimate Iter Pr(Prob)
## site1      0.126   604     0.14
## 
## Residual standard error: 0.329 on 17 degrees of freedom
## Multiple R-Squared: 0.141, Adjusted R-squared: 0.0902
## F-statistic: 2.79 on 1 and 17 DF,  p-value: 0.113

```

The permutation test gives a p-value of 0.1424 which is a bit different from our previously calculated 0.1229, and is different each time you run the routine (since it is based on a sampling approximation, just as we coded). The function also provides, as a reference, the pooled two-sample test at the bottom.

For the emisions data, the permutation test gives this result.

```

# permutation test version
library(lmPerm)
aovp.summary <- aovp(hc ~ year, data = emis.long)
## [1] "Settings: unique SS"
aovp.summary

```

```

## Call:
##   aovp(formula = hc ~ year, data = emis.long)
##
## Terms:
##           year Residuals
## Sum of Squares 4226834 17759968
## Deg. of Freedom     4         73
##
## Residual standard error: 493.2
## Estimated effects may be unbalanced

summary(aovp.summary)

## Component 1 :
##              Df R Sum Sq R Mean Sq Iter Pr(Prob)
## year          4 4226834 1056709 5000  0.0084 **
## Residuals    73 17759968    243287
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

```

Thus the overall ANOVA rejects the null hypothesis of all equal means. A followup set of pairwise tests can be done with `lmp()`.

```

# these are the levels of the factor, the first is the baseline group
levels(emis.long$year)
## [1] "Pre.y63" "y63.7"   "y68.9"   "y70.1"   "y72.4"

# permutation test version
library(lmPerm)
lmp.summary <- lmp(hc ~ year, data = emis.long)
## [1] "Settings: unique SS"
summary(lmp.summary)

##
## Call:
## lmp(formula = hc ~ year, data = emis.long)
##
## Residuals:
##      Min    1Q Median    3Q   Max
## -543.6 -224.1 -126.4   44.2 2492.7
##
## Coefficients:
##             Estimate Iter Pr(Prob)
## year1      325.8 5000  <2e-16 ***
## year2      236.7  798   0.112
## year3     -58.5  221   0.312
## year4     -183.3 1793   0.053 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 493 on 73 degrees of freedom
## Multiple R-Squared: 0.192, Adjusted R-squared: 0.148
## F-statistic: 4.34 on 4 and 73 DF, p-value: 0.00331

```

The year1, year2, year3, and year4 correspond to the 2nd through 5th groups. The Pr(Prob) column gives the p-values for the permutation test comparing each of the 2nd through 5th groups to the 1st group (year0). For the other comparisons, reorder the factor levels and rerun.

(Note, I thought `lmp()` would use the baseline factor level as the intercept, which would result in the year1 ... year4 estimates being the pairwise differences from the intercept group. Therefore, by changing which group is the baseline intercept group, I thought you could get pairwise tests between all pairs. However, `lmp()` must be performing some internal factor level ordering ordering because it always seems to use the third group in the `emis.long$year` factor as the baseline intercept. Unfortunately, I don't know how to use `lmp()` to get the other pairwise comparisons.)

6.7 Density estimation

Density estimation is like a histogram: It is a method for visualizing the shape of a univariate distribution (there are methods for doing multivariate density estimation as well, but we will ignore those for the time being). In fact, I snuck in density estimation in the first chapter and have been using it all along! Let's experiment with Newcombe's speed-of-light data (excluding the two outliers).

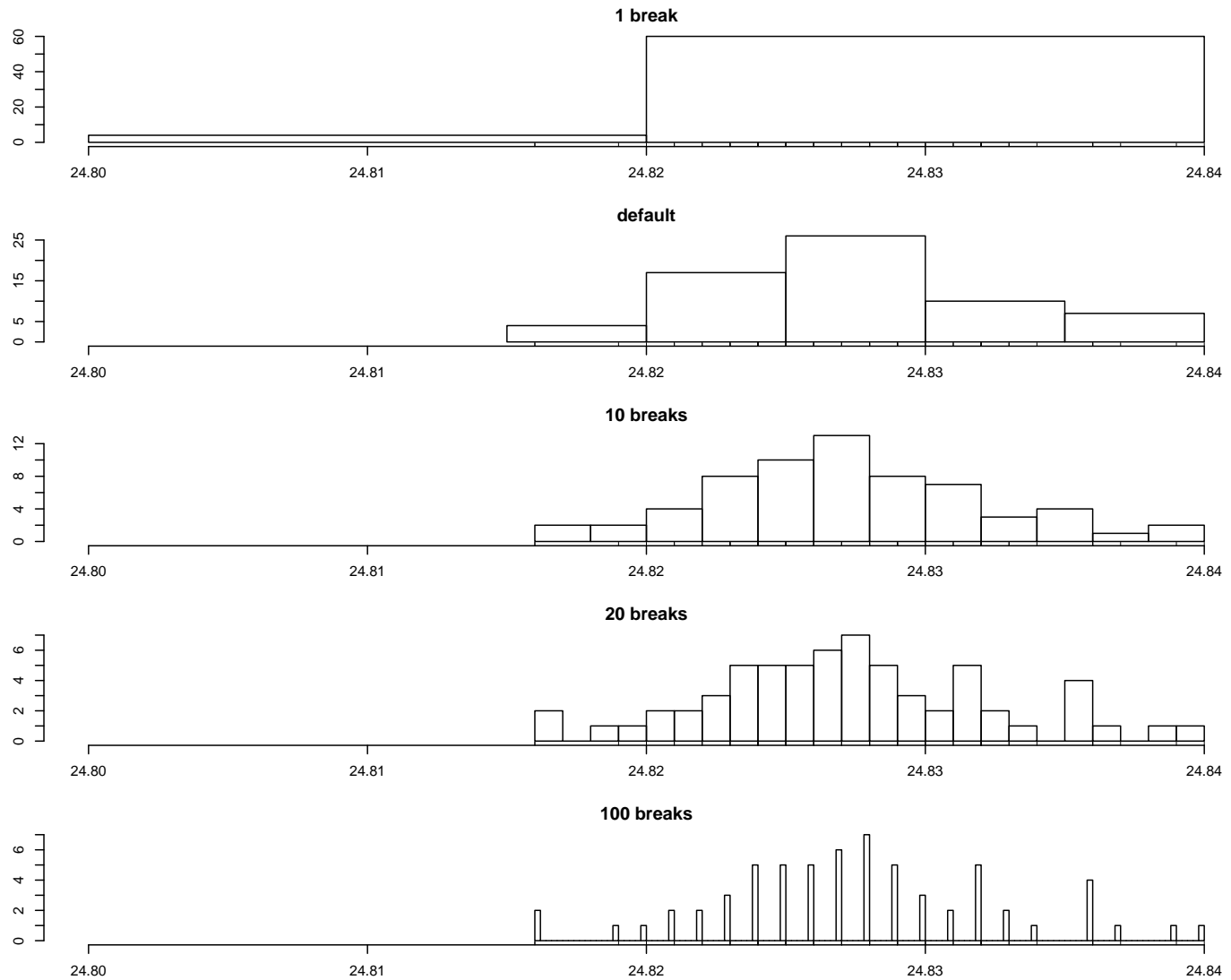
Consider the shape of the histogram for different numbers of bins.

```
#### Density estimation
# include time ranks 3 and above, that is, remove the lowest two values
time2 <- time[(rank(time) >= 3)]

old.par <- par(no.readonly = TRUE)
# make smaller margins
par(mfrow=c(5,1), mar=c(3,2,2,1), oma=c(1,1,1,1))

hist(time2, breaks=1      , main="1 break"      , xlim=c(24.80,24.84), xlab=""); rug(time2)
hist(time2,               main="default"      , xlim=c(24.80,24.84), xlab=""); rug(time2)
hist(time2, breaks=10     , main="10 breaks"    , xlim=c(24.80,24.84), xlab=""); rug(time2)
hist(time2, breaks=20     , main="20 breaks"    , xlim=c(24.80,24.84), xlab=""); rug(time2)
hist(time2, breaks=100    , main="100 breaks"   , xlim=c(24.80,24.84), xlab=""); rug(time2)

# restore par() settings
par(old.par)
```



Notice that we are starting to see more and more bins that include only a single observation (or multiple observations at the precision of measurement). Taken to its extreme, this type of exercise gives in some sense a “perfect” fit to the data but is useless as an estimator of shape.

On the other hand, it is obvious that a single bin would also be completely useless. So we try in some sense to find a middle ground between these two extremes: “Oversmoothing” by using only one bin and “undersmoothing” by using too many. This same paradigm occurs for density estimation, in which the amount of smoothing is determined by a quantity called the bandwidth. By default, R uses an optimal (in some sense) choice of bandwidth.

We’ve already used the `density()` function to provide a smooth curve to our histograms. So far, we’ve taken the default “bandwidth”. Let’s see what happens when we use different bandwidths.

```

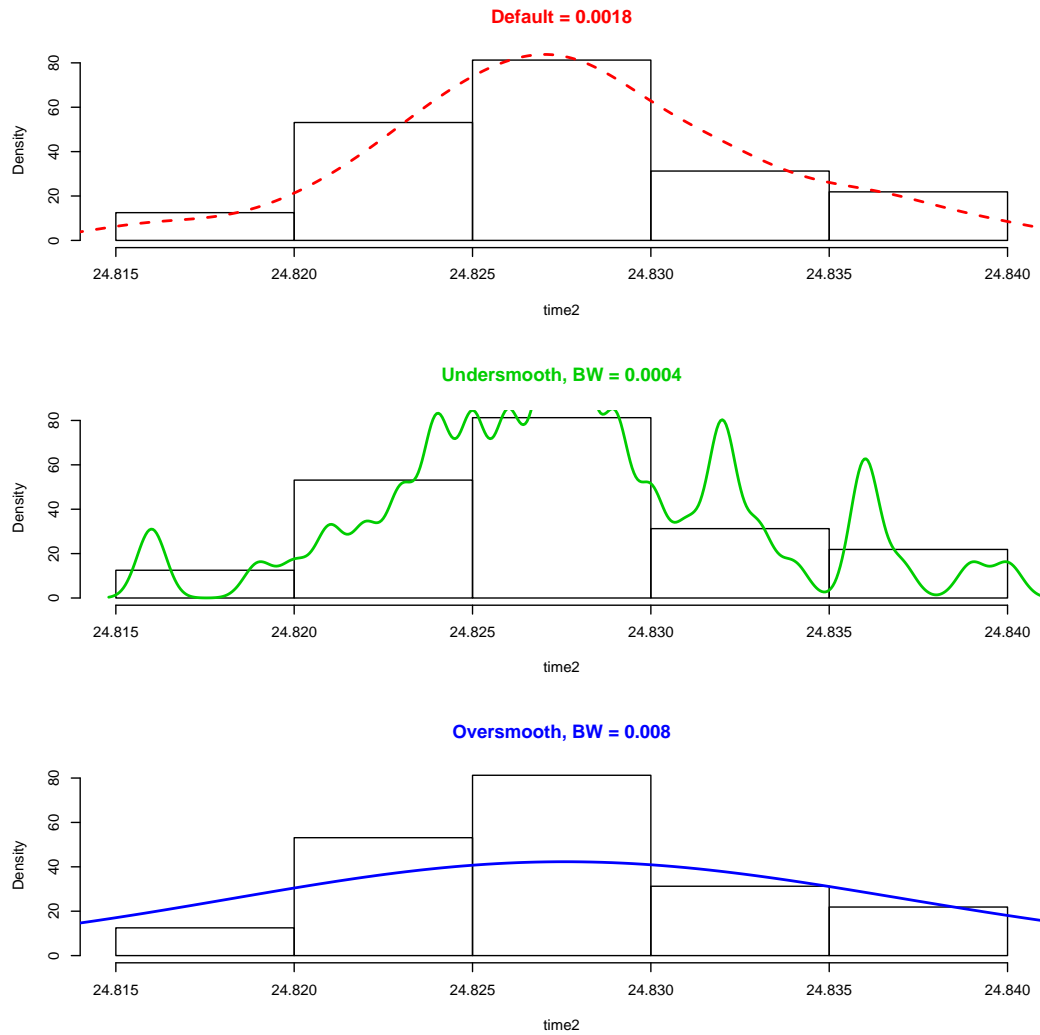
par(mfrow=c(3,1))

# prob=TRUE scales the y-axis like a density function, total area = 1
hist(time2, prob=TRUE, main="")
# apply a density function, store the result
den = density(time2)
# plot density line over histogram
lines(den, col=2, lty=2, lwd=2)
# extract the bandwidth (bw) from the density line
b = round(den$bw, 4)
title(main=paste("Default =", b), col.main=2)

# undersmooth
hist(time2, prob=TRUE, main="")
lines(density(time2, bw=0.0004), col=3, lwd=2)
text(17.5, .35, "", col=3, cex=1.4)
title(main=paste("Undersmooth, BW = 0.0004"), col.main=3)

# oversmooth
hist(time2, prob=TRUE, main="")
lines(density(time2, bw=0.008), col=4, lwd=2)
title(main=paste("Oversmooth, BW = 0.008"), col.main=4)

```



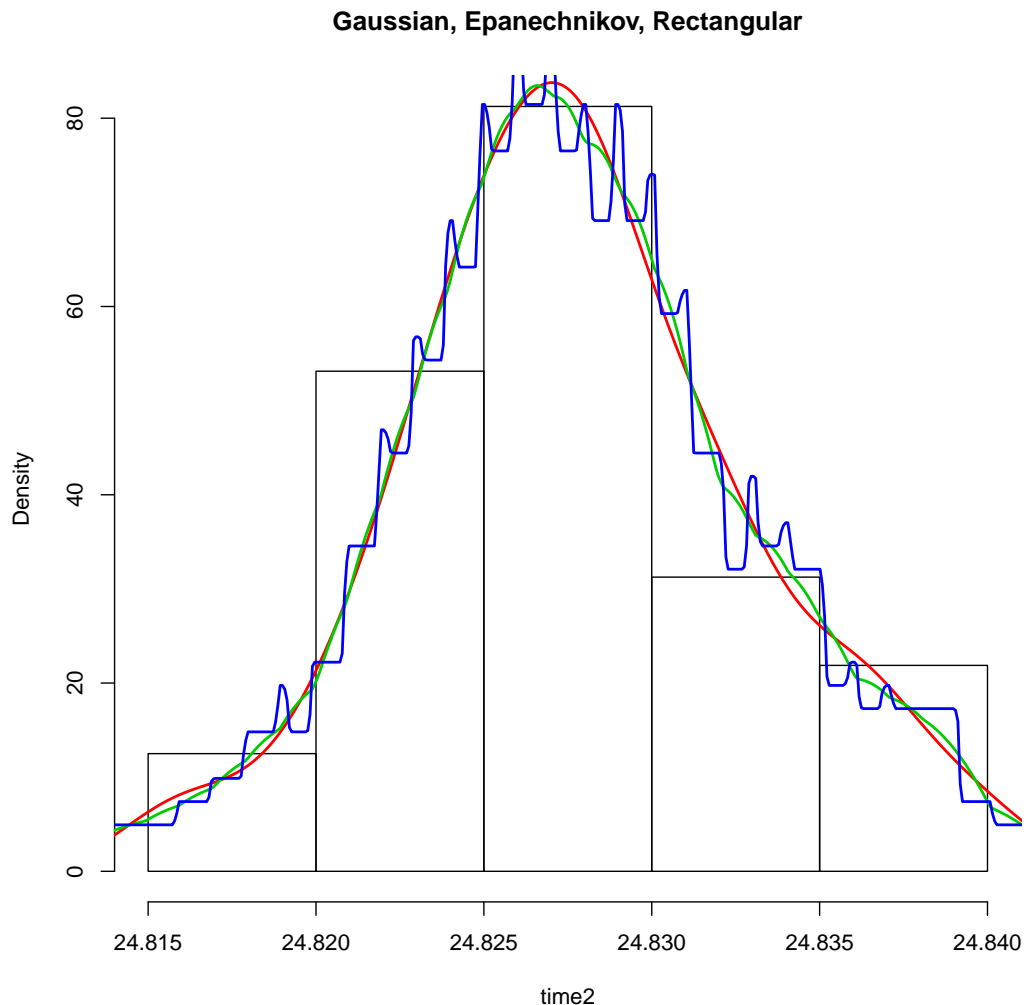
The other determining factor is the kernel, which is the shape each individual point takes before all the shapes are added up for a final density line. While the choice of bandwidth is very important, the choice of kernel is not. Choosing a kernel with hard edges (such as "rect") will result in jagged artifacts, so smoother kernels are often preferred.

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

hist(time2, prob=TRUE, main="")

# default kernel is Gaussian ("Normal")
lines(density(time2) , col=2, lty=1, lwd=2)
lines(density(time2, ker="epan"), col=3, lty=1, lwd=2)
lines(density(time2, ker="rect"), col=4, lty=1, lwd=2)
title(main="Gaussian, Epanechnikov, Rectangular")

# other kernels include: "triangular", "biweight", "cosine", "optcosine"
```



Chapter 7

Categorical Data Analysis

Learning objectives

After completing this topic, you should be able to:

select the appropriate statistical method to compare summaries from categorical variables.

assess the assumptions of one-way and two-way tests of proportions and independence.

decide whether the proportions between populations are different, including in stratified and cross-sectional studies.

recommend action based on a hypothesis test.

Achieving these goals contributes to mastery in these course learning outcomes:

1. organize knowledge.
5. define parameters of interest and hypotheses in words and notation.
6. summarize data visually, numerically, and descriptively.
8. use statistical software.
12. make evidence-based decisions.

7.1 Categorical data

When the response variable is categorical, the interesting questions are often about the probability of one possible outcome versus another, and whether

these probabilities depend on other variables (continuous or categorical).

Example: Titanic The sinking of the Titanic is a famous event, and new books are still being published about it. Many well-known facts — from the proportions of first-class passengers to the “women and children first” policy, and the fact that that policy was not entirely successful in saving the women and children in the third class — are reflected in the survival rates for various classes of passenger. The source provides a data set recording class, sex, age, and survival status for each person on board of the Titanic, and is based on data originally collected by the British Board of Trade¹.

```
# The Titanic dataset is a 4-dimensional table: Class, Sex, Age, Survived
library(datasets)
data(Titanic)

Titanic
## , , Age = Child, Survived = No
##
##       Sex
##   Class Male Female
##   1st     0     0
##   2nd     0     0
##   3rd    35    17
##   Crew    0     0
##
## , , Age = Adult, Survived = No
##
##       Sex
##   Class Male Female
##   1st   118      4
##   2nd   154     13
##   3rd   387     89
##   Crew  670      3
##
## , , Age = Child, Survived = Yes
##
##       Sex
##   Class Male Female
##   1st     5     1
##   2nd    11    13
##   3rd    13    14
##   Crew    0     0
##
```

¹British Board of Trade (1990), Report on the Loss of the “Titanic” (S.S.). British Board of Trade Inquiry Report (reprint). Gloucester, UK: Allan Sutton Publishing. Note that there is not complete agreement among primary sources as to the exact numbers on board, rescued, or lost.

```
## , , Age = Adult, Survived = Yes
##
##      Sex
## Class  Male Female
##   1st    57    140
##   2nd    14     80
##   3rd    75     76
##  Crew   192     20

# reshape into long data.frame
library(reshape2)
df.titanic <- melt(Titanic, value.name = "Freq")
df.titanic

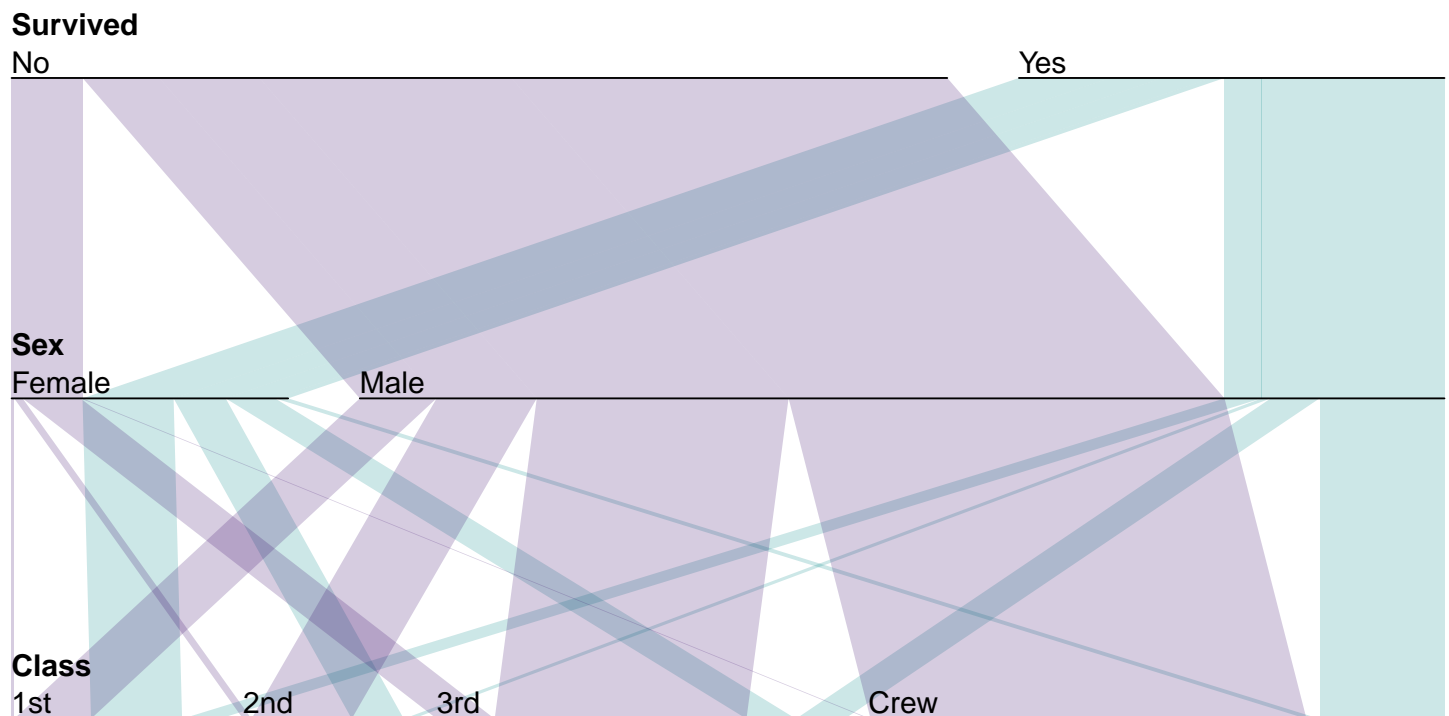
##   Class   Sex   Age Survived Freq
## 1  1st   Male Child     No     0
## 2  2nd   Male Child     No     0
## 3  3rd   Male Child     No    35
## 4  Crew   Male Child     No     0
## 5  1st Female Child     No     0
## 6  2nd Female Child     No     0
## 7  3rd Female Child     No    17
## 8  Crew Female Child     No     0
## 9  1st   Male Adult    No   118
## 10 2nd   Male Adult    No   154
## 11 3rd   Male Adult    No   387
## 12 Crew   Male Adult    No   670
## 13 1st Female Adult   No     4
## 14 2nd Female Adult   No    13
## 15 3rd Female Adult   No    89
## 16 Crew Female Adult   No     3
## 17 1st   Male Child   Yes     5
## 18 2nd   Male Child   Yes    11
## 19 3rd   Male Child   Yes    13
## 20 Crew   Male Child   Yes     0
## 21 1st Female Child  Yes     1
## 22 2nd Female Child  Yes    13
## 23 3rd Female Child  Yes    14
## 24 Crew Female Child  Yes     0
## 25 1st   Male Adult  Yes    57
## 26 2nd   Male Adult  Yes    14
## 27 3rd   Male Adult  Yes    75
## 28 Crew   Male Adult  Yes   192
## 29 1st Female Adult Yes   140
## 30 2nd Female Adult Yes    80
## 31 3rd Female Adult Yes    76
## 32 Crew Female Adult Yes    20

# Total number of people
sum(df.titanic$Freq)
## [1] 2201
```

```
# create colors based on survival
df.titanic$Color <- ifelse(df.titanic$Survived == "Yes", "#008888", "#330066")

# subset only the adults (since there were so few children)
df.titanic.adult <- subset(df.titanic, Age == "Adult")

# see R code on website for function parallelset()
# see help for with(), it allows temporary direct reference to columns in a data.frame
# otherwise, we'd need to specify df.titanic.adult$Survived, ...
with(df.titanic.adult
, parallelset(Survived, Sex, Class, freq = Freq, col = Color, alpha=0.2)
)
```



There are many questions that can be asked of this dataset. How likely were people to survive such a ship sinking in cold water? Is the survival proportion dependent on sex, class, or age, or a combination of these? How different is the survival proportions for 1st class females versus 3rd class males?

7.2 Single Proportion Problems

Assume that you are interested in estimating the proportion p of individuals in a population with a certain characteristic or attribute based on a random or representative sample of size n from the population. The **sample proportion** $\hat{p} = (\# \text{ with attribute in the sample})/n$ is the best guess for p based on the data.

This is the simplest **categorical data** problem. Each response falls into

one of two exclusive and exhaustive categories, called “success” and “failure”. Individuals with the attribute of interest are in the success category. The rest fall into the failure category. Knowledge of the **population proportion** p of successes characterizes the distribution across both categories because the population proportion of failures is $1 - p$.

As an aside, note that the probability that a randomly selected individual has the attribute of interest is the population proportion p with the attribute, so the terms population proportion and probability can be used interchangeably with random sampling.

7.2.1 A CI for p

A two-sided CI for p is a range of plausible values for the unknown population proportion p , based on the observed data. To compute a two-sided CI for p :

1. Specify the confidence level as the percent $100(1 - \alpha)\%$ and solve for the error rate α of the CI.
2. Compute $z_{\text{crit}} = z_{0.5\alpha}$ (i.e., area under the standard normal curve to the left and to the right of z_{crit} are $1 - 0.5\alpha$ and 0.5α , respectively).
 $\text{qnorm}(1-0.05/2)=1.96$.
3. The $100(1 - \alpha)\%$ CI for p has endpoints $L = \hat{p} - z_{\text{crit}}SE$ and $U = \hat{p} + z_{\text{crit}}SE$, respectively, where the “CI standard error” is

$$SE = \sqrt{\frac{\hat{p}(1 - \hat{p})}{n}}.$$

The CI is often written as $\hat{p} \pm z_{\text{crit}}SE$.

Reminder of CI interpretation. The CI is determined once the confidence level is specified and the data are collected. Prior to collecting data, the CI is unknown and can be viewed as random because it will depend on the actual sample selected. Different samples give different CIs. The “confidence” in, say, the 95% CI (which has a 0.05 or 5% error rate) can be interpreted as follows. If you repeatedly sample the population and construct 95% CIs for p , then 95% of the intervals will contain p , whereas 5% (the error rate) will not. The CI you get from your data either covers p , or it does not.

The length of the CI

$$U - L = 2z_{\text{crit}}SE$$

depends on the accuracy of the estimate \hat{p} , as measured by the standard error SE . For a given \hat{p} , this standard error decreases as the sample size n increases, yielding a narrower CI. For a fixed sample size, this standard error is maximized at $\hat{p} = 0.5$, and decreases as \hat{p} moves towards either 0 or 1. In essence, sample proportions near 0 or 1 give narrower CIs for p . However, the normal approximation used in the CI construction is less reliable for extreme values of \hat{p} .

■ CLICKER Qs — CI for proportions STT.08.01.010 ■

Example: Tamper resistant packaging The 1983 Tylenol poisoning episode highlighted the desirability of using tamper-resistant packaging. The article “Tamper Resistant Packaging: Is it Really?” (Packaging Engineering, June 1983) reported the results of a survey on consumer attitudes towards tamper-resistant packaging. A sample of 270 consumers was asked the question: “Would you be willing to pay extra for tamper resistant packaging?” The number of yes respondents was 189. Construct a 95% CI for the proportion p of all consumers who were willing in 1983 to pay extra for such packaging.

Here $n = 270$ and $\hat{p} = 189/270 = 0.700$. The critical value for a 95% CI for p is $z_{0.025} = 1.96$. The CI standard error is given by

$$SE = \sqrt{\frac{0.7 \times 0.3}{270}} = 0.028,$$

so $z_{\text{crit}}SE = 1.96 \times 0.028 = 0.055$. The 95% CI for p is 0.700 ± 0.055 . You are 95% confident that the proportion of consumers willing to pay extra for better packaging is between 0.645 and 0.755. (Willing to pay how much extra?)

Appropriateness of the CI

The standard CI is based on a **large-sample** standard normal approximation to

$$z = \frac{\hat{p} - p}{SE}.$$

A simple rule of thumb requires $n\hat{p} \geq 5$ and $n(1 - \hat{p}) \geq 5$ for the method to be suitable. Given that $n\hat{p}$ and $n(1 - \hat{p})$ are the observed numbers of successes and failures, you should have at least 5 of each to apply the large-sample CI.

In the packaging example, $n\hat{p} = 270 \times (0.700) = 189$ (the number who support the new packaging) and $n(1 - \hat{p}) = 270 \times (0.300) = 81$ (the number who oppose) both exceed 5. The normal approximation is appropriate here.

7.2.2 Hypothesis Tests on Proportions

The following example is typical of questions that can be answered using a hypothesis test for a population proportion.

Example Environmental problems associated with leaded gasolines are well-known. Many motorists have tampered with the emission control devices on their cars to save money by purchasing leaded rather than unleaded gasoline. A *Los Angeles Times* article on March 17, 1984 reported that 15% of all California motorists have engaged in emissions tampering. A random sample of 200 cars from L.A. county was obtained, and the emissions devices on 21 are found to be tampered with. Does this suggest that the proportion of cars in L.A. county with tampered devices differs from the statewide proportion?

Two-Sided Hypothesis Test for p

Suppose you are interested in whether the population proportion p is equal to a prespecified value, say p_0 . This question can be formulated as a two-sided test. To carry out the test:

1. Define the null hypothesis $H_0 : p = p_0$ and alternative hypothesis $H_A : p \neq p_0$.
2. Choose the size or significance level of the test, denoted by α .

3. Using the standard normal probability table, find the critical value z_{crit} such that the areas under the normal curve to the left and right of z_{crit} are $1 - 0.5\alpha$ and 0.5α , respectively. That is, $z_{\text{crit}} = z_{0.5\alpha}$.
4. Compute the test statistic (often to be labelled z_{obs})

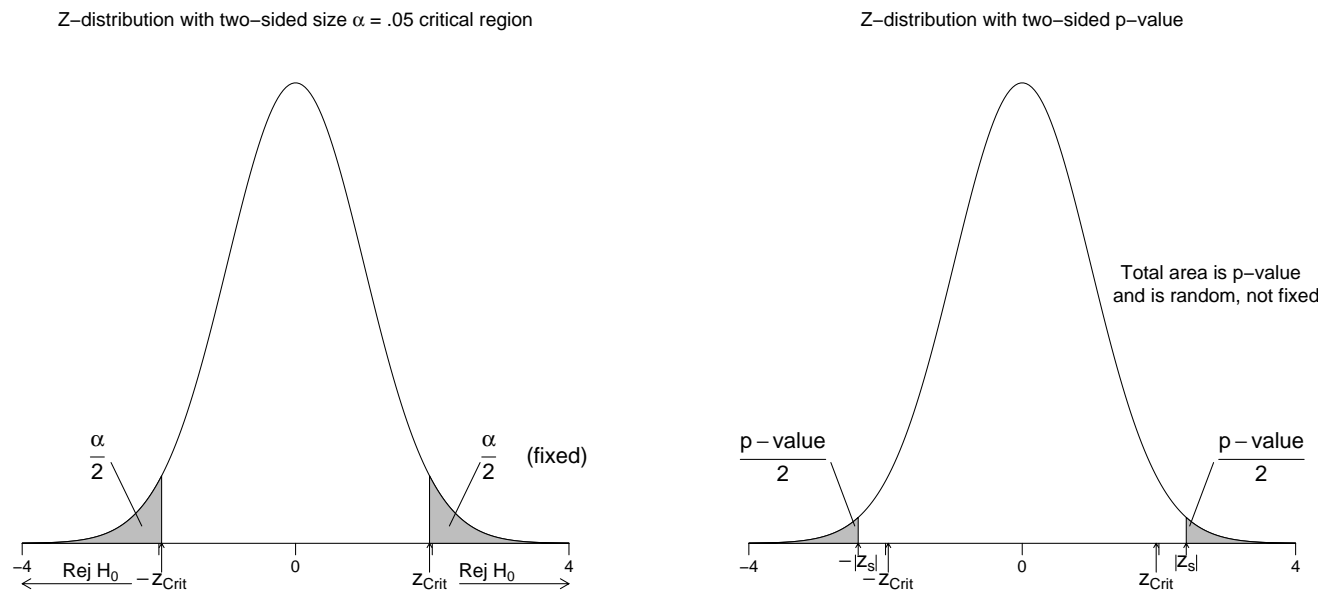
$$z_s = \frac{\hat{p} - p_0}{SE},$$

where the “test standard error” (based on the hypothesized value) is

$$SE = \sqrt{\frac{p_0(1 - p_0)}{n}}.$$

5. Reject H_0 in favor of H_A if $|z_{\text{obs}}| \geq z_{\text{crit}}$. Otherwise, do not reject H_0 .

The rejection rule is easily understood visually. The area under the normal curve outside $\pm z_{\text{crit}}$ is the size α of the test. One-half of α is the area in each tail. You reject H_0 in favor of H_A if the test statistic exceeds $\pm z_{\text{crit}}$. This occurs when \hat{p} is significantly different from p_0 , as measured by the standardized distance z_{obs} between \hat{p} and p_0 .



7.2.3 The p-value for a two-sided test

To compute the p-value (not to be confused with the value of the proportion p) for a two-sided test:

1. Compute the test statistic $z_s = z_{\text{obs}}$.
2. Evaluate the area under the normal probability curve outside $\pm|z_s|$.

Recall that the null hypothesis for a size α test is rejected if and only if the p-value is less than or equal to α .

Example: Emissions data Each car in the target population (L.A. county) either has been tampered with (a success) or has not been tampered with (a failure). Let p = the proportion of cars in L.A. county with tampered emissions control devices. You want to test $H_0 : p = 0.15$ against $H_A : p \neq 0.15$ (here $p_0 = 0.15$). The critical value for a two-sided test of size $\alpha = 0.05$ is $z_{\text{crit}} = 1.96$.

The data are a sample of $n = 200$ cars. The sample proportion of cars that have been tampered with is $\hat{p} = 21/200 = 0.105$. The test statistic is

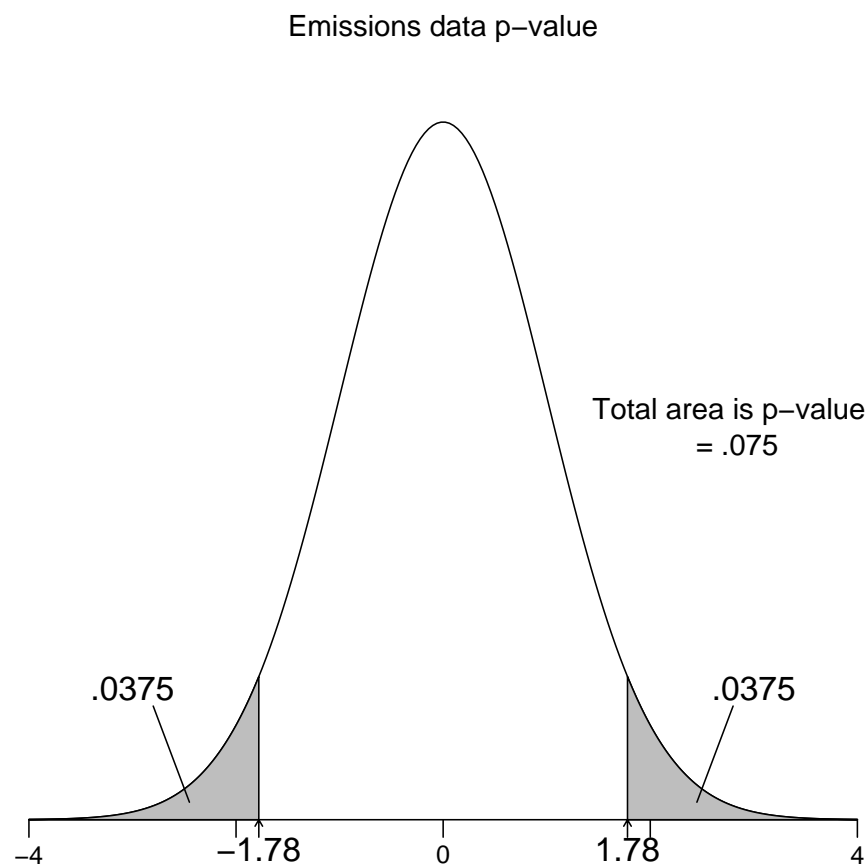
$$z_s = \frac{0.105 - 0.15}{0.02525} = -1.78,$$

where the test standard error satisfies

$$SE = \sqrt{\frac{0.15 \times 0.85}{200}} = 0.02525.$$

Given that $|z_s| = 1.78 < 1.96$, you have insufficient evidence to reject H_0 at the 5% level. That is, you have insufficient evidence to conclude that the proportion of cars in L.A. county that have been tampered with differs from the statewide proportion.

This decision is reinforced by the p-value calculation. The p-value is the area under the standard normal curve outside ± 1.78 . This is $2 \times 0.0375 = 0.075$, which exceeds the test size of 0.05.



Remark The SE used in the test and CI are different. This implies that a hypothesis test and CI could potentially lead to different decisions. That is, a 95% CI for a population proportion might cover p_0 when the p-value for testing $H_0 : p = p_0$ is smaller than 0.05. This will happen, typically, only in cases where the decision is “borderline.”

7.2.4 Appropriateness of Test

The z -test is based on a large-sample normal approximation, which works better for a given sample size when p_0 is closer to 0.5. The sample size needed for an accurate approximation increases dramatically the closer p_0 gets to 0 or to 1. A simple rule of thumb is that the test is appropriate when (the expected number of successes) $np_0 \geq 5$ and (the expected number of failures) $n(1 - p_0) \geq 5$.

In the emissions example, $np_0 = 200 \times (0.15) = 30$ and $n(1 - p_0) = 200 \times (0.85) = 170$ exceed 5, so the normal approximation is appropriate.

7.2.5 R Implementation

```
##### Single Proportion Problems
# Approximate normal test for proportion, without Yates' continuity correction
prop.test(21, 200, p = 0.15, correct = FALSE)

##
## 1-sample proportions test without continuity correction
##
## data: 21 out of 200, null probability 0.15
## X-squared = 3.176, df = 1, p-value = 0.07471
## alternative hypothesis: true p is not equal to 0.15
## 95 percent confidence interval:
## 0.06971 0.15518
## sample estimates:
## p
## 0.105

# Approximate normal test for proportion, with Yates' continuity correction
#prop.test(21, 200, p = 0.15)
```

■ CLICKER Qs — Parachute null hypothesis STT.08.01.040 ■

■ CLICKER Qs — Parachute conclusion STT.08.01.050 ■

■ CLICKER Qs — Parachute p-value STT.08.01.060 ■

7.2.6 One-Sided Tests and One-Sided Confidence Bounds

The mechanics of tests on proportions are similar to tests on means, except we use a different test statistic and a different probability distribution for critical values. This applies to one-sided and two-sided procedures. The example below illustrates a one-sided test and bound.

Example: brain hemispheres An article in the April 6, 1983 edition of *The Los Angeles Times* reported on a study of 53 learning-impaired youngsters at the Massachusetts General Hospital. The right side of the brain was found to be larger than the left side in 22 of the children. The proportion of the general population with brains having larger right sides is known to be 0.25. Does the data provide strong evidence for concluding, as the article claims, that the proportion of learning impaired youngsters with brains having larger right sides exceeds the proportion in the general population?

I will answer this question by computing a p -value for a one-sided test. Let p be the population proportion of learning disabled children with brains having larger right sides. I am interested in testing $H_0 : p = 0.25$ against $H_A : p > 0.25$ (here $p_0 = 0.25$).

The proportion of children sampled with brains having larger right sides is $\hat{p} = 22/53 = 0.415$. The test statistic is

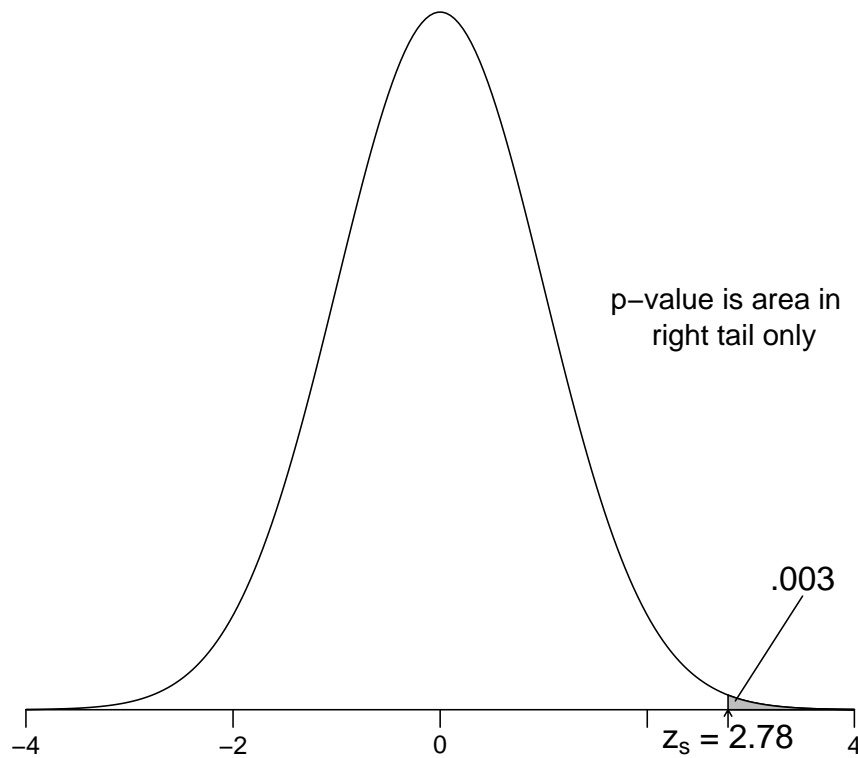
$$z_s = \frac{0.415 - 0.25}{0.0595} = 2.78,$$

where the test standard error satisfies

$$SE = \sqrt{\frac{0.25 \times 0.75}{53}} = 0.0595.$$

The p -value for an upper one-sided test is the area under the standard normal curve to the right of 2.78, which is approximately .003; see the picture below. I would reject H_0 in favor of H_A using any of the standard test levels, say 0.05 or 0.01. The newspaper's claim is reasonable.

Right brain upper one-sided p-value



A sensible next step in the analysis would be to compute a **lower confidence bound** $\hat{p} - z_{\text{crit}}SE$ for p . For illustration, consider a 95% bound. The CI standard error is

$$SE = \sqrt{\frac{\hat{p}(1 - \hat{p})}{n}} = \sqrt{\frac{0.415 \times 0.585}{53}} = 0.0677.$$

The critical value for a one-sided 5% test is $z_{\text{crit}} = 1.645$, so a lower 95% bound on p is $0.415 - 1.645 \times 0.0677 = 0.304$. I am 95% confident that the population proportion of learning disabled children with brains having larger right sides is at least 0.304. Values of p smaller than 0.304 are not supported by the data.

You should verify that the sample size is sufficiently large to use the approximate methods in this example.

```
##### Example: brain hemispheres
# Approximate normal test for proportion, without Yates' continuity correction
prop.test(22, 53, p = 0.25, alternative = "greater", correct = FALSE)
##
```

```
## 1-sample proportions test without continuity correction
##
## data: 22 out of 53, null probability 0.25
## X-squared = 7.704, df = 1, p-value = 0.002754
## alternative hypothesis: true p is greater than 0.25
## 95 percent confidence interval:
## 0.3105 1.0000
## sample estimates:
## p
## 0.4151
```

7.2.7 Small Sample Procedures

Large sample tests and CIs for p should be interpreted with caution in small sized samples because the true error rate usually exceeds the assumed (nominal) value. For example, an assumed 95% CI, with a nominal error rate of 5%, may be only an 80% CI, with a 20% error rate. The large-sample CIs are usually overly optimistic (i.e., too narrow) when the sample size is too small to use the normal approximation.

Alan Agresti suggests the following method for a 95% CI. The standard method computes the sample proportion as $\hat{p} = x/n$ where x is the number of successes in the sample and n is the sample size. Agresti suggested using the estimated proportion $\tilde{p} = (x + 2)/(n + 4)$ with the standard error

$$SE = \sqrt{\frac{\tilde{p}(1 - \tilde{p})}{n + 4}},$$

in the “usual 95% interval” formula: $\tilde{p} \pm 1.96SE$. This appears odd, but amounts to adding two successes and two failures to the observed data, and then computing the standard CI.

This adjustment has little effect when n is large and \hat{p} is not near either 0 or 1, as in the Tylenol example.

Example: swimming segregation This example is based on a case heard before the U.S. Supreme Court. A racially segregated swimming club was ordered to admit minority members. However, it is unclear whether the club has been following the spirit of the mandate. Historically, 85% of the white

applicants were approved. Since the mandate, only 1 of 6 minority applicants has been approved. Is there evidence of continued discrimination?

I examine this issue by constructing a CI and a test for the probability p (or population proportion) that a minority applicant is approved. Before examining the question of primary interest, let me show that the two approximate CIs are very different, due to the small sample size. One minority applicant ($x = 1$) was approved out of $n = 6$ candidates, giving $\hat{p} = 1/6 = 0.167$.

A 95% large-sample CI for p is $(-0.14, 0.46)$. Since a negative proportion is not possible, the CI should be reported as $(0.00, 0.46)$. Agresti's 95% CI (based on 3 successes and 7 failures) is $(0.02, 0.58)$. The big difference between the two intervals coupled with the negative lower endpoint on the standard CI suggests that the normal approximation used with the standard method is inappropriate. This view is reinforced by the rule-of-thumb calculation for using the standard interval. Agresti's CI is wider, which is consistent with my comment that the standard CI is too narrow in small samples. As a comparison, the exact 95% CI is $(0.004, 0.64)$, which agrees more closely with Agresti's interval.

I should emphasize that the exact CI is best to use, but is not available in all statistical packages, so methods based on approximations may be required, and if so, then Agresti's method is clearly better than the standard normal approximation in small sized samples.

Recall that the results of the asymptotic 95% CIs may disagree with the hypothesis test results. Exact methods will not contradict each other this way (neither do these asymptotic methods, usually).

```
#### Example: swimming segregation
## The prop.test() does an additional adjustment, so does not match precisely
##   the results in the above paragraphs

# Approximate normal test for proportion, without Yates' continuity correction
prop.test(1, 6, p = 0.85, correct = FALSE)$conf.int

## Warning: Chi-squared approximation may be incorrect
## [1] 0.03005 0.56350
## attr(),"conf.level")
## [1] 0.95

# Agresti's method
prop.test(1+2, 6+4, p = 0.85, correct = FALSE)$conf.int

## Warning: Chi-squared approximation may be incorrect
## [1] 0.1078 0.6032
## attr(),"conf.level")
```

```
## [1] 0.95
# Exact binomial test for proportion
binom.test(1, 6, p = 0.85)$conf.int
## [1] 0.004211 0.641235
## attr(,"conf.level")
## [1] 0.95
```

Returning to the problem, you might check for discrimination by testing $H_0 : p = 0.85$ against $H_A : p < 0.85$ using an **exact** test. The exact test p-value is 0.000 to three decimal places, and an exact upper bound for p is 0.582. What does this suggest to you?

```
# Exact binomial test for proportion
binom.test(1, 6, alternative = "less", p = 0.85)
##
##  Exact binomial test
##
## data: 1 and 6
## number of successes = 1, number of trials = 6, p-value =
## 0.0003987
## alternative hypothesis: true probability of success is less than 0.85
## 95 percent confidence interval:
##  0.0000 0.5818
## sample estimates:
## probability of success
##                      0.1667
```

7.3 Analyzing Raw Data

In most studies, your data will be stored in a spreadsheet with one observation per case or individual. For example, the data below give the individual responses to the applicants of the swim club.

```
#### Example: swimming segregation, raw data
## read.table options
# sep = default is any white space, but our strings contain a space,
#       so I changed this to a comma
# header = there are no column headers
# stringsAsFactors = default converts strings to factors, but I want them
#                   to just be the plain character text
swim <- read.table(text="
not approved
not approved
not approved
approved
not approved
not approved
", sep = ", ", header=FALSE, stringsAsFactors=FALSE)
```

```
# name the column
names(swim) <- c("application")
# show the structure of the data.frame
str(swim)

## 'data.frame': 6 obs. of 1 variable:
## $ application: chr "not approved" "not approved" "not approved" "approved" ...
# display the data.frame
swim

##      application
## 1    not approved
## 2    not approved
## 3    not approved
## 4     approved
## 5    not approved
## 6    not approved
```

The data were entered as alphabetic strings. We can use `table()` to count frequencies of categorical variables.

```
# count the frequency of each categorical variable
table(swim)

## swim
##     approved not approved
##           1            5
```

You can compute a CI and test for a proportion using raw data, provided the data column includes only two distinct values. The levels can be numerical or alphanumeric.

```
# use the counts from table() for input in binom.test()
# the help for binom.test() says x can be a vector of length 2
# giving the numbers of successes and failures, respectively
# that's exactly what table(swim) gave us
binom.test(table(swim), p = 0.85, alternative = "less")

##
## Exact binomial test
##
## data: table(swim)
## number of successes = 1, number of trials = 6, p-value =
## 0.0003987
## alternative hypothesis: true probability of success is less than 0.85
## 95 percent confidence interval:
##  0.0000 0.5818
## sample estimates:
## probability of success
##                  0.1667
```

It is possible that the order (alphabetically) is the wrong order, failures and successes, in which case we'd need to reorder the input to `binom.test()`.

In Chapter 6 we looked at the binomial distribution to obtain an exact Sign

Test confidence interval for the median. Examine the following to see where the exact p-value for this test comes from.

```
n <- 6
x <- 0:n
p0 <- 0.85
bincdf <- pbinom(x, n, p0)
cdf <- data.frame(x, bincdf)
cdf

##   x     bincdf
## 1 0 1.139e-05
## 2 1 3.987e-04
## 3 2 5.885e-03
## 4 3 4.734e-02
## 5 4 2.235e-01
## 6 5 6.229e-01
## 7 6 1.000e+00
```

■ CLICKER Qs — Excess successes STT.05.01.030 ■

7.4 Goodness-of-Fit Tests (Multinomial)

Example: jury pool The following data set was used as evidence in a court case. The data represent a sample of 1336 individuals from the jury pool of a large municipal court district for the years 1975–1977. The fairness of the representation of various age groups on juries was being contested. The strategy for doing this was to challenge the representativeness of the pool of individuals from which the juries are drawn. This was done by comparing the age group distribution within the jury pool against the age distribution in the district as a whole, which was available from census figures.

Age group (yrs)	Obs. Counts	Obs. Prop.	Census Prop.
18-19	23	0.017	0.061
20-24	96	0.072	0.150
25-29	134	0.100	0.135
30-39	293	0.219	0.217
40-49	297	0.222	0.153
50-64	380	0.284	0.182
65-99	113	0.085	0.102
Total:	1336	1.000	1.000

A statistical question here is whether the jury pool population proportions are equal to the census proportions across the age categories. This comparison can be formulated as a **goodness-of-fit test**, which generalizes the large-sample test on a single proportion to a categorical variable (here age) with $r > 2$ levels. For $r = 2$ categories, the goodness-of-fit test and large-sample test on a single proportion are identical. Although this problem compares two populations, only one sample is involved because the census data is a population summary!

In general, suppose each individual in a population is categorized into one and only one of r levels or categories. Let p_1, p_2, \dots, p_r , be the population proportions in the r categories, where each $p_i \geq 0$ and $p_1 + p_2 + \dots + p_r = 1$. The hypotheses of interest in a goodness-of-fit problem are $H_0 : p_1 = p_{01}, p_2 = p_{02}, \dots, p_r = p_{0r}$ and $H_A : \text{not } H_0$, where $p_{01}, p_{02}, \dots, p_{0r}$ are given category proportions.

The plausibility of H_0 is evaluated by comparing the hypothesized category proportions to estimated (i.e., observed) category proportions $\hat{p}_1, \hat{p}_2, \dots, \hat{p}_r$ from a random or representative sample of n individuals selected from the population. The discrepancy between the hypothesized and observed proportions is measured by the Pearson chi-squared statistic:

$$\chi_s^2 = \sum_{i=1}^r \frac{(O_i - E_i)^2}{E_i},$$

where O_i is the **observed** number in the sample that fall into the i^{th} category ($O_i = n\hat{p}_i$), and $E_i = np_{0i}$ is the number of individuals **expected** to be in the

i^{th} category when H_0 is true.

The Pearson statistic can also be computed as the sum of the squared residuals:

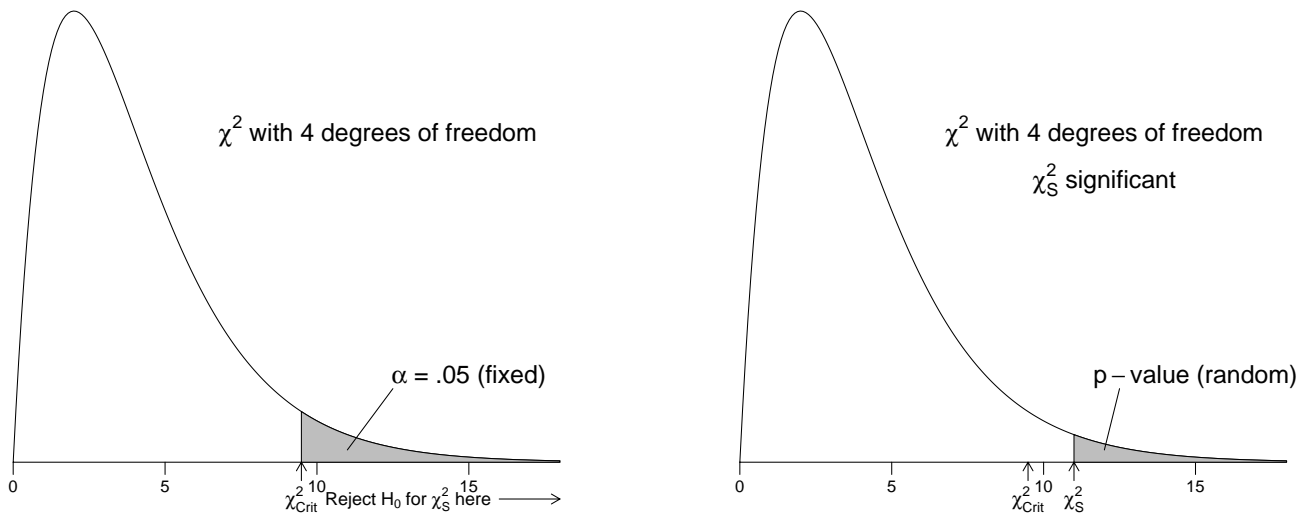
$$\chi_s^2 = \sum_{i=1}^r Z_i^2,$$

where $Z_i = (O_i - E_i)/\sqrt{E_i}$, or in terms of the observed and hypothesized category proportions

$$\chi_s^2 = n \sum_{i=1}^r \frac{(\hat{p}_i - p_{0i})^2}{p_{0i}}.$$

The Pearson statistic χ_s^2 is “small” when all of the observed counts (proportions) are close to the expected counts (proportions). The Pearson χ^2 is “large” when one or more observed counts (proportions) differs noticeably from what is expected when H_0 is true. Put another way, large values of χ_s^2 suggest that H_0 is false.

The critical value χ_{crit} for the test is obtained from a chi-squared probability table with $r - 1$ degrees of freedom. The picture below shows the form of the rejection region. For example, if $r = 5$ and $\alpha = 0.05$, then you reject H_0 when $\chi_s^2 \geq \chi_{\text{crit}}^2 = 9.49$ (`qchisq(0.95, 5-1)`). The p-value for the test is the area under the chi-squared curve with $df = r - 1$ to the right of the observed χ_s^2 value.



Example: jury pool Let p_{18} be the proportion in the jury pool population between ages 18 and 19. Define $p_{20}, p_{25}, p_{30}, p_{40}, p_{50}$, and p_{65} analogously. You are interested in testing that the true jury proportions equal the census proportions, $H_0 : p_{18} = 0.061, p_{20} = 0.150, p_{25} = 0.135, p_{30} = 0.217, p_{40} = 0.153, p_{50} = 0.182$, and $p_{65} = 0.102$ against $H_A : \text{not } H_0$, using the sample of 1336 from the jury pool.

The observed counts, the expected counts, and the category residuals are given in the table below. For example, $E_{18} = 1336 \times (0.061) = 81.5$ and $Z_{18} = (23 - 81.5)/\sqrt{81.5} = -6.48$ in the 18-19 year category.

The Pearson statistic is

$$\chi_s^2 = (-6.48)^2 + (-7.38)^2 + (-3.45)^2 + 0.18^2 + 6.48^2 + 8.78^2 + (-1.99)^2 = 231.26$$

on $r - 1 = 7 - 1 = 6$ degrees of freedom. Here $\chi_{\text{crit}}^2 = 12.59$ at $\alpha = 0.05$. The p-value for the goodness-of-fit test is less than 0.001, which suggests that H_0 is false.

Age group (yrs)	Obs. Counts	Exp. Counts	Residual
18-19	23	81.5	-6.48
20-24	96	200.4	-7.38
25-29	134	180.4	-3.45
30-39	293	289.9	0.18
40-49	297	204.4	6.48
50-64	380	243.2	8.78
65-99	113	136.3	-1.99

7.4.1 Adequacy of the Goodness-of-Fit Test

The chi-squared goodness-of-fit test is a large-sample test. A conservative rule of thumb is that the test is suitable when each **expected** count is at least five. This holds in the jury pool example. There is no widely available alternative method for testing goodness-of-fit with smaller sample sizes. There is evidence, however, that the chi-squared test is **slightly conservative** (the p-values are too large, on average) when the expected counts are smaller. Some statisticians recommend that the chi-squared approximation be used when the minimum expected count is at least one, provided the expected counts are not too variable.

7.4.2 R Implementation

```
#### Example: jury pool
jury <- read.table(text="
Age      Count    CensusProp
18-19     23      0.061
20-24     96      0.150
25-29    134      0.135
30-39    293      0.217
40-49    297      0.153
50-64    380      0.182
65-99    113      0.102
", header=TRUE)

# show the structure of the data.frame
str(jury)
## 'data.frame': 7 obs. of 3 variables:
## $ Age       : Factor w/ 7 levels "18-19","20-24",...: 1 2 3 4 5 6 7
## $ Count     : int  23 96 134 293 297 380 113
## $ CensusProp: num  0.061 0.15 0.135 0.217 0.153 0.182 0.102

# display the data.frame
jury
##      Age Count CensusProp
## 1 18-19   23     0.061
## 2 20-24   96     0.150
## 3 25-29  134     0.135
## 4 30-39  293     0.217
## 5 40-49  297     0.153
## 6 50-64  380     0.182
## 7 65-99  113     0.102

# calculate chi-square goodness-of-fit test
x.summary <- chisq.test(jury$Count, correct = FALSE, p = jury$CensusProp)
# print result of test
x.summary
##
## Chi-squared test for given probabilities
##
## data: jury$Count
## X-squared = 231.3, df = 6, p-value < 2.2e-16

# use output in x.summary and create table
x.table <- data.frame(age = jury$Age
                      , obs = x.summary$observed
                      , exp = x.summary$expected
                      , res = x.summary$residuals
                      , chisq = x.summary$residuals^2
                      , stdres = x.summary$stdres)
x.table
##    age obs    exp    res    chisq    stdres
```

```
## 1 18-19 23 81.5 -6.4797 41.98712 -6.687
## 2 20-24 96 200.4 -7.3748 54.38802 -7.999
## 3 25-29 134 180.4 -3.4520 11.91644 -3.712
## 4 30-39 293 289.9 0.1814 0.03289 0.205
## 5 40-49 297 204.4 6.4763 41.94199 7.037
## 6 50-64 380 243.2 8.7761 77.01921 9.703
## 7 65-99 113 136.3 -1.9936 3.97430 -2.104
```

Plot observed vs expected values to help identify age groups that deviate the most. Plot contribution to chi-square values to help identify age groups that deviate the most. The term “Contribution to Chi-Square” (`chisq`) refers to the values of $\frac{(O-E)^2}{E}$ for each category. χ^2_s is the sum of those contributions.

```
library(reshape2)
x.table.obsexp <- melt(x.table,
  # id.vars: ID variables
  # all variables to keep but not split apart on
  id.vars = c("age"),
  # measure.vars: The source columns
  # (if unspecified then all other variables are measure.vars)
  measure.vars = c("obs", "exp"),
  # variable.name: Name of the destination column identifying each
  # original column that the measurement came from
  variable.name = "stat",
  # value.name: column name for values in table
  value.name = "value"
)

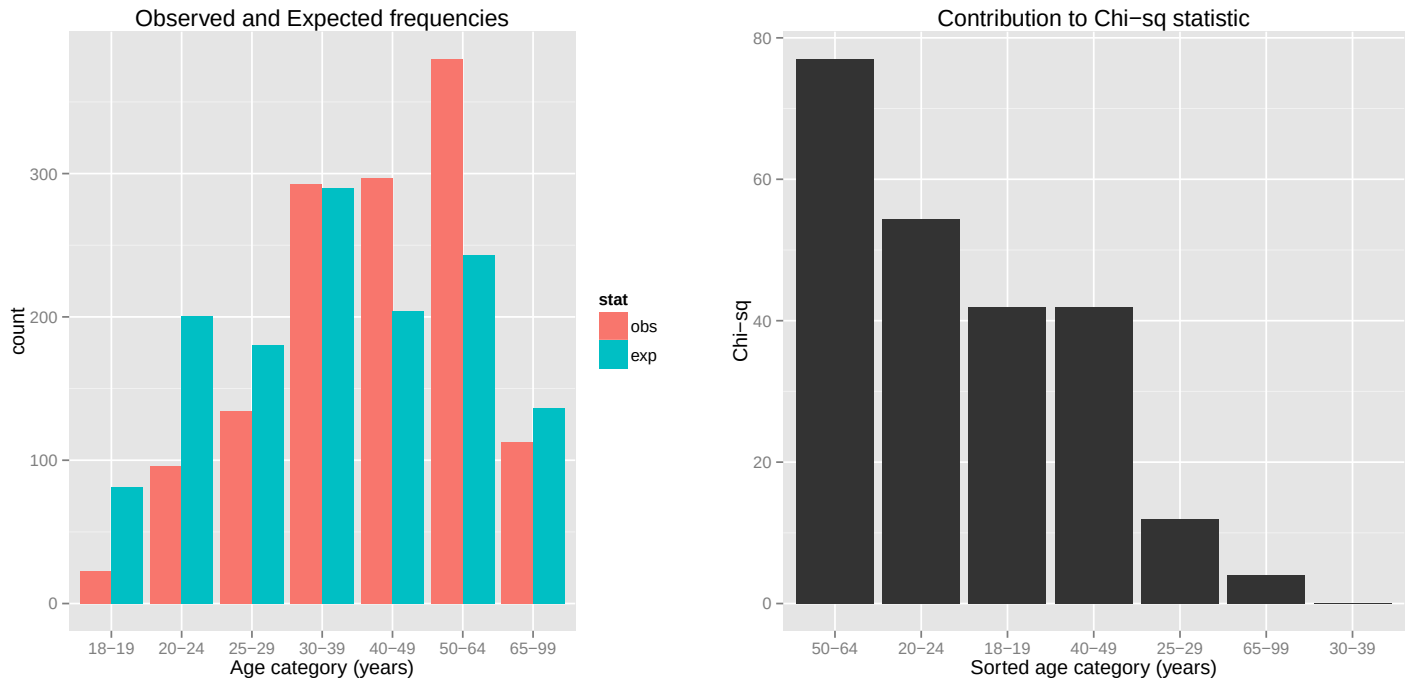
# naming variables manually, the variable.name and value.name not working 11/2012
names(x.table.obsexp) <- c("age", "stat", "value")

# Observed vs Expected counts
library(ggplot2)
p <- ggplot(x.table.obsexp, aes(x = age, fill = stat, weight=value))
p <- p + geom_bar(position="dodge")
p <- p + labs(title = "Observed and Expected frequencies")
p <- p + xlab("Age category (years)")
print(p)

# Contribution to chi-sq
# pull out only the age and chisq columns
x.table.chisq <- x.table[, c("age", "chisq")]
# reorder the age categories to be descending relative to the chisq statistic
x.table.chisq$age <- with(x.table, reorder(age, -chisq))

p <- ggplot(x.table.chisq, aes(x = age, weight = chisq))
p <- p + geom_bar()
p <- p + labs(title = "Contribution to Chi-sq statistic")
p <- p + xlab("Sorted age category (years)")
```

```
p <- p + ylab("Chi-sq")
print(p)
```



7.4.3 Multiple Comparisons in a Goodness-of-Fit Problem

The goodness-of-fit test suggests that at least one of the age category proportions for the jury pool population differs from the census figures. A reasonable next step in the analysis would be to **separately** test the seven hypotheses: $H_0 : p_{18} = 0.061$, $H_0 : p_{20} = 0.150$, $H_0 : p_{25} = 0.135$, $H_0 : p_{30} = 0.217$, $H_0 : p_{40} = 0.153$, $H_0 : p_{50} = 0.182$, and $H_0 : p_{65} = 0.102$ to see which age categories led to this conclusion.

A Bonferroni comparison with a Family Error Rate ≤ 0.05 will be considered for this multiple comparisons problem. The error rates for the seven individual tests are set to $\alpha = 0.05/7 = 0.0071$, which corresponds to 99.29% two-sided CIs for the individual jury pool proportions. The area under the standard normal curve to the right of 2.69 is $0.05/2/7 = 0.0036$, about one-half the error rate for the individual CIs, so the critical value for the CIs, or for the tests, is $z_{\text{crit}} \approx 2.69$. The next table gives individual 99.29% CIs based on the large sample approximation. You can get the individual CIs in R using the `binom.test()` or `prop.test()` function. For example, the CI for Age Group

18-19 is obtained by specifying 23 successes in 1336 trials.

Below I perform exact binomial tests of proportion for each of the seven age categories at the Bonferroni-adjusted significance level. I save the p-values and confidence intervals in a table along with the observed and census proportions in order to display the table below.

```
b.sum1 <- binom.test(jury$Count[1], sum(jury$Count), p = jury$CensusProp[1], alternative = "two.sided", conf.level = 1-0.05/7)
b.sum2 <- binom.test(jury$Count[2], sum(jury$Count), p = jury$CensusProp[2], alternative = "two.sided", conf.level = 1-0.05/7)
b.sum3 <- binom.test(jury$Count[3], sum(jury$Count), p = jury$CensusProp[3], alternative = "two.sided", conf.level = 1-0.05/7)
b.sum4 <- binom.test(jury$Count[4], sum(jury$Count), p = jury$CensusProp[4], alternative = "two.sided", conf.level = 1-0.05/7)
b.sum5 <- binom.test(jury$Count[5], sum(jury$Count), p = jury$CensusProp[5], alternative = "two.sided", conf.level = 1-0.05/7)
b.sum6 <- binom.test(jury$Count[6], sum(jury$Count), p = jury$CensusProp[6], alternative = "two.sided", conf.level = 1-0.05/7)
b.sum7 <- binom.test(jury$Count[7], sum(jury$Count), p = jury$CensusProp[7], alternative = "two.sided", conf.level = 1-0.05/7)
# put the p-value and CI into a data.frame
b.sum <- data.frame(
  rbind( c(b.sum1$p.value, b.sum1$conf.int),
         , c(b.sum2$p.value, b.sum2$conf.int)
         , c(b.sum3$p.value, b.sum3$conf.int)
         , c(b.sum4$p.value, b.sum4$conf.int)
         , c(b.sum5$p.value, b.sum5$conf.int)
         , c(b.sum6$p.value, b.sum6$conf.int)
         , c(b.sum7$p.value, b.sum7$conf.int)
   )
)
names(b.sum) <- c("p.value", "CI.lower", "CI.upper")
b.sum$Age <- jury$Age
b.sum$Observed <- x.table$obs/sum(x.table$obs)
b.sum$CensusProp <- jury$CensusProp
b.sum

##      p.value CI.lower CI.upper Age Observed CensusProp
## 1 8.815e-15 0.009137 0.02920 18-19  0.01722    0.061
## 2 2.695e-18 0.054160 0.09294 20-24  0.07186    0.150
## 3 1.394e-04 0.079398 0.12435 25-29  0.10030    0.135
## 4 8.422e-01 0.189621 0.25120 30-39  0.21931    0.217
## 5 2.383e-11 0.192456 0.25433 40-49  0.22231    0.153
## 6 5.916e-20 0.251744 0.31881 50-64  0.28443    0.182
## 7 3.742e-02 0.065366 0.10708 65-99  0.08458    0.102
```

The CIs for the 30-39 and 65-99 year categories contain the census proportions. In the other five age categories, there are significant differences between the jury pool proportions and the census proportions. In general, young adults appear to be underrepresented in the jury pool whereas older age groups are overrepresented.

	Age	p.value	CI.lower	CI.upper	Observed	CensusProp
1	18-19	0.000	0.009	0.029	0.017	0.061
2	20-24	0.000	0.054	0.093	0.072	0.150
3	25-29	0.000	0.079	0.124	0.100	0.135
4	30-39	0.842	0.190	0.251	0.219	0.217
5	40-49	0.000	0.192	0.254	0.222	0.153
6	50-64	0.000	0.252	0.319	0.284	0.182
7	65-99	0.037	0.065	0.107	0.085	0.102

The residuals also highlight significant differences because the largest residuals correspond to the categories that contribute most to the value of χ^2_s . Some researchers use the residuals for the multiple comparisons, treating the Z_i s as

standard normal variables. Following this approach, you would conclude that the jury pool proportions differ from the proportions in the general population in every age category where $|Z_i| \geq 2.70$ (using the same Bonferroni correction). This gives the same conclusion as before.

The two multiple comparison methods are similar, but not identical. The residuals

$$Z_i = \frac{O_i - E_i}{\sqrt{E_i}} = \frac{\hat{p}_i - p_{0i}}{\sqrt{\frac{p_{0i}}{n}}}$$

agree with the large-sample statistic for testing $H_0 : p_i = p_{0i}$, except that the divisor in Z_i omits a $1 - p_{0i}$ term. The Z_i s are not standard normal random variables as assumed, and the value of Z_i underestimates the significance of the observed differences. Multiple comparisons using the Z_i s will find, on average, fewer significant differences than the preferred method based on the large sample tests. However, the differences between the two methods are usually minor when all of the hypothesized proportions are small.

7.5 Comparing Two Proportions: Independent Samples

The New Mexico state legislature is interested in how the proportion of registered voters that support Indian gaming differs between New Mexico and Colorado. Assuming neither population proportion is known, the state's statistician might recommend that the state conduct a survey of registered voters sampled independently from the two states, followed by a comparison of the sample proportions in favor of Indian gaming.

Statistical methods for comparing two proportions using independent samples can be formulated as follows. Let p_1 and p_2 be the proportion of populations 1 and 2, respectively, with the attribute of interest. Let \hat{p}_1 and \hat{p}_2 be the corresponding sample proportions, based on independent random or representative samples of size n_1 and n_2 from the two populations.

7.5.1 Large Sample CI and Tests for $p_1 - p_2$

A large-sample CI for $p_1 - p_2$ is $(\hat{p}_1 - \hat{p}_2) \pm z_{\text{crit}} SE_{CI}(\hat{p}_1 - \hat{p}_2)$, where z_{crit} is the standard normal critical value for the desired confidence level, and

$$SE_{CI}(\hat{p}_1 - \hat{p}_2) = \sqrt{\frac{\hat{p}_1(1 - \hat{p}_1)}{n_1} + \frac{\hat{p}_2(1 - \hat{p}_2)}{n_2}}$$

is the CI standard error.

A large-sample p-value for a test of the null hypothesis $H_0 : p_1 - p_2 = 0$ against the two-sided alternative $H_A : p_1 - p_2 \neq 0$ is evaluated using tail areas of the standard normal distribution (identical to one sample evaluation) in conjunction with the test statistic

$$z_s = \frac{\hat{p}_1 - \hat{p}_2}{SE_{test}(\hat{p}_1 - \hat{p}_2)},$$

where

$$SE_{test}(\hat{p}_1 - \hat{p}_2) = \sqrt{\frac{\bar{p}(1 - \bar{p})}{n_1} + \frac{\bar{p}(1 - \bar{p})}{n_2}} = \sqrt{\bar{p}(1 - \bar{p}) \left(\frac{1}{n_1} + \frac{1}{n_2} \right)}$$

is the test standard error for $\hat{p}_1 - \hat{p}_2$. The **pooled proportion**

$$\bar{p} = \frac{n_1 \hat{p}_1 + n_2 \hat{p}_2}{n_1 + n_2}$$

is the proportion of successes in the two samples combined. The test standard error has the same functional form as the CI standard error, with \bar{p} replacing the individual sample proportions.

The pooled proportion is the best guess at the common population proportion when $H_0 : p_1 = p_2$ is true. The test standard error estimates the standard deviation of $\hat{p}_1 - \hat{p}_2$ assuming H_0 is true.

Remark: As in the one-sample proportion problem, the test and CI SE's are different. This *can* (but usually does not) lead to some contradiction between the test and CI.

Example, vitamin C Two hundred and seventy nine (279) French skiers were studied during two one-week periods in 1961. One group of 140 skiers receiving a placebo each day, and the other 139 receiving 1 gram of ascorbic acid (Vitamin C) per day. The study was double blind — neither the subjects nor the researchers knew who received which treatment. Let p_1 be the probability that a member of the ascorbic acid group contracts a cold during the study period, and p_2 be the corresponding probability for the placebo group. Linus Pauling (Chemistry and Peace Nobel prize winner) and I are interested in testing whether $H_0 : p_1 = p_2$. The data are summarized below as a two-by-two table of counts (a contingency table)

Outcome	Ascorbic Acid	Placebo
# with cold	17	31
# with no cold	122	109
Totals	139	140

The sample sizes are $n_1 = 139$ and $n_2 = 140$. The sample proportion of skiers developing colds in the placebo and treatment groups are $\hat{p}_2 = 31/140 = 0.221$ and $\hat{p}_1 = 17/139 = 0.122$, respectively. The difference is $\hat{p}_1 - \hat{p}_2 = 0.122 - 0.221 = -0.099$. The pooled proportion is the number of skiers that developed colds divided by the number of skiers in the study: $\bar{p} = 48/279 = 0.172$.

The test standard error is

$$SE_{test}(\hat{p}_1 - \hat{p}_2) = \sqrt{0.172 \times (1 - 0.172) \left(\frac{1}{139} + \frac{1}{140} \right)} = 0.0452.$$

The test statistic is

$$z_s = \frac{0.122 - 0.221}{0.0452} = -2.19.$$

The p-value for a two-sided test is twice the area under the standard normal curve to the right of 2.19 (or twice the area to the left of -2.19), which is $2 \times (0.014) = 0.028$. At the 5% level, we reject the hypothesis that the probability of contracting a cold is the same whether you are given a placebo or Vitamin C.

A CI for $p_1 - p_2$ provides a measure of the size of the treatment effect. For a 95% CI

$$\begin{aligned} z_{\text{crit}} SE_{CI}(\hat{p}_1 - \hat{p}_2) &= 1.96 \sqrt{\frac{0.221 \times (1 - 0.221)}{140} + \frac{0.122 \times (1 - 0.122)}{139}} \\ &= 1.96 \times (0.04472) = 0.088. \end{aligned}$$

The 95% CI for $p_1 - p_2$ is -0.099 ± 0.088 , or $(-0.187, -0.011)$. We are 95% confident that p_2 exceeds p_1 by at least 0.011 but not by more than 0.187.

On the surface, we would conclude that a daily dose of Vitamin C decreases a French skier's chance of developing a cold by between 0.011 and 0.187 (with 95% confidence). This conclusion was somewhat controversial. Several reviews of the study felt that the experimenter's evaluations of cold symptoms were unreliable. Many other studies refute the benefit of Vitamin C as a treatment for the common cold.

```
#### Example, vitamin C
# Approximate normal test for two-proportions, without Yates' continuity correction
prop.test(c(17, 31), c(139, 140), correct = FALSE)

##
## 2-sample test for equality of proportions without continuity
## correction
##
## data: c(17, 31) out of c(139, 140)
## X-squared = 4.811, df = 1, p-value = 0.02827
## alternative hypothesis: two.sided
## 95 percent confidence interval:
## -0.18686 -0.01139
## sample estimates:
## prop 1 prop 2
## 0.1223 0.2214
```

Conditional probability

In probability theory, a conditional probability is the probability that an event will occur, when another event is known to occur or to have occurred. If the events are A and B respectively, this is said to be “the probability of A given B ”. It is commonly denoted by $\Pr(A|B)$. $\Pr(A|B)$ may or may not be equal to $\Pr(A)$, the probability of A . If they are equal, A and B are said to be independent. For example, if a coin is flipped twice, “the outcome of the second flip” is independent of “the outcome of the first flip”.

In the Vitamin C example above, the unconditional observed probability of contracting a cold is $\Pr(\text{cold}) = (17 + 31)/(139 + 140) = 0.172$. The conditional observed probabilities are $\Pr(\text{cold}|\text{ascorbic acid}) = 17/139 = 0.1223$ and $\Pr(\text{cold}|\text{placebo}) = 31/140 = 0.2214$. The two-sample test of $H_0 : p_1 = p_2$ where $p_1 = \Pr(\text{cold}|\text{ascorbic acid})$ and $p_2 = \Pr(\text{cold}|\text{placebo})$ is effectively testing whether $\Pr(\text{cold}) = \Pr(\text{cold}|\text{ascorbic acid}) = \Pr(\text{cold}|\text{placebo})$. This tests whether contracting a cold is independent of the vitamin C treatment.

Example, cervical dysplasia A case-control study was designed to examine risk factors for cervical dysplasia² (Becker et al. 1994). All the women in the study were patients at UNM clinics. The 175 cases were women, aged 18-40, who had cervical dysplasia. The 308 controls were women aged 18-40 who did not have cervical dysplasia. Each woman was classified as positive or negative, depending on the presence of HPV (human papilloma virus). The data are summarized below.

HPV Outcome	Cases	Controls
Positive	164	130
Negative	11	178
Sample size	175	308

Let p_1 be the probability that a case is HPV positive and let p_2 be the probability that a control is HPV positive. The sample sizes are $n_1 = 175$ and $n_2 = 308$. The sample proportions of positive cases and controls are $\hat{p}_1 = 164/175 = 0.937$ and $\hat{p}_2 = 130/308 = 0.422$.

For a 95% CI

$$\begin{aligned} z_{\text{crit}} SE_{CI}(\hat{p}_1 - \hat{p}_2) &= 1.96 \sqrt{\frac{0.937 \times (1 - 0.937)}{175} + \frac{0.422 \times (1 - 0.422)}{308}} \\ &= 1.96 \times (0.03336) = 0.0659. \end{aligned}$$

A 95% CI for $p_1 - p_2$ is $(0.937 - 0.422) \pm 0.066$, or 0.515 ± 0.066 , or $(0.449, 0.581)$. I am 95% confident that p_1 exceeds p_2 by at least 0.45 but not by more than 0.58.

²<http://www.ncbi.nlm.nih.gov/pubmedhealth/PMH0002461/>

```
# Approximate normal test for two-proportions, without Yates' continuity correction
prop.test(c(164, 130), c(175, 308), correct = FALSE)

##
## 2-sample test for equality of proportions without continuity
## correction
##
## data: c(164, 130) out of c(175, 308)
## X-squared = 124.3, df = 1, p-value < 2.2e-16
## alternative hypothesis: two.sided
## 95 percent confidence interval:
## 0.4492 0.5809
## sample estimates:
## prop 1 prop 2
## 0.9371 0.4221
```

Not surprisingly, a two-sided test at the 5% level would reject $H_0 : p_1 = p_2$. In this problem one might wish to do a one-sided test, instead of a two-sided test. Let us carry out this test, as a refresher on how to conduct one-sided tests.

```
# one-sided test, are cases more likely to be HPV positive?
prop.test(c(164, 130), c(175, 308), correct = FALSE, alternative = "greater")

##
## 2-sample test for equality of proportions without continuity
## correction
##
## data: c(164, 130) out of c(175, 308)
## X-squared = 124.3, df = 1, p-value < 2.2e-16
## alternative hypothesis: greater
## 95 percent confidence interval:
## 0.4598 1.0000
## sample estimates:
## prop 1 prop 2
## 0.9371 0.4221
```

Appropriateness of Large Sample Test and CI

The standard two-sample CI and test used above are appropriate when each sample is large. A rule of thumb suggests a minimum of at least five successes (i.e., observations with the characteristic of interest) and failures (i.e., observations without the characteristic of interest) in each sample before using these methods. This condition is satisfied in our two examples.



7.6 Effect Measures in Two-by-Two Tables

Consider a study of a particular disease, where each individual is either exposed or not-exposed to a risk factor. Let p_1 be the proportion diseased among the individuals in the exposed population, and p_2 be the proportion diseased among the non-exposed population. This population information can be summarized as a two-by-two table of population proportions:

Outcome	Exposed population	Non-Exposed population
Diseased	p_1	p_2
Non-Diseased	$1 - p_1$	$1 - p_2$

A standard measure of the difference between the exposed and non-exposed populations is the **absolute difference**: $p_1 - p_2$. We have discussed statistical methods for assessing this difference.

In many epidemiological and biostatistical settings, other measures of the difference between populations are considered. For example, the relative risk

$$\text{RR} = \frac{p_1}{p_2}$$

is commonly reported when the individual risks p_1 and p_2 are small. The odds ratio

$$\text{OR} = \frac{p_1/(1 - p_1)}{p_2/(1 - p_2)}$$

is another standard measure. Here $p_1/(1 - p_1)$ is the odds of being diseased in the exposed group, whereas $p_2/(1 - p_2)$ is the odds of being diseased in the non-exposed group.

I mention these measures because you may see them or hear about them. Note that each of these measures can be easily estimated from data, using the sample proportions as estimates of the unknown population proportions. For example, in the vitamin C study:

Outcome	Ascorbic Acid	Placebo
# with cold	17	31
# with no cold	122	109
Totals	139	140

the proportion with colds in the placebo group is $\hat{p}_2 = 31/140 = 0.221$. The proportion with colds in the vitamin C group is $\hat{p}_1 = 17/139 = 0.122$.

The estimated absolute difference in risk is $\hat{p}_1 - \hat{p}_2 = 0.122 - 0.221 = -0.099$. The estimated risk ratio and odds ratio are

$$\widehat{RR} = \frac{0.122}{0.221} = 0.55$$

and

$$\widehat{OR} = \frac{0.122/(1 - 0.122)}{0.221/(1 - 0.221)} = 0.49,$$

respectively.

7.7 Analysis of Paired Samples: Dependent Proportions

Paired and more general **block analyses** are appropriate with longitudinal data collected over time and in medical studies where several treatments are given to the same patient over time. A key feature of these designs that invalidates the two-sample method discussed earlier is that repeated observations within a unit or individual are likely to be correlated, and not independent.

Example, President performance For example, in a random sample of $n = 1600$ voter-age Americans, 944 said that they approved of the President's performance. One month later, only 880 of the original 1600 sampled approved. The following two-by-two table gives the numbers of individuals with each of the four possible sequences of responses over time. Thus, 150 voter-age Americans approved of the President's performance when initially asked but then disapproved one month later. The row and column totals are the numbers of approvals and disapprovals for the two surveys (Agresti, 1990, p. 350).

(Obs Counts)		Second survey		
First Survey	Approve	Approve	Disapprove	Total
		794	150	944
Approve	86	570	656	
Disapprove				1600
Total	880	720		

Let p_{AA} , p_{AD} , p_{DA} , p_{DD} be the population proportion of voter-age Americans that fall into the four categories, where the subscripts preserve the time ordering and indicate Approval or Disapproval. For example, p_{AD} is the population proportion that approved of the President's performance initially and disapproved one month later. The population proportion that initially approved is $p_{A+} = p_{AA} + p_{AD}$. The population proportion that approved at the time of the second survey is $p_{+A} = p_{AA} + p_{DA}$. The “+” sign used as a subscript means that the replaced subscript has been summed over.

Similarly, let \hat{p}_{AA} , \hat{p}_{AD} , \hat{p}_{DA} , \hat{p}_{DD} be the sample proportion of voter-age Americans that fall into the four categories, and let $\hat{p}_{A+} = \hat{p}_{AA} + \hat{p}_{AD}$ and $\hat{p}_{+A} = \hat{p}_{AA} + \hat{p}_{DA}$ be the sample proportion that approves the first month and the sample proportion that approves the second month, respectively. The table below summarizes the observed proportions. For example, $\hat{p}_{AA} = 794/1600 = 0.496$ and $\hat{p}_{A+} = 944/1600 = 0.496 + 0.094 = 0.590$. The sample proportion of voting-age Americans that approve of the President's performance decreased from one month to the next.

(Obs Proportions)		Second survey		
	First Survey	Approve	Disapprove	Total
Approve		0.496	0.094	0.590
Disapprove		0.054	0.356	0.410
Total		0.550	0.450	1.000

The difference in the population proportions from one month to the next can be assessed by a large-sample CI for $p_{A+} - p_{+A}$, given by $(\hat{p}_{A+} - \hat{p}_{+A}) \pm z_{\text{crit}} SE_{CI}(\hat{p}_{A+} - \hat{p}_{+A})$, where the CI standard error satisfies

$$SE_{CI}(\hat{p}_{A+} - \hat{p}_{+A}) = \sqrt{\frac{\hat{p}_{A+}(1 - \hat{p}_{A+}) + \hat{p}_{+A}(1 - \hat{p}_{+A}) - 2(\hat{p}_{AA}\hat{p}_{DD} - \hat{p}_{AD}\hat{p}_{DA})}{n}}$$

One-sided bounds are constructed in the usual way.

The $-2(\cdot)$ term in the standard error accounts for the dependence between the samples at the two time points. If independent samples of size n had been selected for the two surveys, then this term would be omitted from the standard error, giving the usual two-sample CI.

For a 95% CI in the Presidential survey,

$$\begin{aligned} z_{\text{crit}} SE_{CI}(\hat{p}_{A+} - \hat{p}_{+A}) &= 1.96 \sqrt{\frac{0.590 \times 0.410 + 0.550 \times 0.450 - 2(0.496 \times 0.356 - 0.094 \times 0.054)}{1600}} \\ &= 1.96 \times (0.0095) = 0.0186. \end{aligned}$$

A 95% CI for $p_{A+} - p_{+A}$ is $(0.590 - 0.550) \pm 0.019$, or $(0.021, 0.059)$. You are 95% confident that the population proportion of voter-age Americans that approved of the President's performance the first month was between 0.021 and 0.059 larger than the proportion that approved one month later. This gives evidence of a decrease in the President's approval rating.

A test of $H_0 : p_{A+} = p_{+A}$ can be based on the CI for $p_{A+} - p_{+A}$, or on a standard normal approximation to the test statistic

$$z_s = \frac{\hat{p}_{A+} - \hat{p}_{+A}}{SE_{test}(\hat{p}_{A+} - \hat{p}_{+A})},$$

where the test standard error is given by

$$SE_{test}(\hat{p}_{A+} - \hat{p}_{+A}) = \sqrt{\frac{\hat{p}_{A+}\hat{p}_{+A} - 2\hat{p}_{AA}}{n}}.$$

The test statistic is often written in the simplified form

$$z_s = \frac{n_{AD} - n_{DA}}{\sqrt{n_{AD} + n_{DA}}},$$

where the n_{ij} s are the observed cell counts. An equivalent form of this test, based on comparing the square of z_s to a chi-squared distribution with 1 degree of freedom, is the well-known **McNemar's test** for marginal homogeneity (or symmetry) in the two-by-two table.

For example, in the Presidential survey

$$z_s = \frac{150 - 86}{\sqrt{150 + 86}} = 4.17.$$

The p-value for a two-sided test is, as usual, the area under the standard normal curve outside ± 4.17 . The p-value is less than 0.001, suggesting that H_0 is false.

R can perform this test as McNemar's test.

```

##### Example, President performance
# McNemar's test needs data as a matrix

# Presidential Approval Ratings.
# Approval of the President's performance in office in two surveys,
# one month apart, for a random sample of 1600 voting-age Americans.
pres <-
matrix(c(794, 150, 86, 570),
       nrow = 2, byrow = TRUE,
       dimnames = list("1st Survey" = c("Approve", "Disapprove"),
                      "2nd Survey" = c("Approve", "Disapprove")))
pres
##                2nd Survey
## 1st Survey    Approve Disapprove
##   Approve        794      150
##   Disapprove     86       570
mcnemar.test(pres, correct=FALSE)
##
## McNemar's Chi-squared test
##
## data: pres
## McNemar's chi-squared = 17.36, df = 1, p-value = 3.099e-05
# => significant change (in fact, drop) in approval ratings

```

7.8 Testing for Homogeneity of Proportions

Example, cancer deaths The following two-way table of counts summarizes the location of death and age at death from a study of 2989 cancer deaths (Public Health Reports, 1983).

(Obs Counts)		Location of death				
		Age	Home	Acute Care	Chronic care	Row Total
	15-54		94	418	23	535
	55-64		116	524	34	674
	65-74		156	581	109	846
	75+		138	558	238	934
	Col Total		504	2081	404	2989

The researchers want to compare the age distributions across locations. A one-way ANOVA would be ideal if the actual ages were given. Because the ages are grouped, the data should be treated as categorical. Given the differences in

numbers that died at the three types of facilities, a comparison of proportions or percentages in the age groups is appropriate. A comparison of counts is not.

The table below summarizes the proportion in the four age groups by location. For example, in the acute care facility $418/2081 = 0.201$ and $558/2081 = 0.268$. The **pooled proportions** are the Row Totals divided by the total sample size of 2989. The pooled summary gives the proportions in the four age categories, ignoring location of death.

The age distributions for home and for the acute care facilities are similar, but are very different from the age distribution at chronic care facilities.

To formally compare the observed proportions, one might view the data as representative sample of ages at death from the three locations. Assuming independent samples from the three locations (populations), a chi-squared statistic is used to test whether the population proportions of ages at death are identical (homogeneous) across locations. The **chi-squared test for homogeneity** of population proportions can be defined in terms of proportions, but is traditionally defined in terms of counts.

(Proportions)		Location of death			
	Age	Home	Acute Care	Chronic care	Pooled
	15-54	0.187	0.201	0.057	0.179
	55-64	0.230	0.252	0.084	0.226
	65-74	0.310	0.279	0.270	0.283
	75+	0.273	0.268	0.589	0.312
	Total	1.000	1.000	1.000	1.000

In general, assume that the data are independent samples from c populations (strata, groups, sub-populations), and that each individual is placed into one of r levels of a categorical variable. The raw data will be summarized as a $r \times c$ **contingency table** of counts, where the columns correspond to the samples, and the rows are the levels of the categorical variable. In the age distribution problem, $r = 4$ and $c = 3$.

To implement the test:

1. Compute the (estimated) **expected** count for each cell in the table as follows:

$$E = \frac{\text{Row Total} \times \text{Column Total}}{\text{Total Sample Size}}$$

2. Compute the Pearson test statistic

$$\chi_s^2 = \sum_{\text{all cells}} \frac{(O - E)^2}{E},$$

where O is the **observed** count.

3. For a size α test, reject the hypothesis of homogeneity if $\chi_s^2 \geq \chi_{\text{crit}}^2$, where χ_{crit}^2 is the upper α critical value from the chi-squared distribution with $df = (r - 1)(c - 1)$.

The p-value for the chi-squared test of homogeneity is equal to the area under the chi-squared curve to the right of χ_s^2 .

For a two-by-two table of counts, the chi-squared test of homogeneity of proportions is identical to the two-sample proportion test we discussed earlier.

The (estimated) expected counts for the (15-54, Home) cell and for the (75+, Acute Care) cell in the age distribution data are $E = 535 \times 504/2989 = 90.21$ and $E = 934 \times 2081/2989 = 650.27$, respectively. The other expected counts were computed similarly, and are summarized below. The row and column sums on the tables of observed and expected counts always agree.

(Exp Counts)		Location of death			Row Total
Age		Home	Acute Care	Chronic care	
15-54		90.21	372.48	72.31	535
55-64		113.65	469.25	91.10	674
65-74		142.65	589	114.35	846
75-		157.49	650.27	126.24	934
Col Total		504	2081	404	2989

Why is a comparison of the observed and expected counts relevant for testing homogeneity? To answer this question, first note that the expected cell count can be expressed

$$E = \text{Col Total} \times \text{Pooled proportion for category}.$$

For example, $E = 504 \times 0.179 = 90.21$ in the (15-54, Home) cell. A comparison of the observed and expected counts is a comparison of the observed category proportions in a location with the pooled proportions, taking the size of each

sample into consideration. Thinking of the pooled proportion as a weighted average of the sample proportions for a category, the Pearson χ^2_s statistic is an aggregate measure of variation in the observed proportions across samples. If the category proportions are similar across samples then the category and pooled proportions are similar, resulting in a “small” value of χ^2_s . Large values of χ^2_s occur when there is substantial variation in the observed proportions across samples, in one or more categories. In this regard, the Pearson statistic is similar to the F -statistic in a one-way ANOVA (where large differences between groups result in a large F -statistic).

```
#### Example, cancer deaths
canddeath <-
matrix(c( 94, 418, 23,
       116, 524, 34,
       156, 581, 109,
       138, 558, 238),
      nrow = 4, byrow = TRUE,
      dimnames = list("Age" = c("15-54", "55-64", "65-74", "75+"),
                      "Location of death" = c("Home", "Acute Care", "Chronic care")))
canddeath
##          Location of death
## Age      Home Acute Care Chronic care
## 15-54    94     418      23
## 55-64   116     524      34
## 65-74   156     581      109
## 75+     138     558      238

chisq.summary <- chisq.test(canddeath, correct=FALSE)
chisq.summary
##
##  Pearson's Chi-squared test
##
## data:  canddeath
## X-squared = 197.6, df = 6, p-value < 2.2e-16

# The Pearson residuals
chisq.summary$residuals
##          Location of death
## Age      Home Acute Care Chronic care
## 15-54  0.3990   2.3587   -5.7989
## 55-64  0.2206   2.5274   -5.9824
## 65-74  1.1177  -0.3297   -0.5001
## 75+   -1.5530  -3.6183    9.9467

# The sum of the squared residuals is the chi-squared statistic:
chisq.summary$residuals^2
##          Location of death
```

```

## Age      Home Acute Care Chronic care
## 15-54  0.15916    5.5636   33.6274
## 55-64  0.04865    6.3875   35.7888
## 65-74  1.24916    0.1087   0.2501
## 75+    2.41184    13.0924  98.9369
sum(chisq.summary$residuals^2)
## [1] 197.6

```

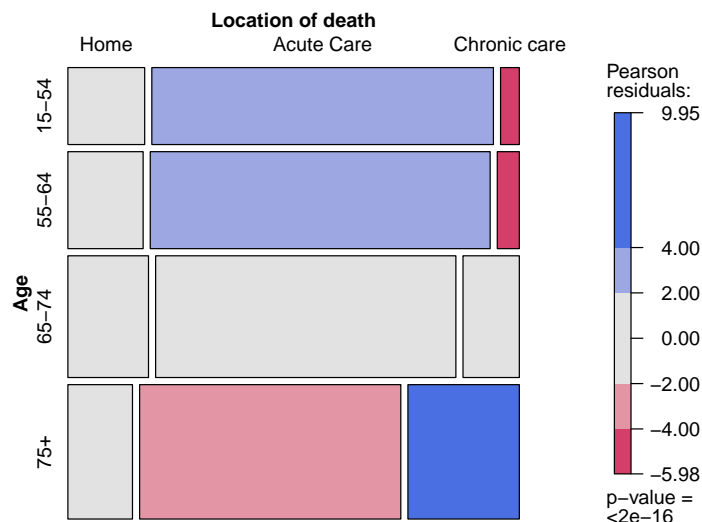
A visualization of the Pearson residuals is available with a `mosaic()` plot in the `vcd` package. Extended mosaic and association plots are each helpful methods of visualizing complex data and evaluating deviations from a specified independence model. For extended mosaic plots, use `mosaic(x, condvar=, data=)` where `x` is a table or formula, `condvar=` is an optional conditioning variable, and `data=` specifies a data frame or a table. Include `shade=TRUE` to color the figure, and `legend=TRUE` to display a legend for the Pearson residuals.

```

# mosaic plot
library(vcd)
mosaic(canddeath, shade=TRUE, legend=TRUE)

## association plot (10/28/2013 worked before, not working now)
#library(vcd)
#assoc(canddeath, shade=TRUE)

```



The `vcd` package provides a variety of methods for visualizing multivariate categorical data, inspired by Michael Friendly's wonderful "Visualizing Cate-

gorical Data". For more details, see The Strucplot Framework³.

7.8.1 Adequacy of the Chi-Square Approximation

The chi-squared tests for homogeneity and independence are large-sample tests. As with the goodness-of-fit test, a simple rule-of-thumb is that the approximation is adequate when the expected (not observed) cell counts are 5 or more. This rule is conservative, and some statisticians argue that the approximation is valid for expected counts as small as one.

In practice, the chi-squared approximation to χ_s^2 tends to be a bit conservative, meaning that statistically significant results would likely retain significance had a more accurate approximation been used.

R may not print out a warning message whenever a noticeable percentage of cells have expected counts less than 5. Ideally, one would use Fisher's exact test (`fisher.test()`) for tables with small counts, and can be used for larger than 2-by-2 tables when the frequencies are not too large.

7.9 Testing for Homogeneity in Cross-Sectional and Stratified Studies

Two-way tables of counts are often collected using either **stratified sampling** or **cross-sectional** sampling.

In a **stratified design**, distinct groups, strata, or sub-populations are identified. Independent samples are selected from each group, and the sampled individuals are classified into categories. The Indian gaming example is an illustration of a stratified design (where the two strata were NM and CO voters). Stratified designs provide estimates for the strata (population) proportion in each of the categories. A test for **homogeneity of proportions** is used to compare the strata.

In a **cross-sectional design**, individuals are randomly selected from a population and classified by the levels of **two** categorical variables. With cross-

³<http://cran.r-project.org/web/packages/vcd/vignettes/strucplot.pdf>

sectional samples you can test homogeneity of proportions by comparing either the row proportions or by comparing the column proportions.

Example, antismoking adverts The following data (*The Journal of Advertising*, 1983, pp. 34–42) are from a cross-sectional study that involved soliciting opinions on anti-smoking advertisements. Each subject was asked whether they smoked and their reaction (on a five-point ordinal scale) to the ad. The data are summarized as a two-way table of counts, given below:

	Str. Dislike	Dislike	Neutral	Like	Str. Like	Row Tot
Smoker	8	14	35	21	19	97
Non-smoker	31	42	78	61	69	281
Col Total	39	56	113	82	88	378

The row proportions (i.e., fix a row and compute the proportions for the column categories) are

(Row Prop)	Str. Dislike	Dislike	Neutral	Like	Str. Like	Row Tot
Smoker	0.082	0.144	0.361	0.216	0.196	1.000
Non-smoker	0.110	0.149	0.278	0.217	0.245	1.000

For example, the entry for the (Smoker, Str. Dislike) cell is: $8/97 = 0.082$.

Similarly, the column proportions are

(Col Prop)	Str. Dislike	Dislike	Neutral	Like	Str. Like
Smoker	0.205	0.250	0.310	0.256	0.216
Non-smoker	0.795	0.750	0.690	0.744	0.784
Total	1.000	1.000	1.000	1.000	1.000

Although it may be more natural to compare the smoker and non-smoker row proportions, the column proportions can be compared across ad responses. There is no advantage to comparing “rows” instead of “columns” in a formal test of homogeneity of proportions with cross-sectional data. The Pearson chi-squared test treats the rows and columns interchangeably, so you get the same result regardless of how you view the comparison. However, one of the two comparisons may be more natural to interpret.

Note that checking for homogeneity of proportions is meaningful in stratified studies only when the comparison is between strata! Further, if the strata correspond to columns of the table, then the

column proportions or percentages are meaningful whereas the row proportions are not.

7.9.1 Testing for Independence in a Two-Way Contingency Table

The row and column classifications for a population where each individual is cross-classified by two categorical variables are said to be **independent** if each **population** cell proportion in the two-way table is the product of the proportion in a given row and the proportion in a given column. One can show that independence is equivalent to homogeneity of proportions. In particular, the two-way table of population cell proportions satisfies independence if and only if the population column proportions are homogeneous. If the population column proportions are homogeneous then so are the population row proportions.

This suggests that a test for independence or **no association** between two variables based on a cross-sectional study can be implemented using the chi-squared test for homogeneity of proportions. This suggestion is correct. If independence is not plausible, I interpret the dependence as a deviation from homogeneity, using the classification for which the interpretation is most natural.

The Pearson chi-squared test of independence is not significant (p -value = 0.56). The observed association between smoking status and the ad reaction is not significant. This suggests, for example, that the smoker's reactions to the ad were not statistically significantly different from the non-smoker's reactions, which is consistent with the smokers and non-smokers attitudes being fairly similar.

```
#### Example, antismoking adverts
antismokead <-
matrix(c( 8, 14, 35, 21, 19,
       31, 42, 78, 61, 69),
      nrow = 2, byrow = TRUE,
      dimnames = list(
        "Status" = c("Smoker", "Non-smoker"),
        "Reaction" = c("Str. Dislike", "Dislike", "Neutral", "Like", "Str. Like")))
antismokead
##          Reaction
## Status     Str. Dislike Dislike Neutral Like Str. Like
```

```

##   Smoker          8      14     35    21      19
## Non-smoker       31      42     78    61      69
chisq.summary <- chisq.test(antismokead, correct=FALSE)
chisq.summary

##
## Pearson's Chi-squared test
##
## data: antismokead
## X-squared = 2.991, df = 4, p-value = 0.5594

# All expected frequencies are at least 5
chisq.summary$expected

##           Reaction
## Status      Str. Dislike Dislike Neutral Like Str. Like
## Smoker        10.01 14.37     29 21.04    22.58
## Non-smoker    28.99 41.63     84 60.96    65.42

# Contribution to chi-squared statistic
chisq.summary$residuals^2

##           Reaction
## Status      Str. Dislike Dislike Neutral      Like Str. Like
## Smoker        0.4029 0.009546  1.2426 8.515e-05    0.5682
## Non-smoker    0.1391 0.003295  0.4289 2.939e-05    0.1961

```

7.9.2 Further Analyses in Two-Way Tables

The χ_s^2 statistic is a **summary measure** of independence or homogeneity. A careful look at the data usually reveals the nature of the **association** or **heterogeneity** when the test is significant. There are numerous meaningful ways to explore two-way tables to identify sources of association or heterogeneity. For example, in the comparison of age distributions across locations, you might consider the 4×2 tables comparing all possible pairs of locations. Another possibility would be to compare the proportion in the 75+ age category across locations. For the second comparison you need a 2×3 table of counts, where the two rows correspond to the individuals less than 75 years old and those 75+ years old, respectively (i.e., collapse the first three rows of the original 4×2 table). The possibilities are almost limitless in large tables. Of course, theoretically generated comparisons are preferred to data dredging.

Example: drugs and nausea, testing for homogeneity A randomized double-blind experiment compared the effectiveness of several drugs in reducing postoperative nausea. All patients were anesthetized with nitrous oxide and ether. The following table shows the incidence of nausea during the first four postoperative hours of four drugs and a placebo. Compare the drugs to each other and to the placebo.

Drug	# with Nausea	# without Nausea	Sample Size
Placebo	96	70	166
Chlorpromazine	52	100	152
Dimenhydrinate	52	33	85
Pentobarbitol (100mg)	35	32	67
Pentobarbitol (150mg)	37	48	85

Let p_{PL} be the probability of developing nausea given a placebo, and define p_{CH} , p_{DI} , p_{PE100} , and p_{PE150} analogously. A simple initial analysis would be to test homogeneity of proportions: $H_0 : p_{PL} = p_{CH} = p_{DI} = p_{PE100} = p_{PE150}$ against $H_A : \text{not } H_0$.

The data were entered as frequencies. The output shows that the proportion of patients exhibiting nausea (see the **column** percents — the cell and row percentages are not interpretable, so they are omitted) is noticeably different across drugs. In particular, Chlorpromazine is the most effective treatment with $\hat{p}_{CH} = 0.34$ and Dimenhydrinate is the least effective with $\hat{p}_{DI} = 0.61$.

The p-value for the chi-squared test is 0.00005, which leads to rejecting H_0 at the 0.05 or 0.01 levels. The data strongly suggest there are differences in the effectiveness of the various treatments for postoperative nausea.

```
#### Example: drugs and nausea, testing for homogeneity
nausea <-
matrix(c(96, 70, 52, 100, 52, 33, 35, 32, 37, 48),
      nrow = 5, byrow = TRUE,
      dimnames = list("Drug" = c("PL", "CH", "DI", "PE100", "PE150"),
                     "Result" = c("Nausea", "No Nausea")))
nausea

##          Result
## Drug    Nausea No Nausea
## PL      96     70
## CH      52     100
## DI      52      33
## PE100   35      32
## PE150   37      48
```

```
# Sorted proportions of nausea by drug
nausea.prop <- sort(nausea[,1]/rowSums(nausea))
nausea.prop
##      CH   PE150   PE100      PL      DI
## 0.3421 0.4353 0.5224 0.5783 0.6118

# chi-sq test of association
chisq.summary <- chisq.test(nausea, correct=FALSE)
chisq.summary
##
## Pearson's Chi-squared test
##
## data: nausea
## X-squared = 24.83, df = 4, p-value = 5.451e-05

# All expected frequencies are at least 5
chisq.summary$expected
##
##      Result
## Drug    Nausea No Nausea
##   PL     81.35    84.65
##   CH     74.49    77.51
##   DI     41.66    43.34
##   PE100  32.84    34.16
##   PE150  41.66    43.34
```

A sensible follow-up analysis is to identify which treatments were responsible for the significant differences. For example, the placebo and chlorpromazine can be compared using a test of $p_{PL} = p_{CH}$ or with a CI for $p_{PL} - p_{CH}$.

In certain experiments, specific comparisons are of interest, for example a comparison of the drugs with the placebo. Alternatively, all possible comparisons might be deemed relevant. The second case is suggested here based on the problem description. I will use a Bonferroni adjustment to account for the multiple comparisons. The Bonferroni adjustment accounts for data dredging, but at a cost of less sensitive comparisons.

There are 10 possible comparisons here. The Bonferroni analysis with an overall Family Error Rate of 0.05 (or less) tests the 10 individual hypotheses at the $0.05/10=0.005$ level.

```
nausea.table <- data.frame(Interval = rep(NA,10)
                           , CI.lower = rep(NA,10)
                           , CI.upper = rep(NA,10)
                           , Z = rep(NA,10)
                           , p.value = rep(NA,10)
                           , sig.temp = rep(NA,10)
                           , sig = rep(NA,10))
# row names for table
nausea.table[,1] <- c("p_PL - p_CH")
```

```

, "p_PL - p_DI"
, "p_PL - p_PE100"
, "p_PL - p_PE150"
, "p_CH - p_DI"
, "p_CH - p_PE100"
, "p_CH - p_PE150"
, "p_DI - p_PE100"
, "p_DI - p_PE150"
, "p_PE100 - p_PE150")
# test results together in a table
i.tab <- 0
for (i in 1:4) {
  for (j in (i+1):5) {
    i.tab <- i.tab + 1
    nausea.summary <- prop.test(nausea[c(i,j),], correct = FALSE, conf.level = 1-0.05/10)
    nausea.table[i.tab, 2:6] <- c(NA,
      , nausea.summary$conf.int[1]
      , nausea.summary$conf.int[2]
      , sign(-diff(NAUSEA.SUMMARY$ESTIMATE)) * NAUSEA.SUMMARY$STATISTIC^0.5
      , NAUSEA.SUMMARY$P.VALUE
      , (NAUSEA.SUMMARY$P.VALUE < 0.05/10))
    if (NAUSEA.TABLE$SIG.TEMP[i.tab] == 1) { NAUSEA.TABLE$SIG[i.tab] <- "*"}
      else { NAUSEA.TABLE$SIG[i.tab] <- " " }
  }
}
# remove temporary sig indicator
NAUSEA.TABLE <- subset(NAUSEA.TABLE, select = -SIG.TEMP)
#NAUSEA.TABLE

```

The following table gives two-sample tests of proportions with nausea and 99.5% CIs for the differences between the ten pairs of proportions. The only two p-values are less than 0.005 corresponding to $p_{PL} - p_{CH}$ and $p_{CH} - p_{DI}$. I am 99.5% confident that p_{CH} is between 0.084 and 0.389 less than p_{PL} , and I am 99.5% confident that p_{CH} is between 0.086 and 0.453 less than p_{DI} . The other differences are not significant.

	Interval	CI.lower	CI.upper	Z	p.value	sig
1	$p_{PL} - p_{CH}$	0.0838	0.3887	4.2182	0.0000	*
2	$p_{PL} - p_{DI}$	-0.2167	0.1498	-0.5099	0.6101	
3	$p_{PL} - p_{PE100}$	-0.1464	0.2582	0.7788	0.4361	
4	$p_{PL} - p_{PE150}$	-0.0424	0.3284	2.1485	0.0317	
5	$p_{CH} - p_{DI}$	-0.4532	-0.0861	-4.0122	0.0001	*
6	$p_{CH} - p_{PE100}$	-0.3828	0.0222	-2.5124	0.0120	
7	$p_{CH} - p_{PE150}$	-0.2788	0.0924	-1.4208	0.1554	
8	$p_{DI} - p_{PE100}$	-0.1372	0.3160	1.1058	0.2688	
9	$p_{DI} - p_{PE150}$	-0.0352	0.3881	2.3034	0.0213	
10	$p_{PE100} - p_{PE150}$	-0.1412	0.3154	1.0677	0.2857	

Using ANOVA-type groupings, and arranging the treatments from most to least effective (low proportions to high), we get:

CH (0.34) PE150 (0.44) PE100 (0.52) PL (0.58) DI (0.61)

Chapter 8

Correlation and Regression

Learning objectives

After completing this topic, you should be able to:

select graphical displays that reveal the relationship between two continuous variables.

summarize model fit.

interpret model parameters, such as slope and R^2 .

assess the model assumptions visually and numerically.

Achieving these goals contributes to mastery in these course learning outcomes:

1. organize knowledge.
5. define parameters of interest and hypotheses in words and notation.
6. summarize data visually, numerically, and descriptively.
8. use statistical software.
12. make evidence-based decisions.

8.1 Introduction

Suppose we select $n = 10$ people from the population of college seniors who plan to take the medical college admission test (MCAT) exam. Each takes the test, is coached, and then retakes the exam. Let X_i be the pre-coaching score and let Y_i be the post-coaching score for the i^{th} individual, $i = 1, 2, \dots, n$.

There are several questions of potential interest here, for example: Are Y and X related (associated), and how? Does coaching improve your MCAT score? Can we use the data to develop a mathematical model (formula) for predicting post-coaching scores from the pre-coaching scores? These questions can be addressed using **correlation** and **regression** models.

The **correlation coefficient** is a standard measure of **association** or relationship between two features Y and X . Most scientists equate Y and X being correlated to mean that Y and X are associated, related, or **dependent** upon each other. However, correlation is only a measure of the strength of a **linear relationship**. For later reference, let ρ be the correlation between Y and X in the population and let r be the sample correlation. I define r below. The population correlation is defined analogously from population data.

Suppose each of n sampled individuals is measured on two quantitative characteristics called Y and X . The data are pairs of observations (X_1, Y_1) , (X_2, Y_2) , \dots , (X_n, Y_n) , where (X_i, Y_i) is the (X, Y) pair for the i^{th} individual in the sample. The sample correlation between Y and X , also called the **Pearson product moment correlation coefficient**, is

$$r = \frac{S_{XY}}{S_X S_Y} = \frac{\sum_i^n (X_i - \bar{X})(Y_i - \bar{Y})}{\sqrt{\sum_i^n (X_i - \bar{X})^2} \sqrt{\sum_i^n (Y_i - \bar{Y})^2}},$$

where

$$S_{XY} = \frac{\sum_{i=1}^n (X_i - \bar{X})(Y_i - \bar{Y})}{n - 1}$$

is the **sample covariance** between Y and X , and $S_Y = \sqrt{\sum_i^n (Y_i - \bar{Y})^2 / (n - 1)}$ and $S_X = \sqrt{\sum_i^n (X_i - \bar{X})^2 / (n - 1)}$ are the standard deviations for the Y and X samples.

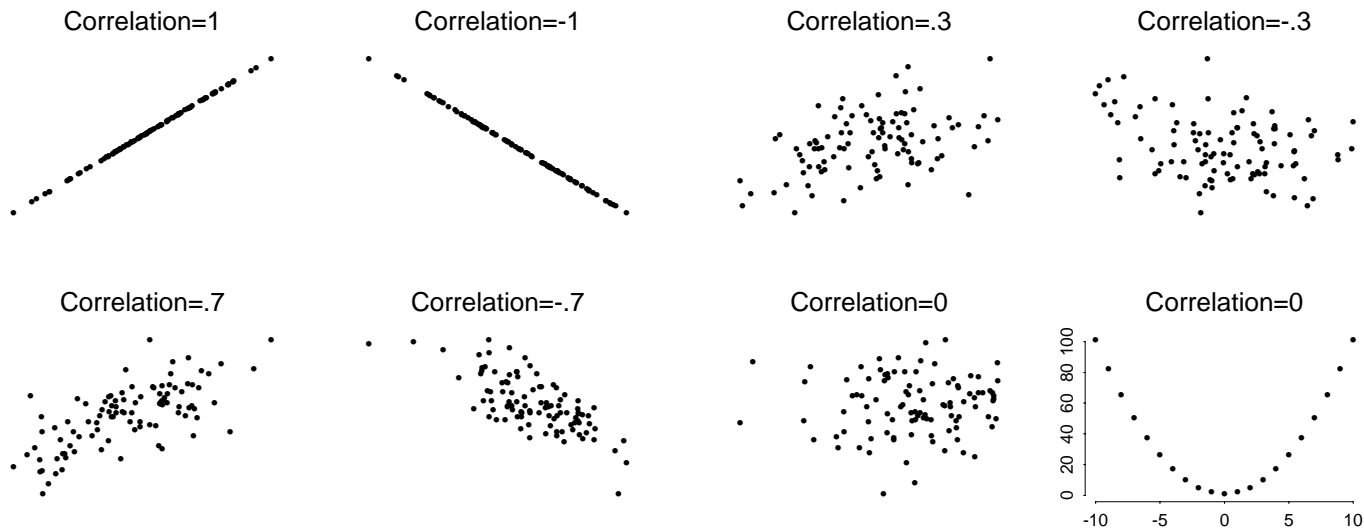
Important properties of r :

1. $-1 \leq r \leq 1$.
2. If Y_i tends to increase linearly with X_i then $r > 0$.
3. If Y_i tends to decrease linearly with X_i then $r < 0$.
4. If there is a perfect linear relationship between Y_i and X_i with a positive slope then $r = +1$.

5. If there is a perfect linear relationship between Y_i and X_i with a negative slope then $r = -1$.
6. The closer the points (X_i, Y_i) come to forming a straight line, the closer r is to ± 1 .
7. The magnitude of r is unchanged if either the X or Y sample is transformed linearly (such as feet to inches, pounds to kilograms, Celsius to Fahrenheit).
8. The correlation does not depend on which variable is called Y and which is called X .

If r is near ± 1 , then there is a strong linear relationship between Y and X in the sample. This suggests we might be able to accurately predict Y from X with a linear equation (i.e., linear regression). If r is near 0, there is a weak linear relationship between Y and X , which suggests that a linear equation provides little help for predicting Y from X . The pictures below should help you develop a sense about the size of r .

Note that $r = 0$ does not imply that Y and X are not related in the sample. It only implies they are not linearly related. For example, in the last plot $r = 0$ yet $Y_i = X_i^2$, exactly.



■ CLICKER Qs — Correlation coefficients STT.02.02.010 ■

■ CLICKER Qs — Strong correlation STT.02.02.020 ■

8.2 Testing that $\rho = 0$

Suppose you want to test $H_0 : \rho = 0$ against $H_A : \rho \neq 0$, where ρ is the population correlation between Y and X . This test is usually interpreted as a test of no association, or relationship, between Y and X in the population. Keep in mind, however, that ρ measures the strength of a **linear** relationship.

The standard test of $H_0 : \rho = 0$ is based on the magnitude of r . If we let

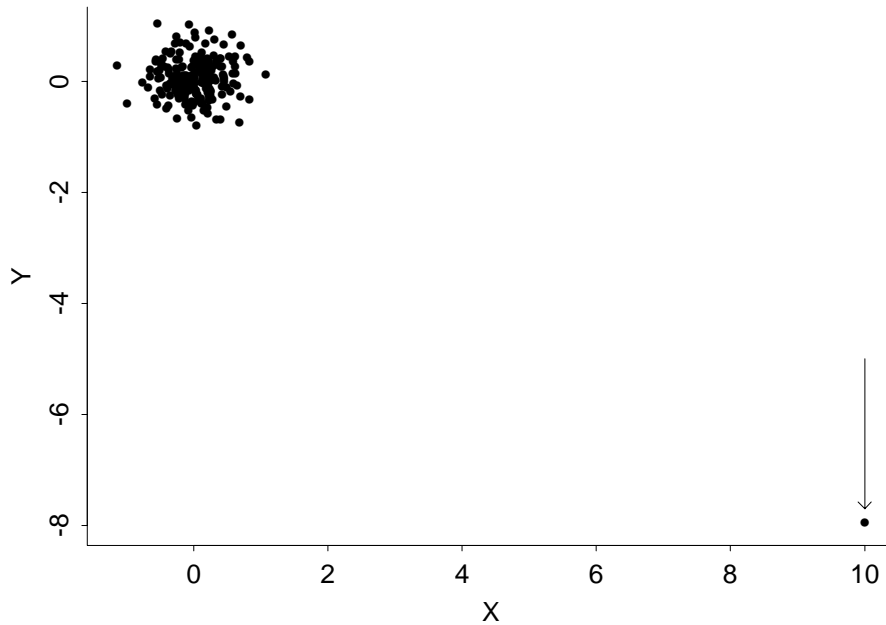
$$t_s = r \sqrt{\frac{n - 2}{1 - r^2}},$$

then the test rejects H_0 in favor of H_A if $|t_s| \geq t_{\text{crit}}$, where t_{crit} is the two-sided test critical value from a t -distribution with $df = n - 2$. The p-value for the test is the area under the t -curve outside $\pm t_s$ (i.e., two-tailed test p-value).

This test assumes that the data are a random sample from a **bivariate normal population** for (X, Y) . This assumption implies that all linear combinations of X and Y , say $aX + bY$, are normal. In particular, the (marginal) population frequency curves for X and Y are normal. At a minimum, you should make boxplots of the X and Y samples to check marginal normality. For large-sized samples, a plot of Y against X should be roughly an elliptical cloud, with the density of the points decreasing as the points move away from the center of the cloud.

8.3 The Spearman Correlation Coefficient

The Pearson correlation r can be highly influenced by outliers in one or both samples. For example, $r \approx -1$ in the plot below. If you delete the one extreme case with the largest X and smallest Y value then $r \approx 0$. The two analyses are contradictory. The first analysis (ignoring the plot) suggests a strong linear relationship, whereas the second suggests the lack of a linear relationship. I will not strongly argue that you should (must?) delete the extreme case, but I am concerned about any conclusion that depends heavily on the presence of a single observation in the data set.



Spearman's rank correlation coefficient r_S is a sensible alternative to r when normality is unreasonable or outliers are present. Most books give a computational formula for r_S . I will verbally describe how to compute r_S . First, order the X_i s and assign them ranks. Then do the same for the Y_i s and replace the original data pairs by the pairs of ranked values. The Spearman rank correlation is the Pearson correlation computed from the pairs of ranks.

The Spearman correlation r_S estimates the **population rank correlation coefficient**, which is a measure of the strength of linear relationship between population ranks. The Spearman correlation, as with other rank-based methods, is not sensitive to the presence of outliers in the data (or any information about the marginal distribution of X or Y). In the plot above, $r_S \approx 0$ whether the unusual point is included or excluded from the analysis. In samples without unusual observations and a linear trend, you often find that the Spearman and Pearson correlations are similar, $r_S \approx r$.

An important point to note is that the magnitude of the Spearman correlation does not change if either X or Y or both are transformed (monotonically). Thus, if r_S is noticeably greater than r , a transformation of the data might provide a stronger linear relationship.

Example: Blood loss Eight patients underwent a thyroid operation. Three variables were measured on each patient: weight in kg, time of operation in minutes, and blood loss in ml. The scientists were interested in the factors that influence blood loss.

```
#### Example: Blood loss
thyroid <- read.table(text="
weight time blood_loss
44.3 105      503
40.6  80      490
69.0  86      471
43.7 112      505
50.3 109      482
50.2 100      490
35.4  96      513
52.2 120      464
", header=TRUE)

# show the structure of the data.frame
str(thyroid)

## 'data.frame': 8 obs. of  3 variables:
## $ weight    : num 44.3 40.6 69 43.7 50.3 50.2 35.4 52.2
## $ time      : int 105 80 86 112 109 100 96 120
## $ blood_loss: int 503 490 471 505 482 490 513 464

# display the data.frame
#thyroid
```

	weight	time	blood_loss
1	44.3	105	503
2	40.6	80	490
3	69.0	86	471
4	43.7	112	505
5	50.3	109	482
6	50.2	100	490
7	35.4	96	513
8	52.2	120	464

Below, we calculate the Pearson correlations between all pairs of variables (left), as well as the p-values (right) for testing whether the correlation is equal to zero.

```
p.corr <- cor(thyroid);
#p.corr

# initialize pvalue table with dim names
p.corr.pval <- p.corr;
for (i1 in 1:ncol(thyroid)) {
  for (i2 in 1:ncol(thyroid)) {
    p.corr.pval[i1,i2] <- cor.test(thyroid[, i1], thyroid[, i2])$p.value
  }
}
#p.corr.pval
```

	weight	time	blood_loss		weight	time	blood_loss
weight	1.000	-0.066	-0.772	weight	0.0000	0.8761	0.0247
time	-0.066	1.000	-0.107	time	0.8761	0.0000	0.8003
blood_loss	-0.772	-0.107	1.000	blood_loss	0.0247	0.8003	0.0000

Similarly, we calculate the Spearman (rank) correlation table (left), as well as the p-values (right) for testing whether the correlation is equal to zero.

```
s.corr <- cor(thyroid, method = "spearman");
#s.corr

# initialize pvalue table with dim names
s.corr.pval <- p.corr;
for (i1 in 1:ncol(thyroid)) {
  for (i2 in 1:ncol(thyroid)) {
    s.corr.pval[i1,i2] <- cor.test(thyroid[, i1], thyroid[, i2],
                                    method = "spearman")$p.value
  }
}
## Warning: Cannot compute exact p-value with ties
#s.corr.pval
```

	weight	time	blood_loss		weight	time	blood_loss
weight	1.000	0.286	-0.874	weight	0.0000	0.5008	0.0045
time	0.286	1.000	-0.156	time	0.5008	0.0000	0.7128
blood_loss	-0.874	-0.156	1.000	blood_loss	0.0045	0.7128	0.0000

Here are scatterplots for the original data and the ranks of the data using `ggpairs()` from the `GGally` package with `ggplot2`.

```
# Plot the data using ggplot
library(ggplot2)
library(GGally)

## Loading required package: reshape
##
## Attaching package: 'reshape'
## The following object is masked from 'package:plyr':
##
##     rename, round_any
## The following object is masked from 'package:reshape2':
##
##     colsplit, melt, recast
## The following object is masked from 'package:class':
##
##     condense
## Warning: replacing previous import 'rename' when loading 'reshape'
## Warning: replacing previous import 'round_any' when loading 'reshape'
p1 <- ggpairs(thyroid)
print(p1)

thyroid$rank_weight <- rank(thyroid$weight )
```

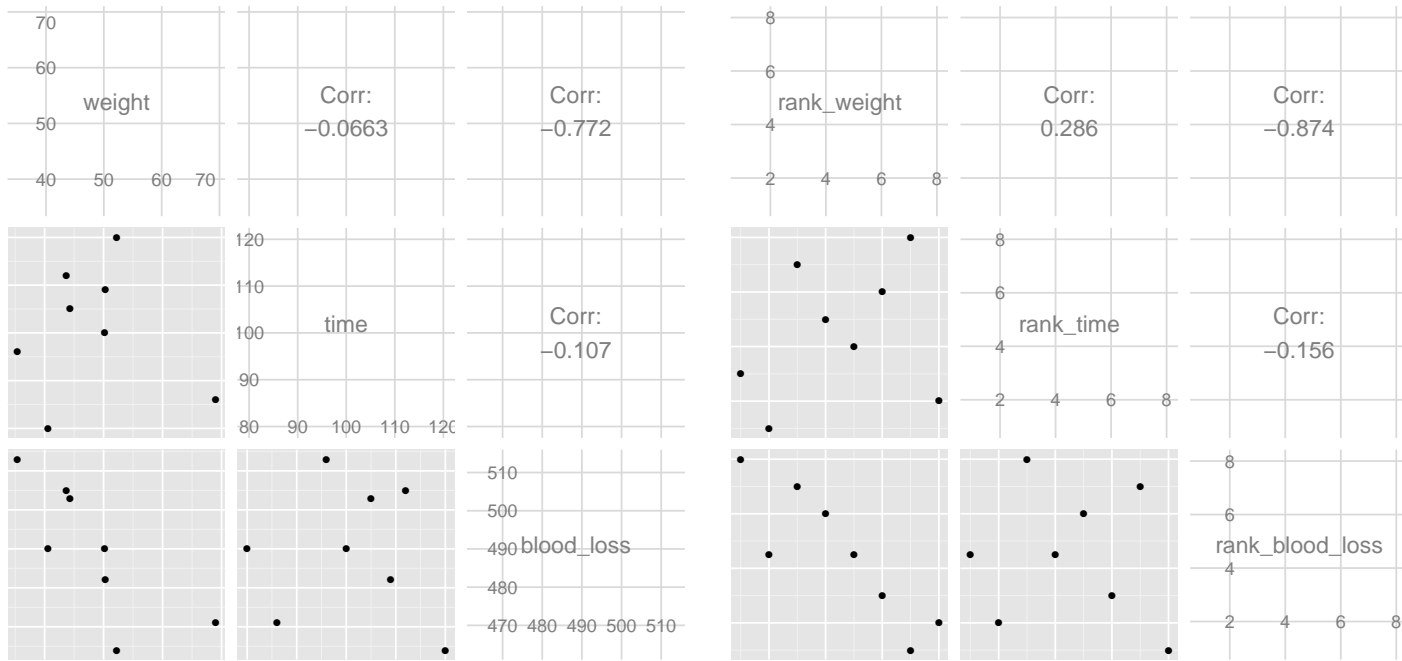
```

thyroid$rank_time      <- rank(thyroid$time      )
thyroid$rank_blood_loss <- rank(thyroid$blood_loss)

p2 <- ggpairs(thyroid[,4:6])
print(p2)

# detach package after use so reshape2 works (old reshape (v.1) conflicts)
detach("package:GGally", unload=TRUE)
detach("package:reshape", unload=TRUE)

```



Comments:

- (Pearson correlations). Blood loss tends to decrease linearly as weight increases, so r should be negative. The output gives $r = -0.77$. There is not much of a linear relationship between blood loss and time, so r should be close to 0. The output gives $r = -0.11$. Similarly, weight and time have a weak negative correlation, $r = -0.07$.
- The Pearson and Spearman correlations are fairly consistent here. Only the correlation between blood loss and weight is significantly different from zero at the $\alpha = 0.05$ level (the p-values are given below the correlations).
- (Spearman p-values) R gives the correct p-values. Calculating the p-value using the Pearson correlation on the ranks is not correct, strictly speaking.

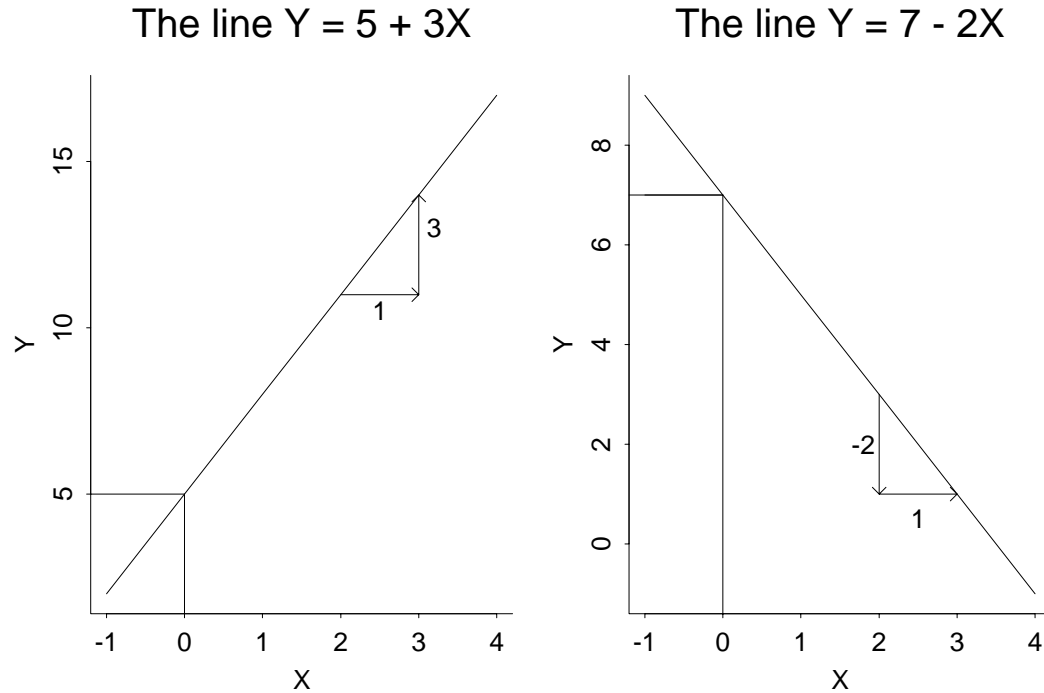


8.4 Simple Linear Regression

In linear regression, we are interested in developing a linear equation that best summarizes the relationship in a sample between the **response variable** Y and the **predictor variable** (or **independent variable**) X . The equation is also used to predict Y from X . The variables are not treated symmetrically in regression, but the appropriate choice for the response and predictor is usually apparent.

8.4.1 Linear Equation

If there is a perfect linear relationship between Y and X then $Y = \beta_0 + \beta_1 X$ for some β_0 and β_1 , where β_0 is the Y -intercept and β_1 is the slope of the line. Two plots of linear relationships are given below. The left plot has $\beta_0 = 5$ and $\beta_1 = 3$. The slope is positive, which indicates that Y increases linearly when X increases. The right plot has $\beta_0 = 7$ and $\beta_1 = -2$. The slope is negative, which indicates that Y decreases linearly when X increases.



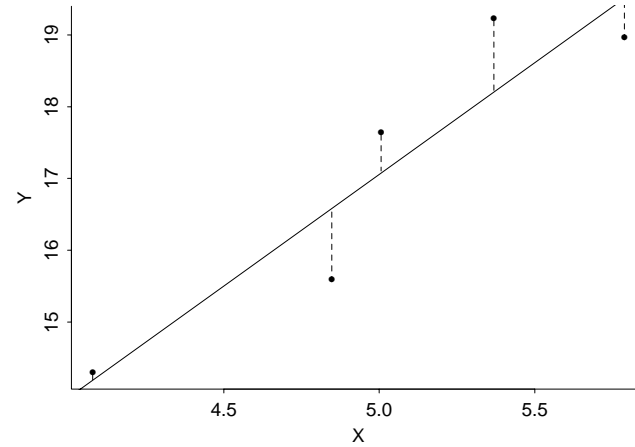
8.4.2 Least Squares

Data rarely, if ever, fall on a straight line. However, a straight line will often describe the **trend** for a set of data. Given a data set, (X_i, Y_i) , $i = 1, \dots, n$, with a **linear trend**, what linear equation “best” summarizes the observed relationship between Y and X ? There is no universally accepted definition of “best”, but many researchers accept the **Least Squares** line (LS line) as a reasonable summary.

Mathematically, the LS line chooses the values of β_0 and β_1 that minimize

$$\sum_{i=1}^n \{Y_i - (\beta_0 + \beta_1 X_i)\}^2$$

over all possible choices of β_0 and β_1 . These values can be obtained using calculus. Rather than worry about this calculation, note that the LS line makes the sum of squared (vertical) deviations between the responses Y_i and the line as small as possible, over all possible lines. The LS line goes through the mean point, (\bar{X}, \bar{Y}) , which is typically in the “the heart” of the data, and is often closely approximated by an eye-ball fit to the data.



The equation of the LS line is

$$\hat{y} = b_0 + b_1 X$$

where the intercept b_0 satisfies

$$b_0 = \bar{Y} - b_1 \bar{X}$$

and the slope is

$$b_1 = \frac{\sum_i(Y_i - \bar{Y})(X_i - \bar{X})}{\sum_i(X_i - \bar{X})^2} = r \frac{S_Y}{S_X}.$$

As before, r is the Pearson correlation between Y and X , whereas S_Y and S_X are the sample standard deviations for the Y and X samples, respectively. The **sign of the slope** and the **sign of the correlation** are **identical** (i.e., + correlation implies + slope).

Special symbols b_0 and b_1 identify the LS intercept and slope to distinguish the LS line from the generic line $Y = \beta_0 + \beta_1 X$. You should think of \hat{Y} as the **fitted value** at X , or the value of the LS line at X .

Fit a regression for the equation estimates from `summary()`. Note that we'll reuse the output of `lm()` over and over again.

```
lm.blood.wt <- lm(blood_loss ~ weight, data = thyroid)
lm.blood.wt

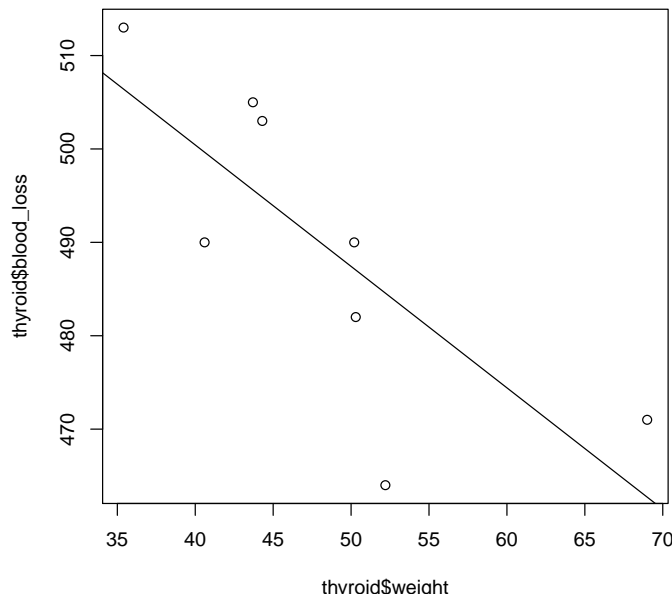
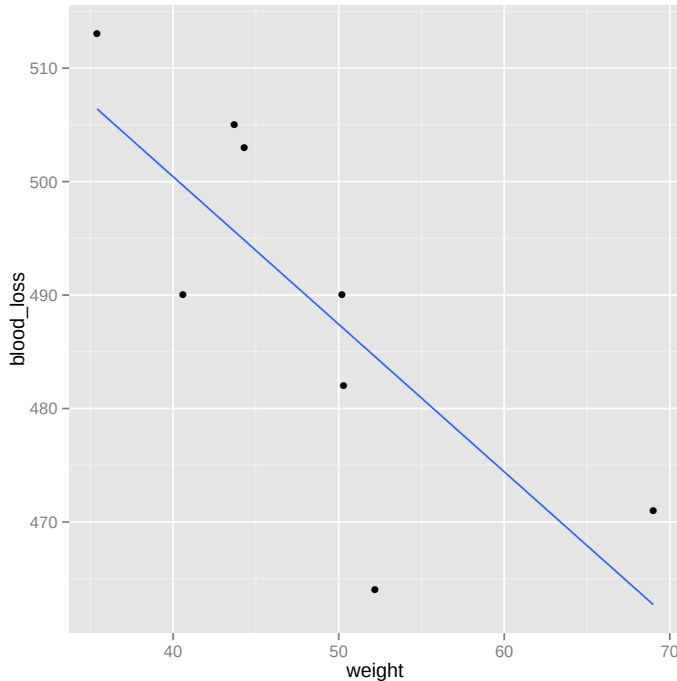
##
## Call:
## lm(formula = blood_loss ~ weight, data = thyroid)
##
## Coefficients:
## (Intercept)      weight
##      552.4        -1.3
#
# use summary() to get t-tests of parameters (slope, intercept)
summary(lm.blood.wt)

##
## Call:
## lm(formula = blood_loss ~ weight, data = thyroid)
##
## Residuals:
##    Min     1Q Median     3Q    Max 
## -20.57  -6.19   4.71   8.19   9.38 
##
## Coefficients:
##             Estimate Std. Error t value Pr(>|t|)    
## (Intercept) 552.442    21.441   25.77  2.3e-07 ***
## weight       -1.300     0.436   -2.98   0.025 *  
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 11.7 on 6 degrees of freedom
## Multiple R-squared:  0.597, Adjusted R-squared:  0.529 
## F-statistic: 8.88 on 1 and 6 DF,  p-value: 0.0247
```

Create a scatterplot with regression fit.

```
# ggplot: Plot the data with linear regression fit and confidence bands
library(ggplot2)
p <- ggplot(thyroid, aes(x = weight, y = blood_loss))
p <- p + geom_point()
p <- p + geom_smooth(method = lm, se = FALSE)
print(p)

# Base graphics: Plot the data with linear regression fit and confidence bands
# scatterplot
plot(thyroid$weight, thyroid$blood_loss)
# regression line from lm() fit
abline(lm.blood.wt)
```



For the **thyroid operation data** with Y = Blood loss in ml and X = Weight in kg , the LS line is $\hat{Y} = 552.44 - 1.30X$, or Predicted Blood Loss = $552.44 - 1.30 \text{ Weight}$. For an $86kg$ individual, the Predicted Blood Loss = $552.44 - 1.30 \times 86 = 440.64ml$.

The LS regression coefficients for this model are interpreted as follows. The intercept b_0 is the predicted blood loss for a $0\ kg$ individual. The intercept has no meaning here. The slope b_1 is the predicted increase in blood loss for each additional kg of weight. The slope is -1.30 , so the predicted *decrease* in blood loss is $1.30\ ml$ for each increase of $1\ kg$ in weight.

Any fitted linear relationship holds only approximately and does not necessarily extend outside the range of the data. In particular, nonsensical predicted blood losses of less than zero are obtained at very large weights outside the

range of data.

8.5 ANOVA Table for Regression

The LS line minimizes

$$\sum_{i=1}^n \{Y_i - (\beta_0 + \beta_1 X_i)\}^2$$

over all choices for β_0 and β_1 . Inserting the LS estimates b_0 and b_1 into this expression gives

$$\text{Residual Sums of Squares} = \sum_{i=1}^n \{Y_i - (b_0 + b_1 X_i)\}^2.$$

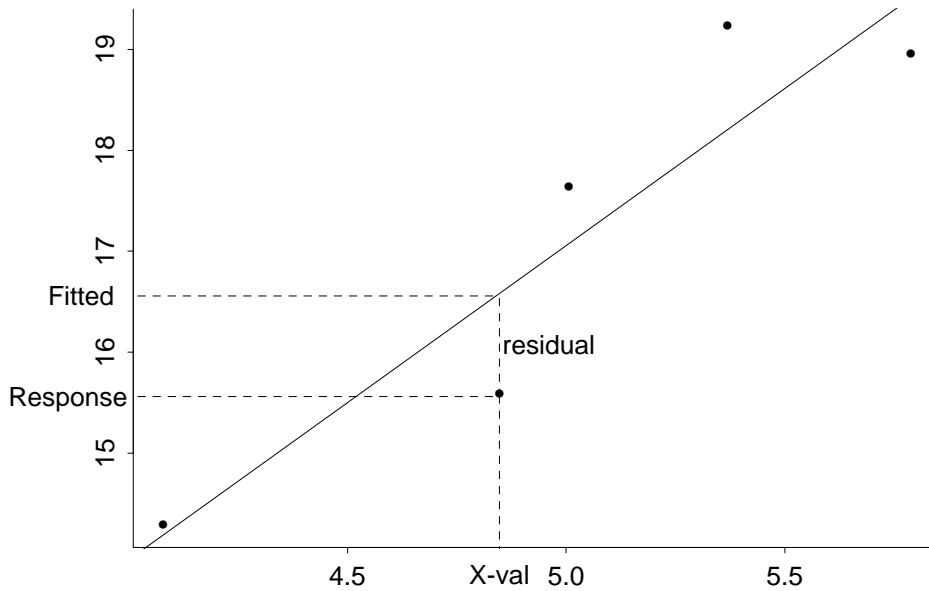
Several bits of notation are needed. Let

$$\hat{Y}_i = b_0 + b_1 X_i$$

be the **predicted** or fitted Y -value for an X -value of X_i and let $e_i = Y_i - \hat{Y}_i$. The fitted value \hat{Y}_i is the value of the LS line at X_i whereas the **residual** e_i is the distance that the observed response Y_i is from the LS line. Given this notation,

$$\text{Residual Sums of Squares} = \text{Res SS} = \sum_{i=1}^n (Y_i - \hat{y}_i)^2 = \sum_{i=1}^n e_i^2.$$

Here is a picture to clarify matters:



The Residual SS, or sum of squared residuals, is *small* if each \hat{Y}_i is *close to* Y_i (i.e., the line closely fits the data). It can be shown that

$$\text{Total SS in } Y = \sum_{i=1}^n (Y_i - \bar{Y})^2 \geq \text{Res SS} \geq 0.$$

Also define

$$\text{Regression SS} = \text{Reg SS} = \text{Total SS} - \text{Res SS} = b_1 \sum_{i=1}^n (Y_i - \bar{Y})(X_i - \bar{X}).$$

The Total SS measures the variability in the Y -sample. Note that

$$0 \leq \text{Regression SS} \leq \text{Total SS}.$$

The percentage of the variability in the Y -sample that is **explained by the linear relationship** between Y and X is

$$R^2 = \text{coefficient of determination} = \frac{\text{Reg SS}}{\text{Total SS}}.$$

Given the definitions of the Sums of Squares, we can show $0 \leq R^2 \leq 1$ and

$$R^2 = \text{square of Pearson correlation coefficient} = r^2.$$

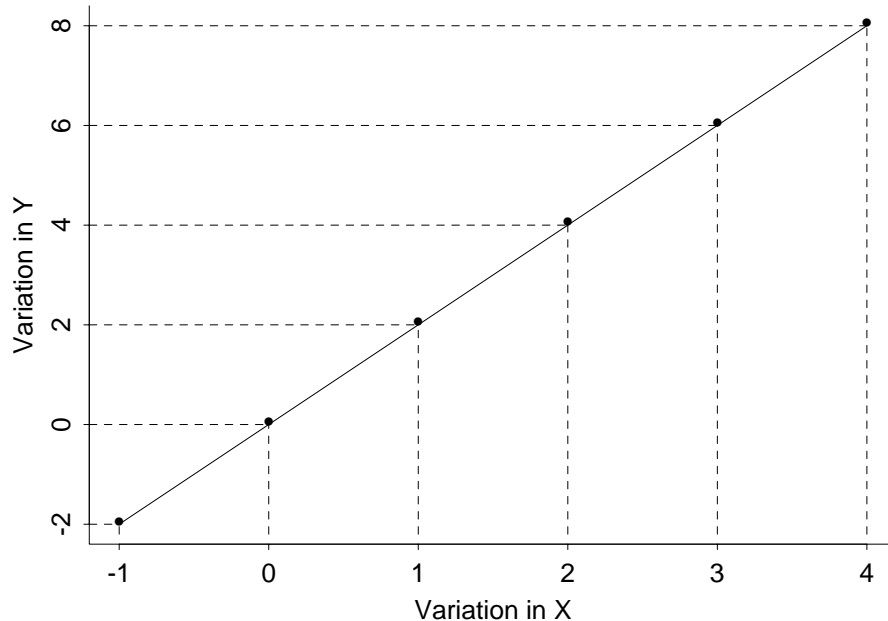
To understand the interpretation of R^2 , at least in two extreme cases, note that

$$\text{Reg SS} = \text{Total SS} \Leftrightarrow \text{Res SS} = 0$$

\Leftrightarrow all the data points fall on a straight line

\Leftrightarrow all the variability in Y is explained by the linear relationship
(which has variation)

$\Leftrightarrow R^2 = 1.$ (see the picture below)



Furthermore,

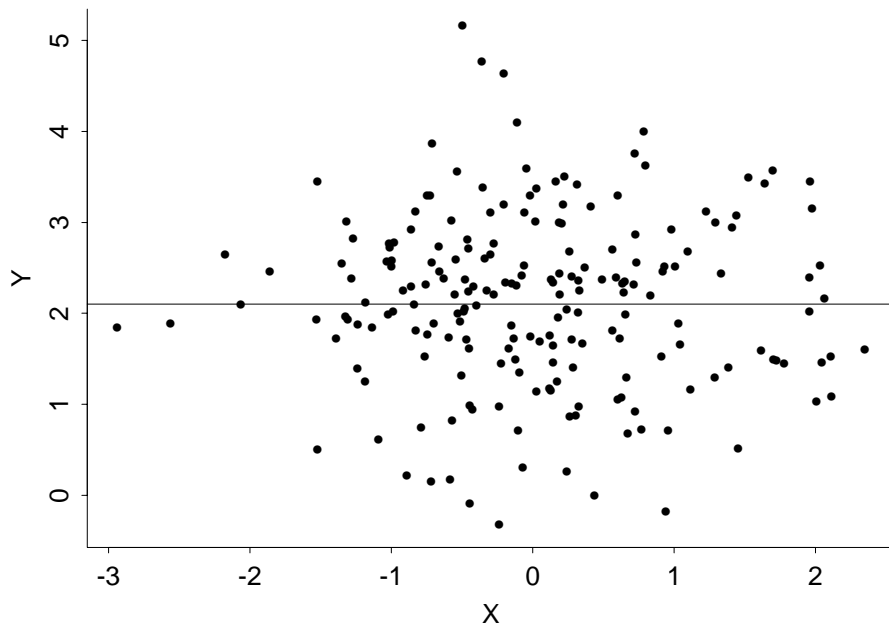
$$\text{Reg SS} = 0 \Leftrightarrow \text{Total SS} = \text{Res SS}$$

$\Leftrightarrow b_1 = 0$

\Leftrightarrow LS line is $\hat{Y} = \bar{Y}$

\Leftrightarrow none of the variability in Y is explained by a linear relationship

$\Leftrightarrow R^2 = 0.$



Each Sum of Squares has a corresponding df (degrees of freedom). The Sums of Squares and df are arranged in an analysis of variance (ANOVA) table:

Source	df	SS	MS
Regression	1	SSR	MSR
Residual (Error)	$n - 2$	SSE	MSE
Total	$n - 1$		

The Total df is $n - 1$. The Residual df is n minus the number of parameters (2) estimated by the LS line. The Regression df is the number of predictor variables (1) in the model. A Mean Square is always equal to the Sum of Squares divided by the df . Sometime the following notation is used for the Residual MS: $s_{Y|X}^2 = \text{Resid}(SS)/(n - 2)$.

```
# ANOVA table of the simple linear regression fit
anova(lm.blood.wt)

## Analysis of Variance Table
##
## Response: blood_loss
##           Df  Sum Sq  Mean Sq F value Pr(>F)
## weight      1  1207    1207     8.88  0.025 *
## Residuals   6   816     136
## ---
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

8.5.1 Brief discussion of the output for blood loss problem

1. Identify fitted line: Blood Loss = $552.44 - 1.30 \text{ Weight}$ (i.e., $b_0 = 552.44$ and $b_1 = -1.30$).

What is the line of best fit? What is the direction of the relationship?

2. Locate Analysis of Variance Table.

This tests the hypothesis $H_0 : \beta_j = 0, j = 0, 1, \dots, p$ (for all $p + 1$ beta coefficients), against H_A : not H_0 , i.e., at least one $\beta_j \neq 0$. More on this later.

3. Locate Parameter Estimates Table.

Given that not all the betas are zero from the ANOVA, which parameter betas are different from zero, and what are their estimated values and standard errors? More on this later.

4. Note that $R^2 = 0.5967 = (-0.77247)^2 = r^2$.

R^2 indicates the proportion of the variance explained by the regression model. This indicates the predictive ability of the model, and is *not* a indication of model fit.



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8.6 The regression model

The following statistical model is assumed as a means to provide error estimates for the LS line, regression coefficients, and predictions. Assume that the data (X_i, Y_i) , $i = 1, \dots, n$, are a sample of (X, Y) values from the population of interest, and

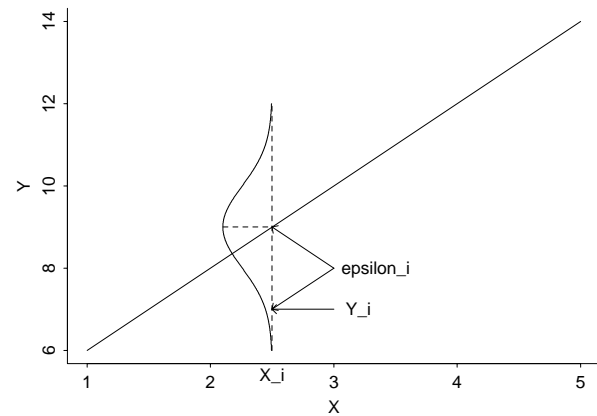
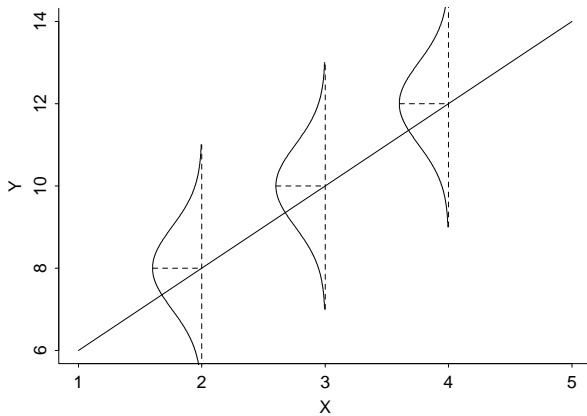
1. The mean in the population of all responses Y at a given X value (sometimes called $\mu_{Y|X}$) falls on a straight line, $\beta_0 + \beta_1 X$, called the population regression line.

2. The variation among responses Y at a given X value is the same for each X , and is denoted by $\sigma_{Y|X}^2$.
3. The population of responses Y at a given X is normally distributed.
4. The pairs (X_i, Y_i) are a random sample from the population. Alternatively, we can think that the X_i s were fixed by the experimenter, and that the Y_i are random responses at the selected predictor values.

The model is usually written in the form

$$Y_i = \beta_0 + \beta_1 X_i + \varepsilon_i$$

(i.e., Response = Mean Response + Residual), where the ε_i s are, by virtue of assumptions 2, 3, and 4, independent normal random variables with mean 0 and variance $\sigma_{Y|X}^2$. The following picture might help see this. Note that the population regression line is unknown, and is estimated from the data using the LS line.



8.6.1 Back to the Data

There are three unknown population parameters in the model: β_0 , β_1 and $\sigma_{Y|X}^2$. Given the data, the LS line

$$\hat{Y} = b_0 + b_1 X$$

estimates the population regression line $Y = \beta_0 + \beta_1 X$. The LS line is our best guess about the unknown population regression line. Here b_0 estimates the

intercept β_0 of the population regression line and b_1 estimates the slope β_1 of the population regression line.

The i^{th} **observed residual** $e_i = Y_i - \hat{Y}_i$, where $\hat{Y}_i = b_0 + b_1 X_i$ is the i^{th} **fitted value**, estimates the **unobservable residual** ε_i (ε_i is unobservable because β_0 and β_1 are unknown). The Residual MS from the ANOVA table is used to estimate $\sigma_{Y|X}^2$:

$$s_{Y|X}^2 = \text{Res SS} = \frac{\text{Res SS}}{\text{Res df}} = \frac{\sum_i (Y_i - \hat{Y}_i)^2}{n - 2}.$$

The denominator $df = n - 2$ is the number of observations minus the number of beta parameters in the model, i.e., β_0 and β_1 .

8.7 CI and tests for β_1

A CI for β_1 is given by $b_1 \pm t_{\text{crit}} SE_{b_1}$, where the standard error of b_1 under the model is

$$SE_{b_1} = \frac{s_{Y|X}}{\sqrt{\sum_i (X_i - \bar{X})^2}},$$

and where t_{crit} is the appropriate critical value for the desired CI level from a t -distribution with $df = \text{Res df}$.

To test $H_0 : \beta_1 = \beta_{10}$ (a given value) against $H_A : \beta_1 \neq \beta_{10}$, reject H_0 if $|t_s| \geq t_{\text{crit}}$, where

$$t_s = \frac{b_1 - \beta_{10}}{SE_{b_1}},$$

and t_{crit} is the t -critical value for a two-sided test, with the desired size and $df = \text{Res df}$. Alternatively, you can evaluate a p-value in the usual manner to make a decision about H_0 .

```
# CI for beta1
sum.lm.blood.wt <- summary(lm.blood.wt)
est.beta1 <- sum.lm.blood.wt$coefficients[2,1]
se.beta1 <- sum.lm.blood.wt$coefficients[2,2]
df.beta1 <- sum.lm.blood.wt$fstatistic[3]
t.crit <- qt(1-0.05/2, df.beta1)
CI.lower <- est.beta1 - t.crit * se.beta1
CI.upper <- est.beta1 + t.crit * se.beta1
c(CI.lower, CI.upper)
## [1] -2.3682 -0.2325
```

The parameter estimates table gives the standard error, t -statistic, and p-value for testing $H_0 : \beta_1 = 0$. Analogous summaries are given for the intercept, β_0 , but these are typically of less interest.

8.7.1 Testing $\beta_1 = 0$

Assuming the mean relationship is linear, consider testing $H_0 : \beta_1 = 0$ against $H_A : \beta_1 \neq 0$. This test can be conducted using a t -statistic, as outlined above, or with an ANOVA F -test, as outlined below.

For the analysis of variance (ANOVA) F -test, compute

$$F_s = \frac{\text{Reg MS}}{\text{Res MS}}$$

and reject H_0 when F_s exceeds the critical value (for the desired size test) from an F -table with numerator $df = 1$ and denominator $df = n - 2$ (see `qf()`). The hypothesis of zero slope (or no relationship) is rejected when F_s is large, which happens when a significant portion of the variation in Y is explained by the linear relationship with X .

The p-values from the t -test and the F -test are always equal. Furthermore this p-value is equal to the p-value for testing no correlation between Y and X , using the t -test described earlier. Is this important, obvious, or disconcerting?

8.8 A CI for the population regression line

I can not overemphasize the **power** of the regression model. The model allows you to estimate the mean response at any X value in the range for which the model is reasonable, even if little or no data is observed at that location.

We estimate the mean population response among individuals with $X = X_p$

$$\mu_p = \beta_0 + \beta_1 X_p,$$

with the fitted value, or the value of the least squares line at X_p :

$$\hat{Y}_p = b_0 + b_1 X_p.$$

X_p is not necessarily one of the observed X_i s in the data. To get a CI for μ_p , use $\hat{Y}_p \pm t_{\text{crit}}SE(\hat{Y}_p)$, where the standard error of \hat{Y}_p is

$$SE(\hat{Y}_p) = s_{Y|X} \sqrt{\frac{1}{n} + \frac{(X_p - \bar{X})^2}{\sum_i (X_i - \bar{X})^2}}.$$

The t -critical value is identical to that used in the subsection on CI for β_1 .

8.8.1 CI for predictions

Suppose a future individual (i.e., someone not used to compute the LS line) has $X = X_p$. The best prediction for the response Y of this individual is the value of the least squares line at X_p :

$$\hat{Y}_p = b_0 + b_1 X_p.$$

To get a CI (prediction interval) for an individual response, use $\hat{Y}_p \pm t_{\text{crit}}SE_{\text{pred}}(\hat{Y}_p)$, where

$$SE_{\text{pred}}(\hat{Y}_p) = s_{Y|X} \sqrt{1 + \frac{1}{n} + \frac{(X_p - \bar{X})^2}{\sum_i (X_i - \bar{X})^2}},$$

and t_{crit} is identical to the critical value used for a CI on β_1 . The prediction variance has two parts: (1) the 1 indicates the variability associated with the data around the mean (regression line), and (2) the rest is the variability associated with estimating the mean.

For example, in the blood loss problem you may want to estimate the blood loss for an $50kg$ individual, and to get a CI for this prediction. This problem is different from computing a CI for the mean blood loss of all $50kg$ individuals!

```
# CI for the mean and PI for a new observation at weight=50
predict(lm.blood.wt, data.frame(weight=50), interval = "confidence", level = 0.95)
##      fit     lwr     upr
## 1 487.4 477.2 497.7

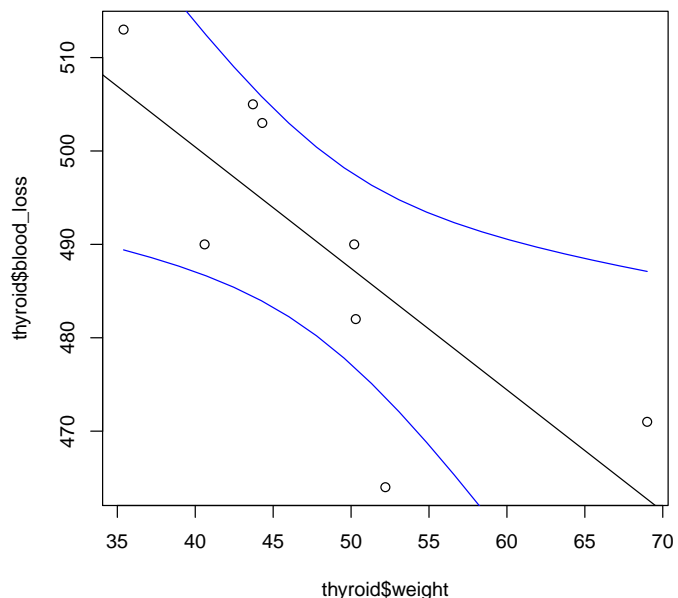
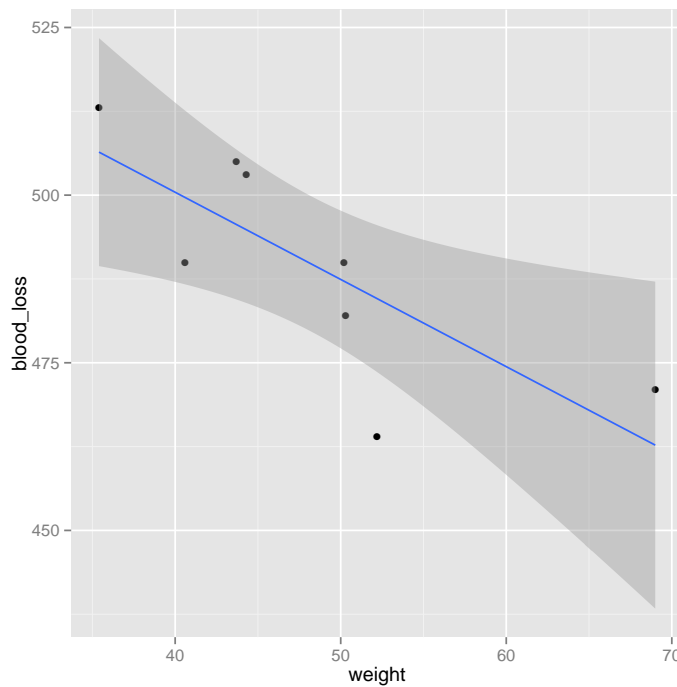
predict(lm.blood.wt, data.frame(weight=50), interval = "prediction", level = 0.95)
##      fit     lwr     upr
## 1 487.4 457.1 517.8
```

Comments

1. The prediction interval is wider than the CI for the mean response. This is reasonable because you are less confident in predicting an individual response than the mean response for all individuals.
2. The CI for the mean response and the prediction interval for an individual response become wider as X_p moves away from \bar{X} . That is, you get a more sensitive CI and prediction interval for X_p s near the center of the data.
3. In plots below include confidence and prediction bands along with the fitted LS line.

```
# ggplot: Plot the data with linear regression fit and confidence bands
library(ggplot2)
p <- ggplot(thyroid, aes(x = weight, y = blood_loss))
p <- p + geom_point()
p <- p + geom_smooth(method = lm, se = TRUE)
print(p)

# Base graphics: Plot the data with linear regression fit and confidence bands
# scatterplot
plot(thyroid$weight, thyroid$blood_loss)
# regression line from lm() fit
abline(lm.blood.wt)
# x values of weight for predictions of confidence bands
x.pred <- data.frame(weight = seq(min(thyroid$weight), max(thyroid$weight),
                                 length = 20))
# draw upper and lower confidence bands
lines(x.pred$weight, predict(lm.blood.wt, x.pred,
                           interval = "confidence")[, "upr"], col = "blue")
lines(x.pred$weight, predict(lm.blood.wt, x.pred,
                           interval = "confidence")[, "lwr"], col = "blue")
```



8.8.2 A further look at the blood loss data

- The LS line is: Predicted Blood Loss = $552.442 - 1.30 \text{ Weight}$.
- The R^2 is 0.597 (i.e., 59.7%).
- The F -statistic for testing $H_0 : \beta_1 = 0$ is $F_{obs} = 8.88$ with a p-value=0.025. The Error MS is $s^2_{Y|X} = 136.0$; see ANOVA table.
- The Parameter Estimates table gives b_0 and b_1 , their standard errors, and t -statistics and p-values for testing $H_0 : \beta_0 = 0$ and $H_0 : \beta_1 = 0$. The t -test and F -test p-values for testing that the slope is zero are identical. We could calculate a 95% CI for β_0 and β_1 . If we did so (using the t critical value) we find we are 95% confident that the slope of the population regression line is between -2.37 and -0.23 .
- Suppose we are interested in estimating the average blood loss among all 50kg individuals. The estimated mean blood loss is $552.442 - 1.30033 \times 50 = 487.43$. Reading off the plot, we are 95% confident that the mean blood loss of all 50kg individuals is between (approximately) 477 and 498 ml. A 95% prediction interval for the blood loss of a single 50 kg person is less precise (about 457 to 518 ml).

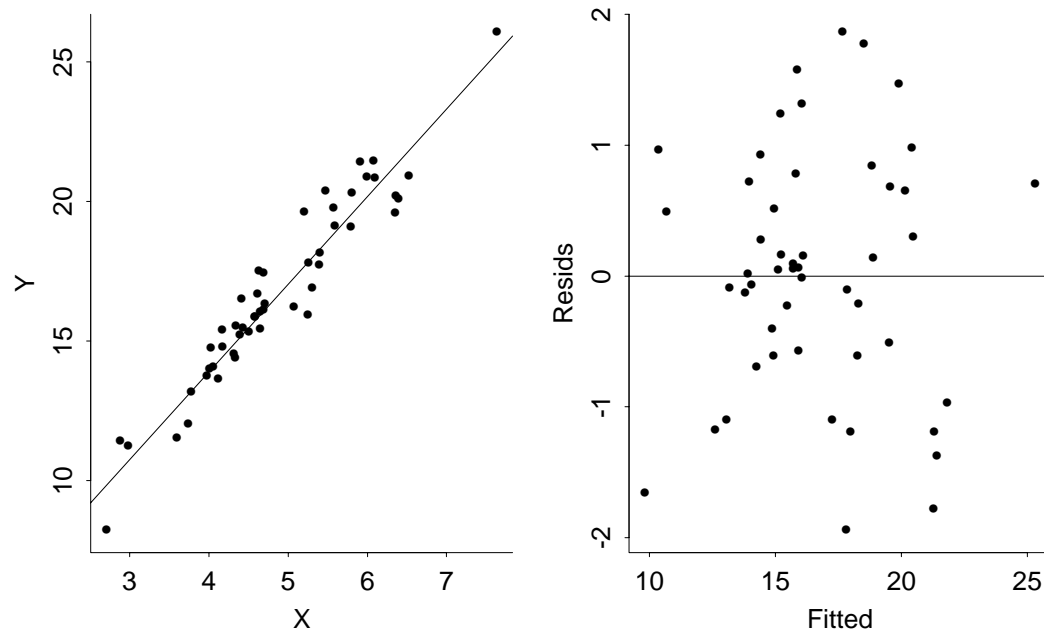
As a summary we might say that weight is important for explaining the variation in blood loss. In particular, the estimated slope of the least squares

line (Predicted Blood loss = $552.442 - 1.30 \text{ Weight}$) is significantly different from zero ($p\text{-value} = 0.0247$), with weight explaining approximately 60% (59.7%) of the variation in blood loss for this sample of 8 thyroid operation patients.

8.9 Checking the regression model

A regression analysis is never complete until the assumptions of the model have been checked. In addition, you need to evaluate whether individual observations, or groups of observations, are unduly influencing the analysis. A first step in any analysis is to plot the data. The plot provides information on the linearity and constant variance assumption. For example, the data plot below shows a linear relationship with roughly constant variance.

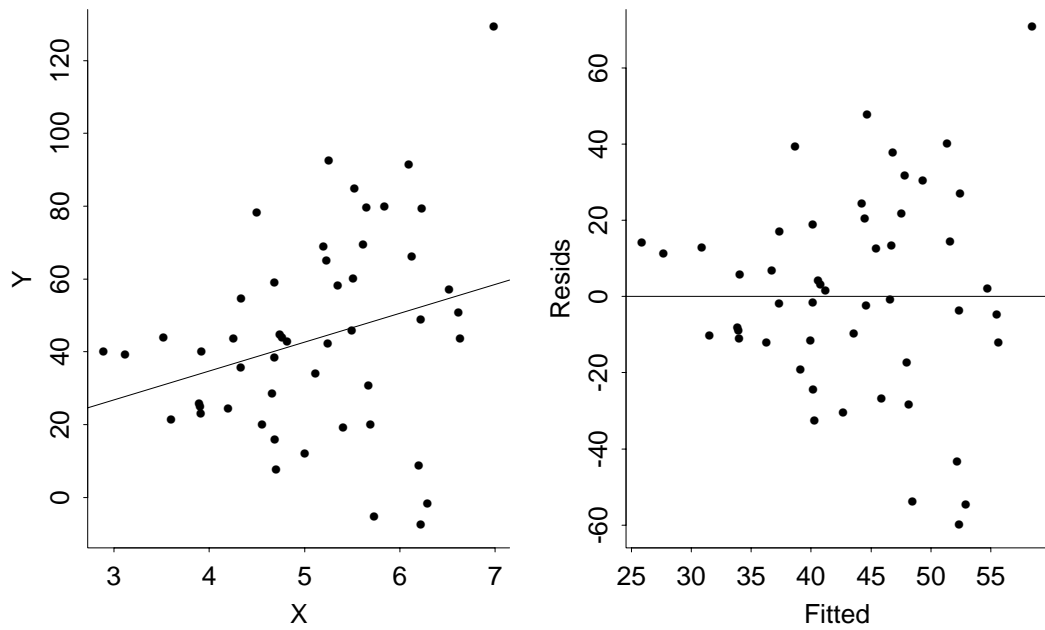
In addition to plotting the data, a variety of methods for assessing model adequacy are based on plots of the residuals, $e_i = Y_i - \hat{Y}_i$ (i.e., Observed–Fitted values). For example, an option is to plot the e_i against the fitted values \hat{Y}_i , as given below. This residual plot should exhibit no systematic dependence of the sign or the magnitude of the residuals on the fitted values.



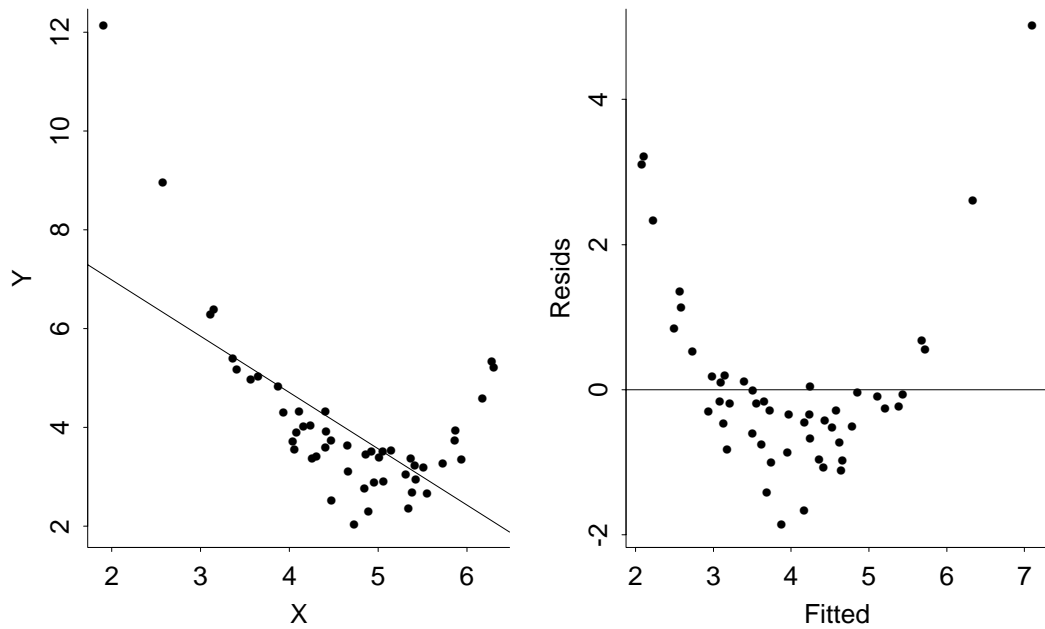
The real power of this plot is with multiple predictor problems (multiple

regression). For simple linear regression, the information in this plot is similar to the information in the original data plot, except that the residual plot eliminates the effect of the trend on your perceptions of model adequacy.

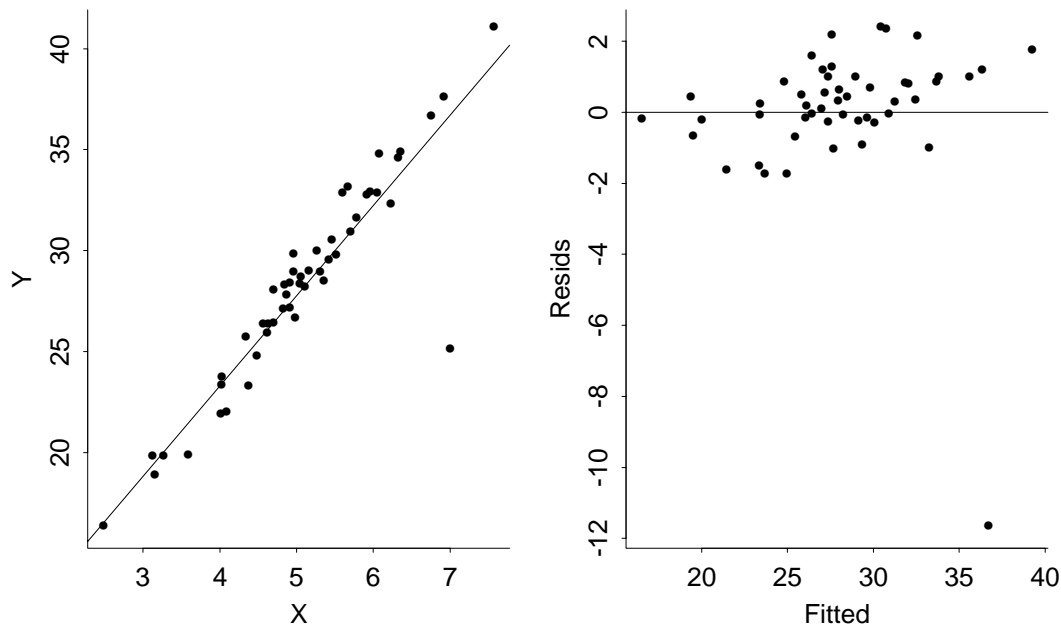
The following plots show how inadequacies in the data plot appear in a residual plot. The first plot shows a roughly linear relationship between Y and X with non-constant variance. The residual plot shows a megaphone shape rather than the ideal horizontal band. A possible remedy is a **weighted least squares** analysis to handle the non-constant variance, or to transform Y to stabilize the variance. Transforming the data may destroy the linearity.



The second plot shows a nonlinear relationship between Y and X . The residual plot shows a systematic dependence of the sign of the residual on the fitted value. A possible remedy is to transform the data.



The last plot shows an outlier. This point has a large residual. A sensible approach is to refit the model after deleting the case and see if any conclusions change.



8.9.1 Checking normality

The normality assumption can be evaluated with a boxplot or a normal quantile plot of the residuals. A formal test of normality using the residuals can be computed as discussed earlier this semester.

8.9.2 Checking independence

Diagnosing dependence among observations usually requires some understanding of the mechanism that generated the data. There are a variety of graphical and inferential tools for checking independence for data collected over time (called a time series). The easiest thing to do is plot the r_i against time index and look for any suggestive patterns.

8.9.3 Outliers

Outliers are observations that are poorly fitted by the regression model. The response for an outlier is far from the fitted line, so outliers have large positive or negative values of the residual e_i .

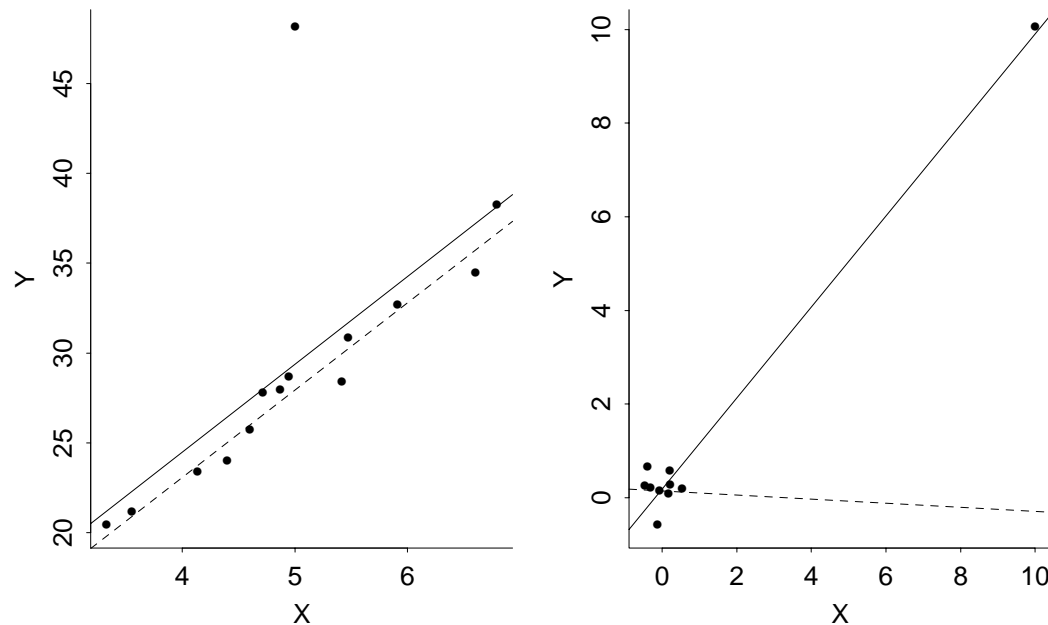
What do you do with outliers? Outliers may be due to incorrect recordings of the data or failure of the measuring device, or indications or a change in the mean or variance structure for one or more cases. Incorrect recordings should be fixed if possible, but otherwise deleted from the analysis.

Routine deletion of outliers from the analysis is not recommended. This practice can have a dramatic effect on the fit of the model and the perceived precision of parameter estimates and predictions. Analysts who routinely omit outliers without cause tend to overstate the significance of their findings and get a false sense of precision in their estimates and predictions. At the very least, a data analyst should repeat the analysis with and without the outliers to see whether any substantive conclusions are changed.

8.9.4 Influential observations

Certain data points can play a very important role in determining the position of the LS line. These data points may or may not be outliers. For example, the

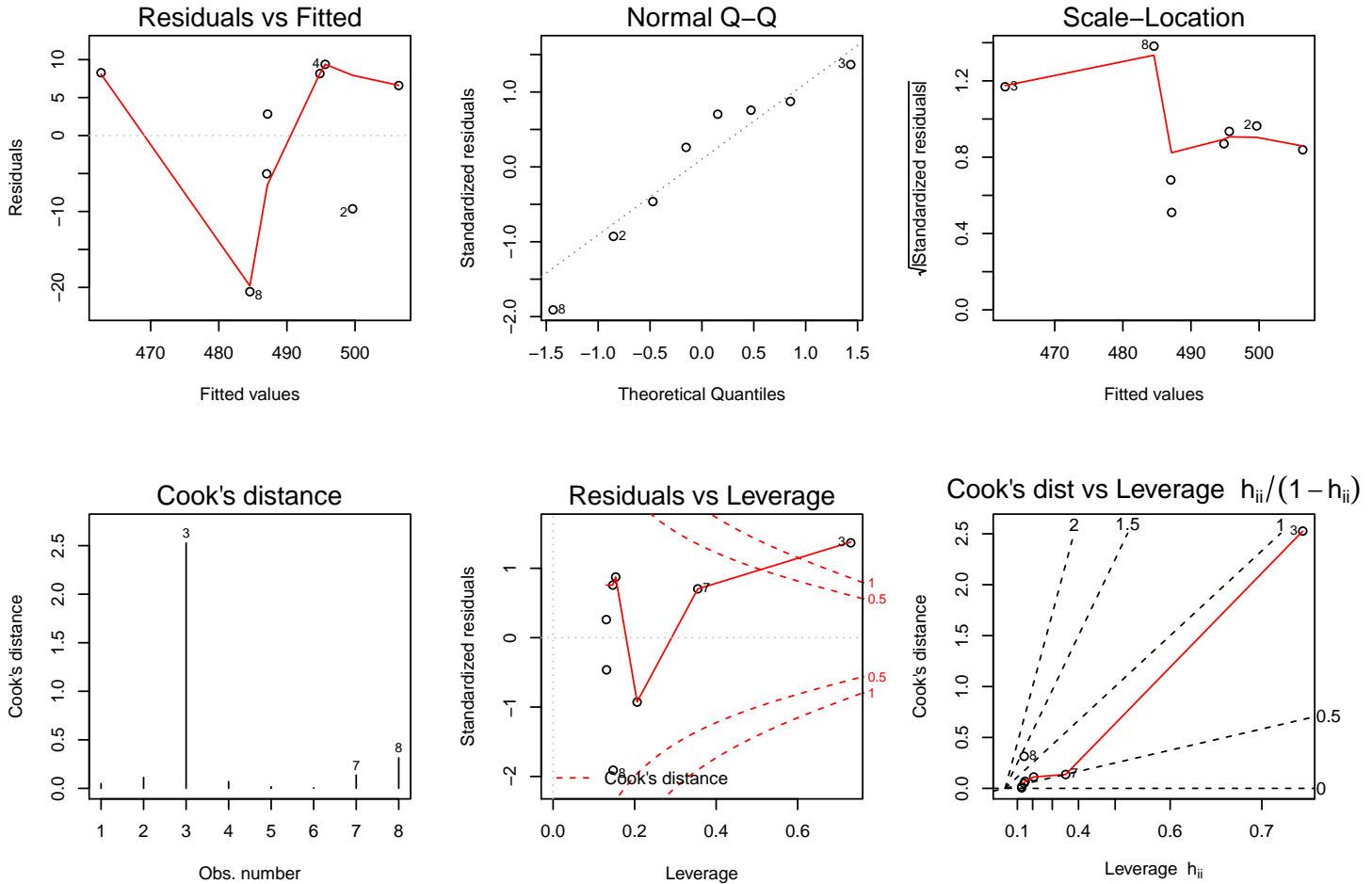
observation with $Y > 45$ in the first plot below is an outlier relative to the LS fit. The extreme observation in the second plot has a very small e_i . Both points are highly **influential observations** — the LS line changes dramatically when these observations are deleted. The influential observation in the second plot is not an outlier because its presence in the analysis determines that the LS line will essentially pass through it! In these plots the solid line is the LS line from the full data set, whereas the dashed line is the LS line after omitting the unusual point.



There are well-defined measures of the influence that individual cases have on the LS line. On the separate output I calculated Cook's D — large values indicate influential values. Which observations are most influential according to this measure? For simple linear regression most influential cases can be easily spotted by carefully looking at the data plot. If you identify cases that you suspect might be influential, you should hold them out (individually) and see if any important conclusions change. If so, you need to think hard about whether the cases should be included or excluded from the analysis.

In the next chapter we'll discuss model assessment, including some of the measures seen in the plots below.

```
# plot diagnostics
par(mfrow=c(2,3))
plot(lm.blood.wt, which = 1:6)
```



■ CLICKER Qs — Outliers STT.02.04.040 ■

8.10 Model Checking and Regression Diagnostics

8.10.1 Introduction

The simple linear regression model is usually written as

$$Y_i = \beta_0 + \beta_1 X_i + \varepsilon_i$$

where the ε_i s are independent normal random variables with mean 0 and variance σ^2 . The model implies (1) The average Y -value at a given X -value is linearly related to X . (2) The variation in responses Y at a given X value is constant. (3) The population of responses Y at a given X is normally distributed. (4) The observed data are a random sample.

A regression analysis is never complete until these assumptions have been checked. In addition, you need to evaluate whether individual observations, or groups of observations, are unduly influencing the analysis. A first step in any analysis is to plot the data. The plot provides information on the linearity and constant variance assumption.

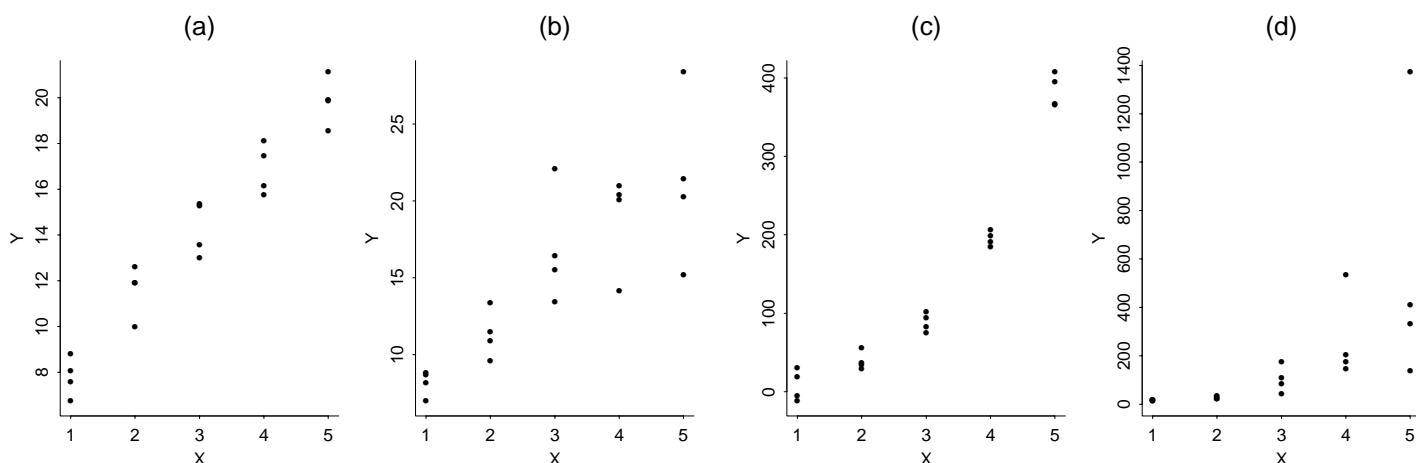


Figure (a) is the only plot that is consistent with the assumptions. The plot shows a linear relationship with constant variance. The other figures show one or more deviations. Figure (b) shows a linear relationship but the variability increases as the mean level increases. In Figure (c) we see a nonlinear relationship with constant variance, whereas (d) exhibits a nonlinear relationship with non-constant variance.

In many examples, nonlinearity or non-constant variability can be addressed by **transforming** Y or X (or both), or by fitting **polynomial models**. These issues will be addressed later.

8.10.2 Residual Analysis

A variety of methods for assessing model adequacy are based on the observed residuals,

$$e_i = Y_i - \hat{Y}_i \quad \text{i.e., Observed} - \text{Fitted values.}$$

The residuals are usually plotted in various ways to assess potential inadequacies. The observed residuals have different variances, depending on X_i . Recall that the standard error of \hat{Y}_i (and therefore e_i) is

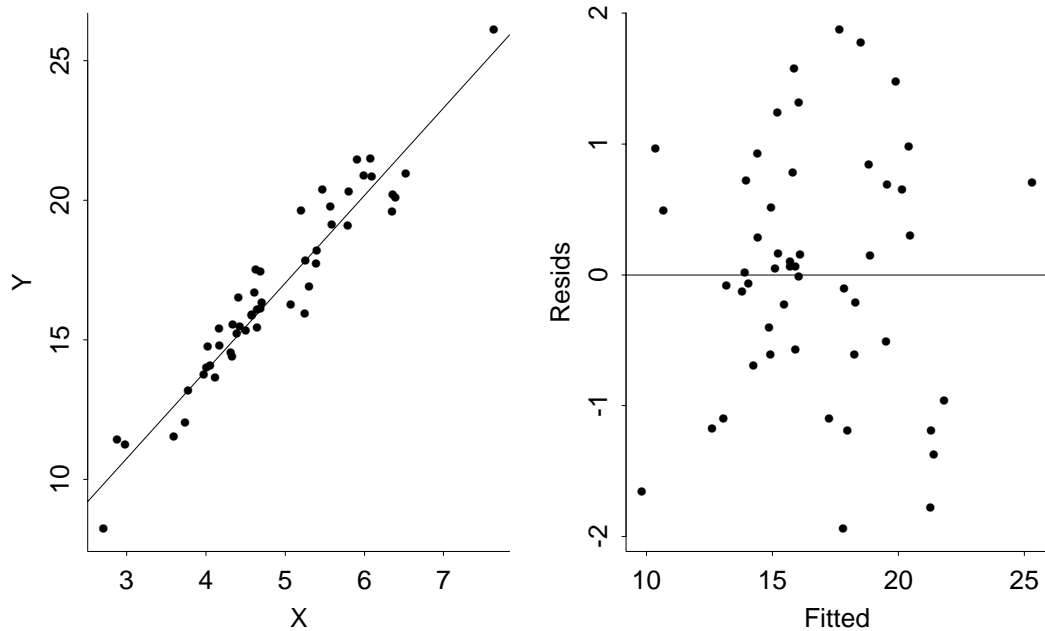
$$\text{SE}(\hat{Y}_i) = \text{SE}(e_i) = s_{Y|X} \sqrt{\frac{1}{n} + \frac{(X_i - \bar{X})^2}{\sum_j (X_j - \bar{X})^2}}.$$

So many statisticians prefer to plot the **studentized residuals** (sometimes called the standardized residuals)

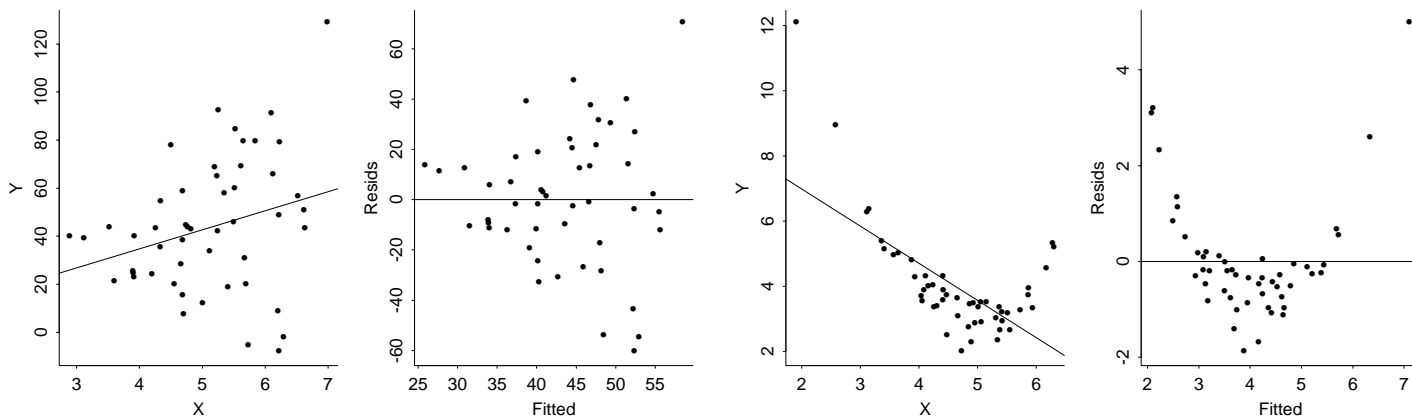
$$r_i = \frac{e_i}{\text{SE}(e_i)}.$$

The studentized residuals have a constant variance of 1 (approximately). I will focus on diagnostic methods using the studentized residuals.

A plot of the studentized residuals r_i against the fitted values \hat{Y}_i often reveals inadequacies with the model. The real power of this plot is with multiple predictor problems (multiple regression). The information contained in this plot with simple linear regression is similar to the information contained in the original data plot, except it is scaled better and eliminates the effect of the trend on your perceptions of model adequacy. The residual plot should exhibit no systematic dependence of the sign or the magnitude of the residuals on the fitted values:



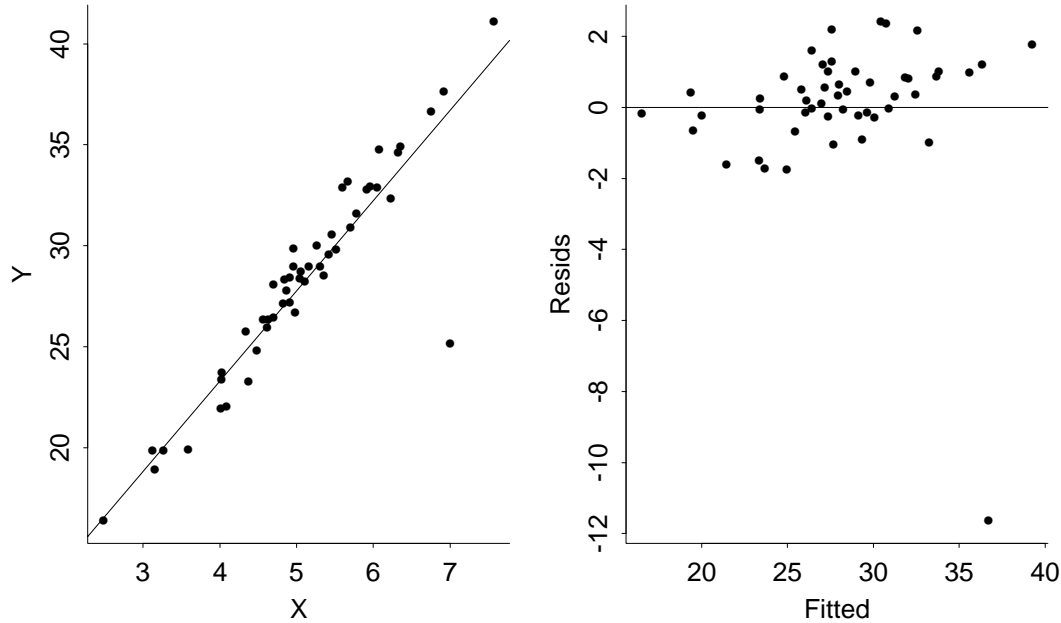
The following sequence of plots show how inadequacies in the data plot appear in a residual plot. The first plot shows a roughly linear relationship between Y and X with non-constant variance. The residual plot shows a megaphone shape rather than the ideal horizontal band. A possible remedy is a **weighted least squares** analysis to handle the non-constant variance (see end of chapter for an example), or to transform Y to stabilize the variance. Transforming the data may destroy the linearity.



The plot above shows a nonlinear relationship between Y and X . The

residual plot shows a systematic dependence of the sign of the residual on the fitted value. Possible remedies were mentioned earlier.

The plot below shows an **outlier**. This case has a large residual and large studentized residual. A sensible approach here is to refit the model after holding out the case to see if any conclusions change.



Nonconstant variance vs sample size Because more extreme observations are more likely to occur with larger sample sizes, sometimes when sample size depends on X it can appear as nonconstant variance. Below sample sizes are either all 25 or (3, 5, 10, 25, 125), and standard deviations are either all 1 or (0.1, 0.5, 1, 1.5, 3). Note that constant variance and different sample sizes appears as though it has nonconstant variance.

```
#### Nonconstant variance vs sample size
dat.var.sam <- function(n, s) {
  dat <- data.frame(
    x = c(rep(0, n[1]),
          rep(1, n[2]),
          rep(2, n[3]),
          rep(3, n[4]),
          rep(4, n[5])),
    y = c(rnorm(n[1], mean = 0, sd = s[1]),
          rnorm(n[2], mean = 0, sd = s[2]),
```

```
        rnorm(n[3], mean = 0, sd = s[3]),
        rnorm(n[4], mean = 0, sd = s[4]),
        rnorm(n[5], mean = 0, sd = s[5]))
    )
  return(dat)
}

library(ggplot2)

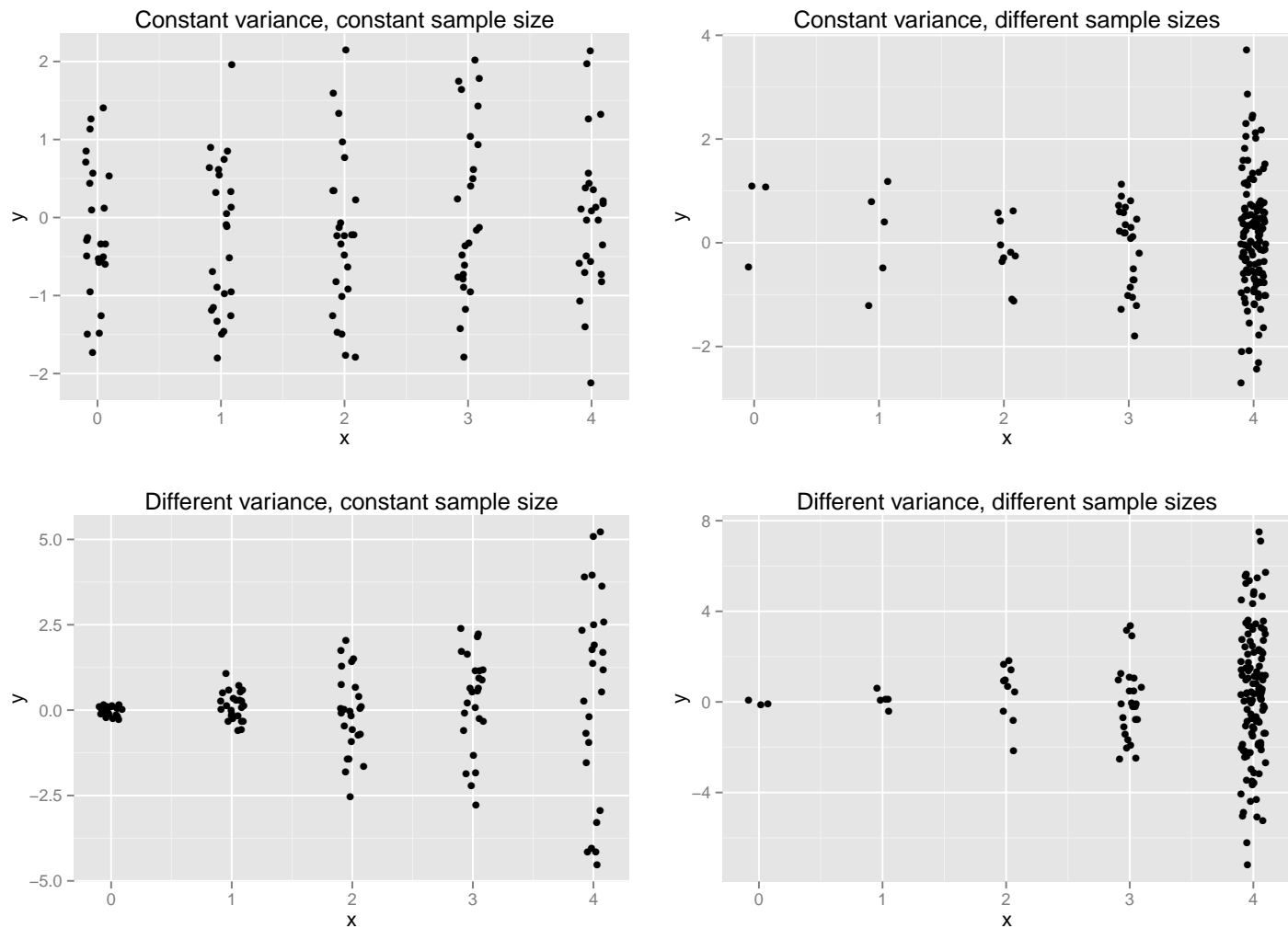
n <- c(25, 25, 25, 25, 25)
s <- c(1, 1, 1, 1, 1)
dat <- dat.var.sam(n, s)
p1 <- ggplot(dat, aes(x = x, y = y))
p1 <- p1 + geom_point(position = position_jitter(width = 0.1))
p1 <- p1 + labs(title = "Constant variance, constant sample size")
#print(p1)

n <- c(3, 5, 10, 25, 125)
s <- c(1, 1, 1, 1, 1)
dat <- dat.var.sam(n, s)
p2 <- ggplot(dat, aes(x = x, y = y))
p2 <- p2 + geom_point(position = position_jitter(width = 0.1))
p2 <- p2 + labs(title = "Constant variance, different sample sizes")
#print(p2)

n <- c(25, 25, 25, 25, 25)
s <- c(0.1, 0.5, 1, 1.5, 3)
dat <- dat.var.sam(n, s)
p3 <- ggplot(dat, aes(x = x, y = y))
p3 <- p3 + geom_point(position = position_jitter(width = 0.1))
p3 <- p3 + labs(title = "Different variance, constant sample size")
#print(p3)

n <- c(3, 5, 10, 25, 125)
s <- c(0.1, 0.5, 1, 1.5, 3)
dat <- dat.var.sam(n, s)
p4 <- ggplot(dat, aes(x = x, y = y))
p4 <- p4 + geom_point(position = position_jitter(width = 0.1))
p4 <- p4 + labs(title = "Different variance, different sample sizes")
#print(p4)

library(gridExtra)
grid.arrange(p1, p2, p3, p4, nrow=2, ncol=2)
```



8.10.3 Checking Normality

The normality assumption for the ε_i s can be evaluated visually with a boxplot or a normal probability plot (rankit plot) of the r_i , or formally with a Shapiro-Wilk test. The normal probability plot often highlights **outliers**, or poorly fitted cases. If an outlier is held out of the data and a new analysis is performed, the resulting normal scores plot may be roughly linear, but often will show a short-tailed distribution. (Why?).

You must interpret regression tests and CI with caution with non-normal data. Statisticians developed robust regression methods for non-normal data which are available in R packages.

8.10.4 Checking Independence

Diagnosing dependence among observations requires an understanding of the data collection process. There are a variety of graphical and inferential tools for checking independence for data collected over time (called a time series). The easiest check is to plot the r_i against time index and look for any suggestive patterns.

8.10.5 Outliers

Outliers are observations that are poorly fitted by the regression model. The response for an outlier is far from the fitted line, so outliers have large positive or negative values of the studentized residual r_i . Usually, $|r_i| > 2$ is considered large. Outliers are often highlighted in residual plots.

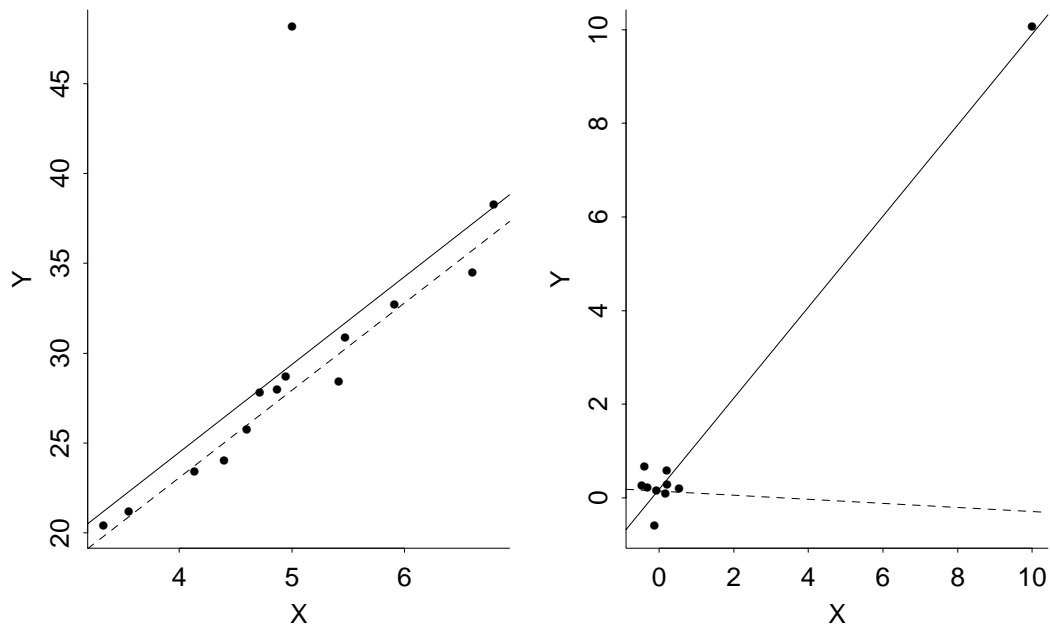
What do you do with outliers? Outliers may be due to incorrect recordings of the data or failure of the measuring device, or indications or a change in the mean or variance structure for one or more cases. Incorrect recordings should be fixed if possible, but otherwise deleted from the analysis.

Routine deletion of outliers from the analysis is not recommended. This practice can have a dramatic effect on the fit of the model and the perceived precision of parameter estimates and predictions. Analysts who routinely omit outliers without cause tend to overstate the significance of their findings and get a false sense of precision in their estimates and predictions. To assess effects of outliers, a data analyst should repeat the analysis holding out the outliers to see whether any substantive conclusions are changed. Very often the only real effect of an outlier is to inflate MSE and hence make p-values a little larger and CIs a little wider than necessary, but without substantively changing conclusions. They can completely mask underlying patterns, however.

8.10.6 Influential Observations

Certain data points can play an important role in determining the position of the LS line. These data points may or may not be outliers. For example, the observation with $Y > 45$ in the first plot is an outlier relative to the LS fit.

The extreme observation in the second plot has a very small r_i . Both points are highly **influential observations** - the LS line changes dramatically when these observations are held out. The influential observation in the second plot is not an outlier because its presence in the analysis determines that the LS line will essentially pass through it! In these plots the solid line is the LS line from the full data set, whereas the dashed line is the LS line after omitting the unusual point.



Dennis Cook developed a measure of the impact that individual cases have on the placement of the LS line. His measure, called **Cook's distance** or Cook's D, provides a summary of how far the LS line changes when each individual point is held out (one at a time) from the analysis. The case with the largest D has the greatest impact on the placement of the LS line. However, the actual influence of this case may be small. In the plots above, the observations I focussed on have the largest Cook's D s.

A simple, but not unique, expression for Cook's distance for the j^{th} case is

$$D_j \propto \sum_i (\hat{Y}_i - \hat{Y}_{i[-j]})^2,$$

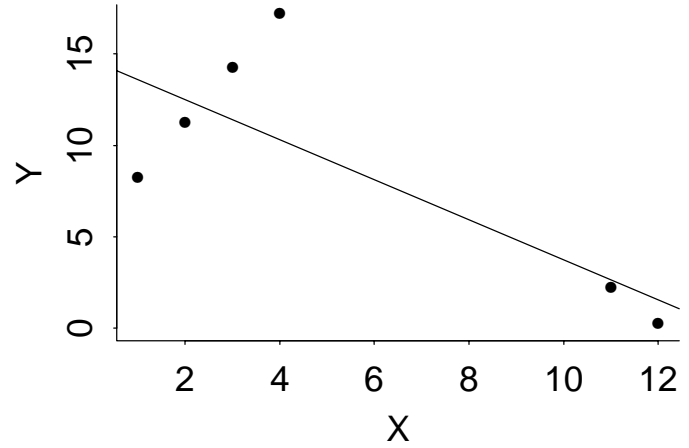
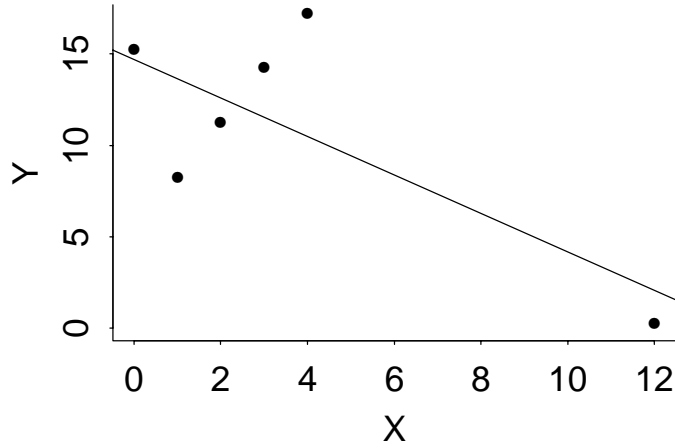
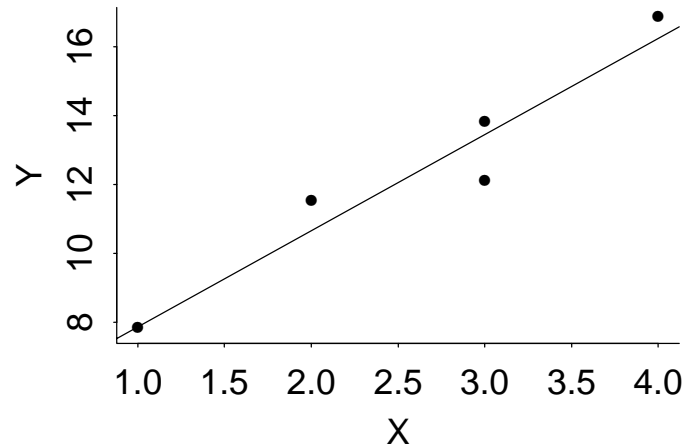
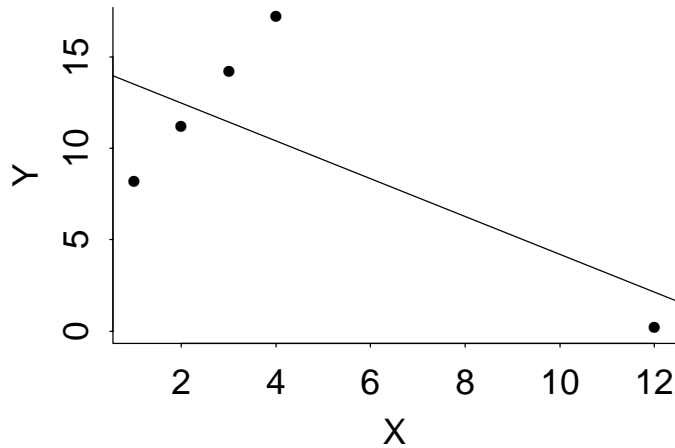
where $\hat{Y}_{i[-j]}$ is the fitted value for the i^{th} case when the LS line is computed from all the data except case j . Here \propto means that D_j is a multiple of $\sum_i (\hat{Y}_i - \hat{Y}_{i[-j]})^2$ where the multiplier does not depend on the case. This expression implies that D_j is also an overall measure of how much the fitted values change when case j is deleted.

Given a regression problem, you should locate the points with the largest D_j s and see whether holding these cases out has a decisive influence on the fit of the model or the conclusions of the analysis. You can examine the relative magnitudes of the D_j s across cases without paying much attention to the actual value of D_j , but there are guidelines (see below) on how large D_j needs to be before you worry about it.

It is difficult to define a good strategy for dealing with outliers and influential observations. Experience is the best guide. I will show you a few examples that highlight some standard phenomena. One difficulty you will find is that certain observations may be outliers because other observations are influential, or vice-versa. If an influential observation is held out, an outlier may remain an outlier, may become influential, or both, or neither. Observations of moderate influence may become more, or less influential, when the most influential observation is held out.

Thus, any sequential refitting of models holding out of observations should not be based on the original (full-data) summaries, but rather on the summaries that result as individual observations are omitted. I tend to focus more on influential observations than outliers.

In the plots below, which cases do you think are most influential, and which are outliers. What happens in each analysis if I delete the most influential case? Are any of the remaining cases influential or poorly fitted?



Many researchers are hesitant to delete points from an analysis. I think this view is myopic, and in certain instances, such as the Gesell example to be discussed, can not be empirically supported. Being rigid about this can lead to some silly analyses of data, but one needs a very good reason and full disclosure if any points are deleted.

8.10.7 Selected diagnostic measures

A number of diagnostic measures can be calculated that are designed to measure the effect of the i^{th} observation on the regression equation. A few are described below.

1. Residuals

The difference between the observed values and predicted or fitted values. The residual is the part of the observation

that is not explained by the fitted model. You can analyze residuals to determine the adequacy of the model.

These are just the raw residuals, e_i , we defined initially. A large residual identifies an observation poorly fit by the model.

2. Standard residuals

Also known as the Studentized residual or internally Studentized residual. The standardized residual is the residual, e_i , divided by an estimate of its standard deviation. This form of the residual takes into account that the residuals may have different variances, which can make it easier to detect outliers. Standardized residuals greater than 2 and less than -2 are usually considered large.

It is possible to have quite different standard errors for residuals, and this puts everything on the same scale, r_i .

3. Deleted t-residuals

Also called externally Studentized residual. Studentized deleted residuals are useful for identifying outliers because the i^{th} residual is calculated when the fitted regression is based on all of the cases except the i^{th} one. The residual is then divided by its estimated standard deviation. Since the Studentized deleted residual for the i^{th} observation estimates all quantities with this observation deleted from the data set, the i^{th} observation cannot influence these estimates. Therefore, unusual Y values clearly stand out. Studentized deleted residuals with large absolute values are considered large.

If the regression model is appropriate, with no outlying observations, each Studentized deleted residual follows the t-distribution with $n - 1 - p$ degrees of freedom.

The problem with residuals is that a highly influential value can force the residual to have a very small value. This measure tries to correct for

that by looking at how well the model fits this observation without using this observation to construct the fit. It is quite possible for the deleted residual to be huge when the raw residual is tiny.

4. h_i (leverages)

Identify observations with unusual or outlying x -values. Observations with large leverage may exert considerable influence on the fitted value and the model. Leverage values fall between 0 and 1. Experts consider a leverage value greater than $2p/n$ or $3p/n$, where p is the number of predictors or factors plus the constant and n is the number of observations, large and suggest you examine the corresponding observation. A rule-of-thumb is to identify observations with leverage over $3p/n$ or 0.99, whichever is smaller, with an X in the table of unusual observations.

High leverage values are basically outliers among the x values. Y does not enter in this calculation. These are values with the *potential* of greatly distorting the fitted model. They may or may not actually have distorted it.

The `hat` variable from the `influence()` function on the object returned from `lm()` fit will give the leverages: `influence(lm.output)$hat`.

5. Cook's distance (D)

An overall measure of the combined impact of each observation on the fitted values. Observations with large D values may be outliers. Because D is calculated using leverage values and standardized residuals, it considers whether an observation is unusual with respect to both x - and y -values.

To interpret D, compare it to the F-distribution with $(p, n - p)$ degrees of freedom to determine the corresponding percentile. If the percentile value is less than 10% or 20%, the observation has little influence on the fitted values. If the percentile value is greater than 50%, the observation has a

major influence on the fitted values and should be examined.

Many statisticians make it a lot simpler than this sounds and use 1 as a cutoff value for large Cook's D . This is a very nice hybrid measure incorporating both x - and y -values. While high leverage values indicate observations that have the *potential* of causing trouble, those with high Cook's D values actually *do* disproportionately affect the overall fit. Using the cutoff of 1 can simplify an analysis, since frequently one or two values will have noticeably larger D values than other observations without actually having much effect, but it can be important to explore any observations that stand out.

`cooks.distance(lm.output)` will give the Cook's distance.

These various measures often flag the same observations as unusual, but they certainly can flag different observations. At the very least I examine standardized residuals and Cooks's D values. They are invaluable diagnostic measures, but nothing is perfect. Observations can be unusual in groups — a pair of unusual high leverage values close to each other will not necessarily be flagged by Cook's D since removing just one may not affect the fit very much. Any analysis takes some careful thought.

These measures and techniques really are designed for multiple regression problems where several predictor variables are used. We are using them in simple linear regression to learn what they do and see what they have to say about data, but in truth it is fairly simple with one variable to see what may be an outlier in the x -direction, to see if the data are poorly fit, etc. With more variables all that becomes quite difficult and these diagnostics are essential parts of those analyses.

8.10.8 Regression analysis suggestion

There are a lot options allowed in R. I will make a few suggestions here on how to start an analysis. What you find once you get started determines what more you might wish to explore.

1. **Plot the data.** With lots of variables the matrix plot is valuable as a quick screen. If you want to see better resolution on an individual scatter

- plot, do the individual scatter plot.
2. Do any obvious **transformations** of the data. We will discuss this in a lot more detail later. Re-plot with transformations.
 3. **Fit the least squares equation.**
 4. **Examine the residual plots and results.** Check for the patterns discussed earlier.
 - (a) Plotting several diagnostic plots together seems convenient to me. This gives you the essential plot (residuals vs. fitted values) plus a quick check on normality and possible violations of independence and influence. If you see something that needs closer investigation, you may need to recreate one of the plots larger by itself.
 - (b) Do you see curvature in the standardized residual plot? If the sign of the residuals has a distinct pattern vs. the fitted values, the linear fit is not adequate and you need some remedy, such as transformations. Note that standardized residuals are more conventional and show you what actually happened, while deleted residuals are probably the better diagnostic tool for identifying problem cases.
 - (c) Does it appear $\sigma_{Y|X}$ depends upon X (we are assuming it does not)? A megaphone pattern in residuals vs. fits is the classic (not the only) pattern to look for. Weighted least squares or transformations may be called for.
 - (d) Do you see obvious outliers? Make sure you do not have a mis-recorded data value. It might be worth refitting the equation without the outlier to see if it affects conclusions substantially.
 - (e) Is the normality assumption reasonable? This can be very closely related to the preceding points.
 - (f) Is there a striking pattern in residuals vs. order of the data? This can be an indication that the independence assumption is not valid.
 5. **Check the Cook's D values.** The Cook's distance plot and Residuals vs. Leverage (with Cook's D) plot are both helpful.
 6. If you found problem observations, omit them from the analysis and see if any conclusions change substantially. There are two good ways to do this.

- (a) Subset the `data.frame` using `subset()`.
- (b) Use `lm()` with the `weights=` option with weights of 0 for the excluded observations, weights of 1 for those included.

You may need to repeat all these steps many times for a complete analysis.

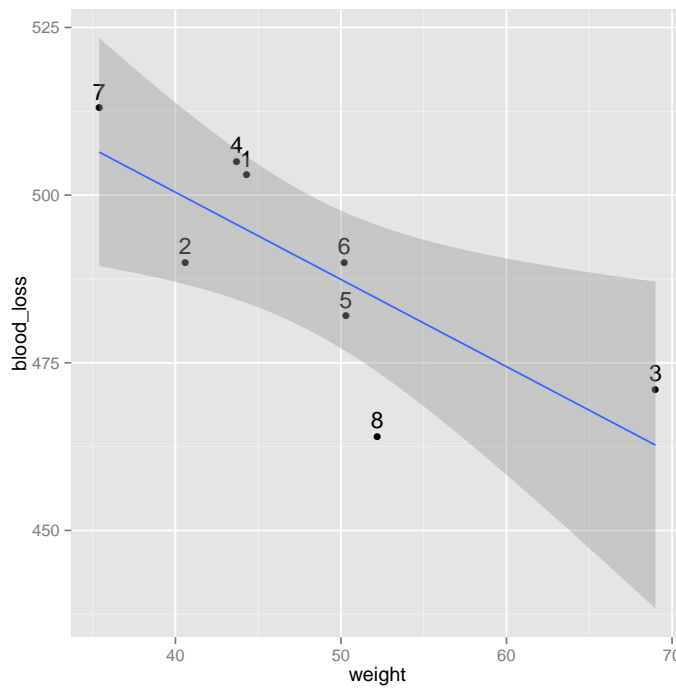
8.10.9 Residual and Diagnostic Analysis of the Blood Loss Data

We looked at much of this before, but let us go through the above steps systematically. Recall the data set (we want to predict blood loss from weight):

	weight	time	blood_loss
1	44.3	105	503
2	40.6	80	490
3	69.0	86	471
4	43.7	112	505
5	50.3	109	482
6	50.2	100	490
7	35.4	96	513
8	52.2	120	464

1. *Plot the data.* Plot blood loss vs. weight.

```
# create data ids
thyroid$id <- 1:nrow(thyroid)
# ggplot: Plot the data with linear regression fit and confidence bands
library(ggplot2)
p <- ggplot(thyroid, aes(x = weight, y = blood_loss, label = id))
p <- p + geom_point()
# plot labels next to points
p <- p + geom_text(hjust = 0.5, vjust = -0.5)
# plot regression line and confidence band
p <- p + geom_smooth(method = lm)
print(p)
```



Clearly the heaviest individual is an unusual value that warrants a closer look (maybe data recording error). I might be inclined to try a transformation here (such as `log(weight)`) to make that point a little less influential.

2. *Do any obvious transformations of the data.* We will look at transformations later.
3. *Fit the least squares equation.* Blood Loss appears significantly negatively associated with weight.

```
lm.blood.wt <- lm(blood_loss ~ weight, data = thyroid)
# use summary() to get t-tests of parameters (slope, intercept)
summary(lm.blood.wt)

##
## Call:
## lm(formula = blood_loss ~ weight, data = thyroid)
##
## Residuals:
##     Min      1Q  Median      3Q     Max 
## -20.57   -6.19    4.71    8.19    9.38 
##
## Coefficients:
##             Estimate Std. Error t value Pr(>|t|)    
## (Intercept) 552.442    21.441   25.77  2.3e-07 ***
## weight       -1.300     0.436   -2.98   0.025 *  
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 11.7 on 6 degrees of freedom
## Multiple R-squared:  0.597, Adjusted R-squared:  0.529 
## F-statistic: 8.88 on 1 and 6 DF,  p-value: 0.0247
```

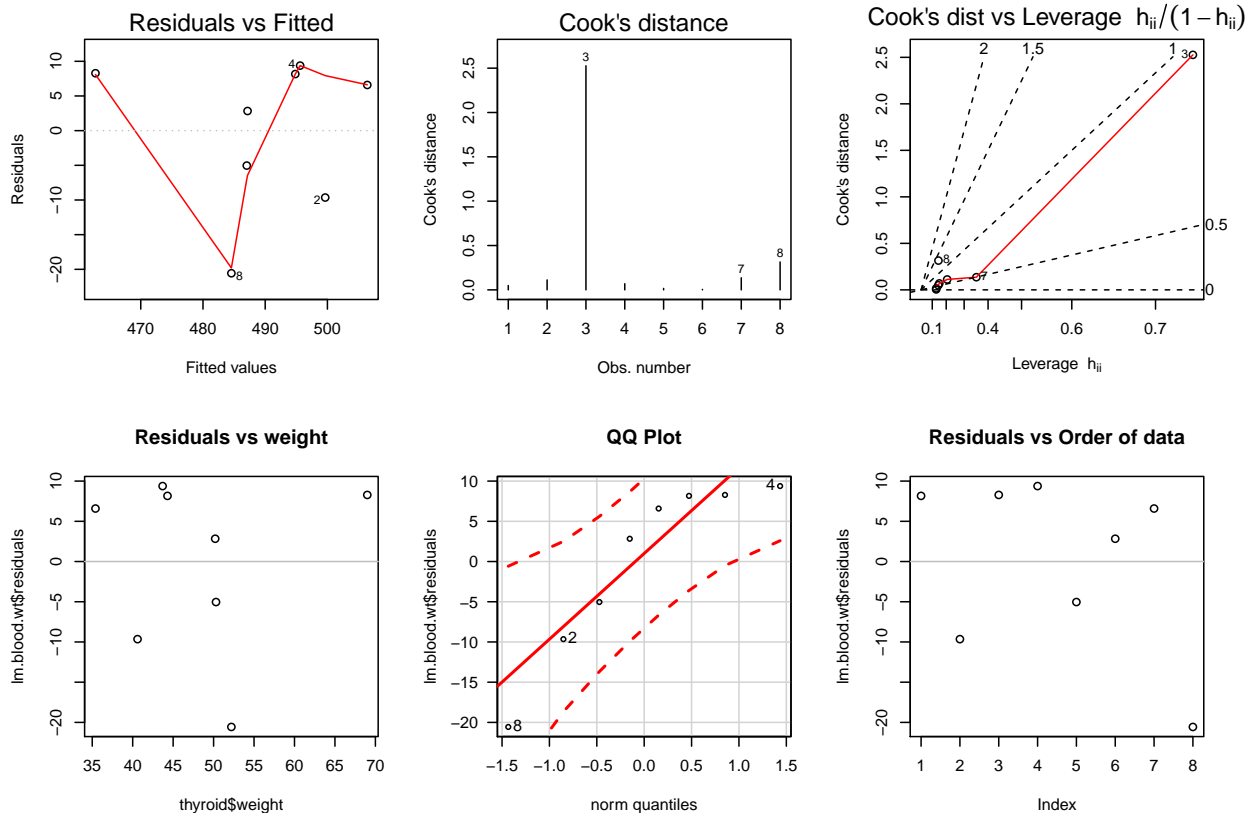
- (a) *Graphs: Check Standardized Residuals (or the Deleted Residuals).* The residual plots:

```
# plot diagnostics
par(mfrow=c(2,3))
plot(lm.blood.wt, which = c(1,4,6))

# residuals vs weight
plot(thyroid$weight, lm.blood.wt$residuals, main="Residuals vs weight")
  # horizontal line at zero
  abline(h = 0, col = "gray75")

# Normality of Residuals
library(car)
# qq plot for studentized resid
#   las    = 1 : turns labels on y-axis to read horizontally
#   id.n   = n : labels n most extreme observations, and outputs to console
#   id.cex = 1 : is the size of those labels
#   lwd    = 1 : line width
qqPlot(lm.blood.wt$residuals, las = 1, id.n = 3, main="QQ Plot")
## 8 2 4
## 1 2 8

# residuals vs order of data
plot(lm.blood.wt$residuals, main="Residuals vs Order of data")
  # horizontal line at zero
  abline(h = 0, col = "gray75")
```



4. Examine the residual plots and results.

- (a) Do you see curvature? There does not appear to be curvature (and it could be hard to detect with so few points).
- (b) Does it appear $\sigma_{Y|X}$ depends upon X ? Not much evidence for this.
- (c) Do you see obvious outliers? Observation 3 is an outlier in the x direction, and therefore possibly a high leverage point and influential on the model fit.
- (d) Is the normality assumption reasonable? There appears to be some skewness, but with so few points normality may be reasonable.
- (e) Is there a striking pattern in residuals vs. order of the data? No striking pattern.

5. Check the Cook's D values. We anticipated that the 3rd observation is affecting the fit by a lot more than any other values. The D -value is much larger than 1. Note that the residual is not large for this value.

6. Omit problem observations from the analysis and see if any conclusions change substantially. Let us refit the equation without observation 3 to see if anything changes drastically. I will use the weighted least squares approach discussed earlier on this example. Define a variable **wt** that is 1 for all observations except obs. 3, and make it 0 for that one.

```
# wt = 1 for all except obs 3 where wt = 0
thyroid$wt <- as.numeric(!(thyroid$id == 3))
thyroid$wt

## [1] 1 1 0 1 1 1 1 1
```

What changes by deleting case 3? The fitted line gets steeper (slope changes from -1.30 to -2.19), adjusted R^2 gets larger (up to 58% from 53%), and S changes from 11.7 to 10.6. Because the Weight values are much less spread out, $SE(\hat{\beta}_1)$ becomes quite a bit larger (to 0.714, up from 0.436) and we lose a degree of freedom for MS Error (which will penalize us on tests and CIs). Just about any quantitative statement we would want to make using CIs would be about the same either way since CIs will overlap a great deal, and our qualitative interpretations are unchanged (Blood Loss drops with Weight). Unless something shows up in the plots, I don't see any very important changes here.

```
lm.blood.wt.no3 <- lm(blood_loss ~ weight, data = thyroid, weights = wt)
# use summary() to get t-tests of parameters (slope, intercept)
summary(lm.blood.wt.no3)

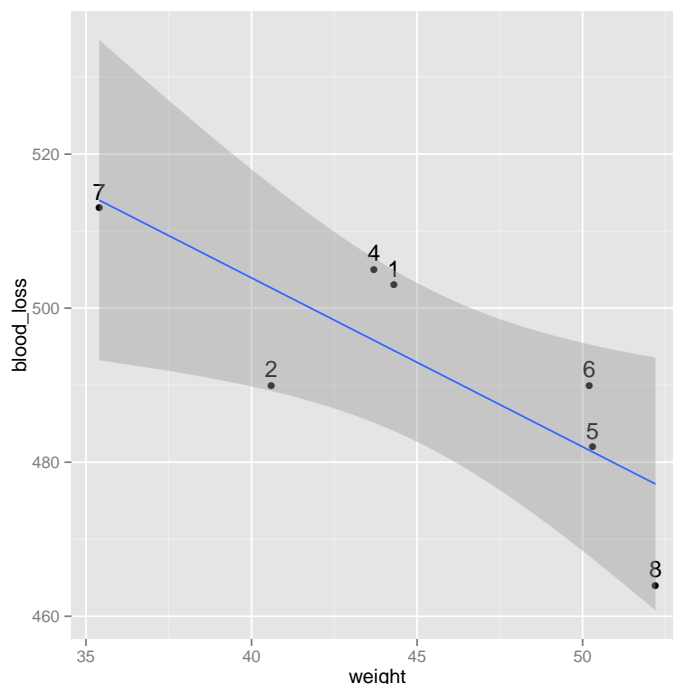
##
## Call:
## lm(formula = blood_loss ~ weight, data = thyroid, weights = wt)
```

```

## 
## Weighted Residuals:
##      1      2      3      4      5      6      7      8
##  8.503 -12.613   0.000   9.187   0.664   8.445 -1.019 -13.168
##
## 
## Coefficients:
##              Estimate Std. Error t value Pr(>|t|)    
## (Intercept) 591.668    32.567   18.17  9.3e-06 ***
## weight       -2.193     0.714   -3.07   0.028 *  
## ---        
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## 
## Residual standard error: 10.6 on 5 degrees of freedom
## Multiple R-squared:  0.653, Adjusted R-squared:  0.584 
## F-statistic: 9.43 on 1 and 5 DF,  p-value: 0.0278

# exclude obs 3
thyroid.no3 <- subset(thyroid, wt == 1)
# ggplot: Plot the data with linear regression fit and confidence bands
library(ggplot2)
p <- ggplot(thyroid.no3, aes(x = weight, y = blood_loss, label = id))
p <- p + geom_point()
# plot labels next to points
p <- p + geom_text(hjust = 0.5, vjust = -0.5)
# plot regression line and confidence band
p <- p + geom_smooth(method = lm)
print(p)

```



Nothing very striking shows up in the residual plots, and no Cook's D values are very large among the remaining observations.

```

# plot diagnostics
par(mfrow=c(2,3))
plot(lm.blood.wt.no3, which = c(1,4,6))

```

```

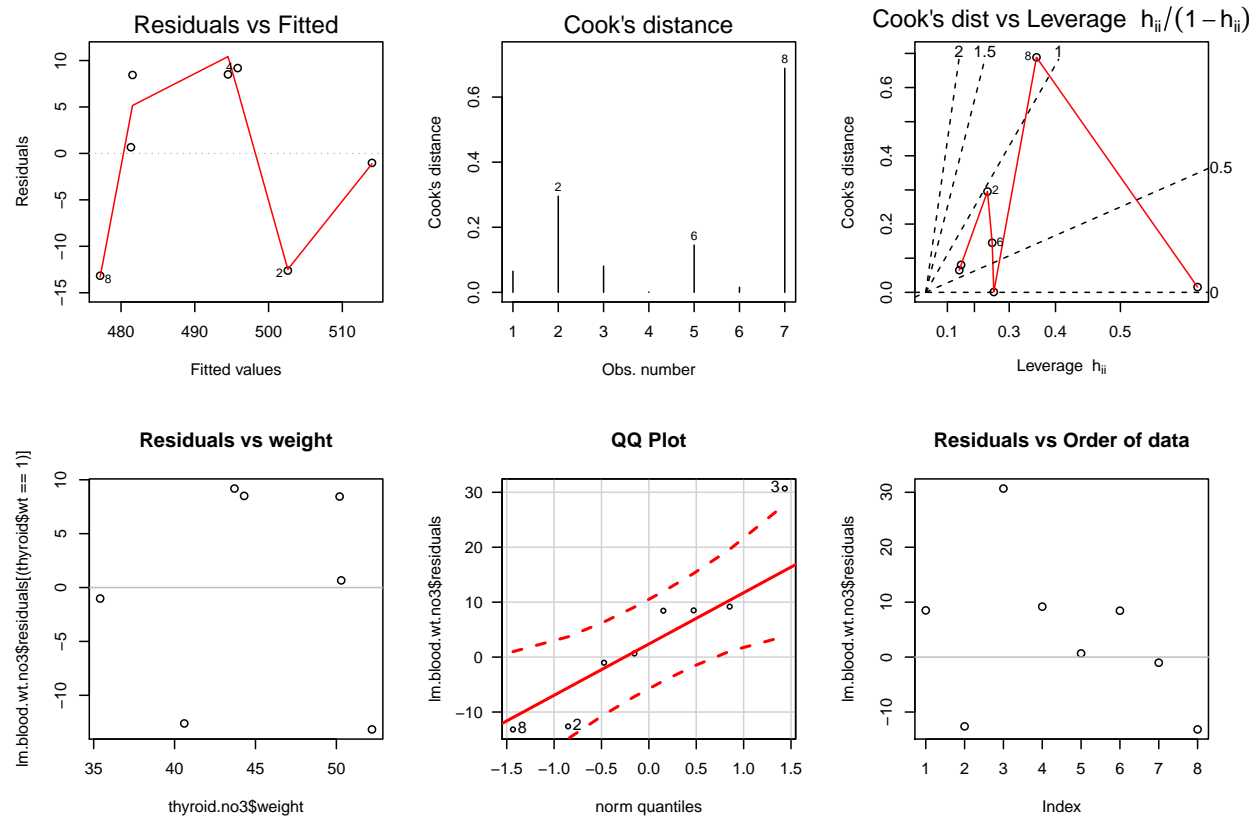
# residuals vs weight
plot(thyroid.no3$weight, lm.blood.wt.no3$residuals[(thyroid$wt == 1)],
      main="Residuals vs weight")
# horizontal line at zero
abline(h = 0, col = "gray75")

# Normality of Residuals
library(car)
# qq plot for studentized resid
#   las    = 1 : turns labels on y-axis to read horizontally
#   id.n   = n : labels n most extreme observations, and outputs to console
#   id.cex = 1 : is the size of those labels
#   lwd    = 1 : line width
qqPlot(lm.blood.wt.no3$residuals, las = 1, id.n = 3, main="QQ Plot")

## 3 8 2
## 8 1 2

# residuals vs order of data
plot(lm.blood.wt.no3$residuals, main="Residuals vs Order of data")
# horizontal line at zero
abline(h = 0, col = "gray75")

```



How much difference is there in a practical sense? Examine the 95% prediction interval for a new observation at Weight = 50kg. Previously we saw that interval based on all 8 observations was from 457.1 to 517.8 ml of Blood Loss.

Based on just the 7 observations the prediction interval is 451.6 to 512.4 ml. There really is no practical difference here.

```
# CI for the mean and PI for a new observation at weight=50
predict(lm.blood.wt      , data.frame(weight=50), interval = "prediction")
##     fit    lwr   upr
## 1 487.4 457.1 517.8

predict(lm.blood.wt.no3, data.frame(weight=50), interval = "prediction")
## Warning: Assuming constant prediction variance even though model fit is weighted
##     fit    lwr   upr
## 1 482 451.6 512.4
```

Therefore, while obs. 3 was potentially influential, whether the value is included or not makes very little difference in the model fit or relationship between Weight and BloodLoss.

8.10.10 Gesell data

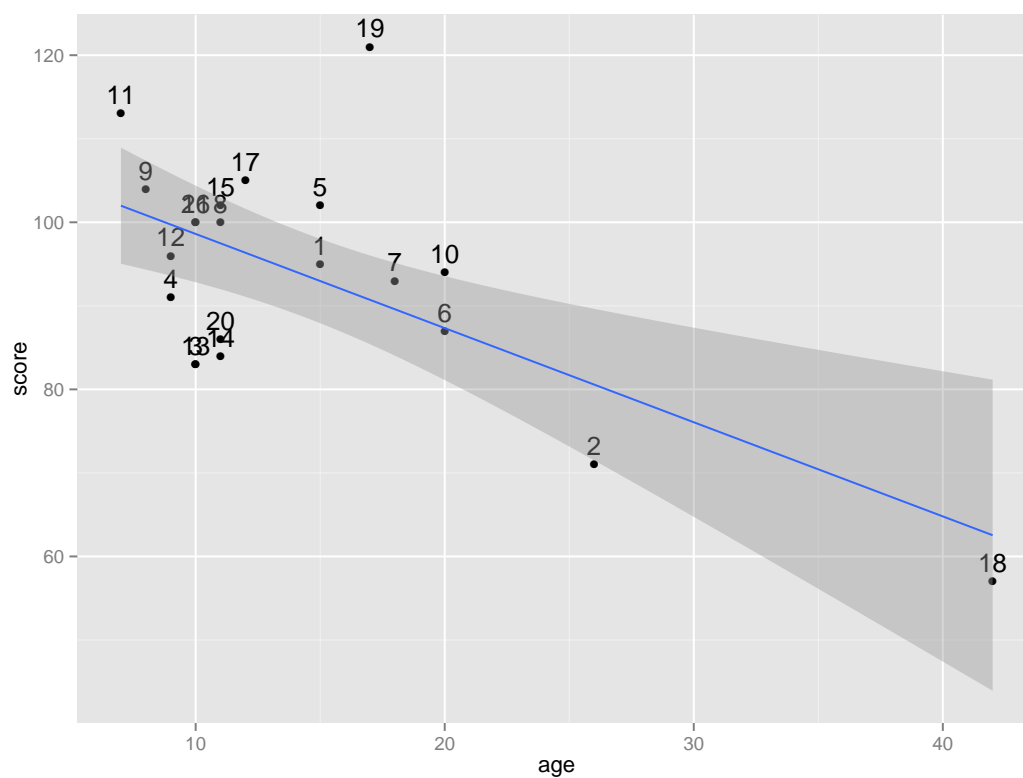
These data are from a UCLA study of cyanotic heart disease in children. The predictor is the age of the child in months at first word and the response variable is the Gesell adaptive score, for each of 21 children.

	id	age	score
1	1	15	95
2	2	26	71
3	3	10	83
4	4	9	91
5	5	15	102
6	6	20	87
7	7	18	93
8	8	11	100
9	9	8	104
10	10	20	94
11	11	7	113
12	12	9	96
13	13	10	83
14	14	11	84
15	15	11	102
16	16	10	100
17	17	12	105
18	18	42	57
19	19	17	121
20	20	11	86
21	21	10	100

Let us go through the same steps as before.

1. Plot Score versus Age. Comment on the relationship between Score and Age.

```
# ggplot: Plot the data with linear regression fit and confidence bands
library(ggplot2)
p <- ggplot(gesell, aes(x = age, y = score, label = id))
p <- p + geom_point()
# plot labels next to points
p <- p + geom_text(hjust = 0.5, vjust = -0.5)
# plot regression line and confidence band
p <- p + geom_smooth(method = lm)
print(p)
```



2. There are no obvious transformations to try here.
3. Fit a simple linear regression model. Provide an equation for the LS line. Does age at first word appear to be an “important predictor” of Gesell adaptive score? (i.e., is the estimated slope significantly different from zero?)

```
lm.score.age <- lm(score ~ age, data = gesell)
# use summary() to get t-tests of parameters (slope, intercept)
summary(lm.score.age)

##
## Call:
## lm(formula = score ~ age, data = gesell)
##
## Residuals:
##     Min      1Q  Median      3Q     Max 
## -15.60   -8.73    1.40    4.52   30.29 
## 
## Coefficients:
##             Estimate Std. Error t value Pr(>|t|)    
## (Intercept) 109.87      5.07   21.68 7.3e-15 ***
## age         -1.13      0.31   -3.63  0.0018 **  
## ---        
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1 
## 
## Residual standard error: 11 on 19 degrees of freedom
## Multiple R-squared:  0.41, Adjusted R-squared:  0.379 
## F-statistic: 13.2 on 1 and 19 DF,  p-value: 0.00177
```

4. Do these plots suggest any inadequacies with the model?

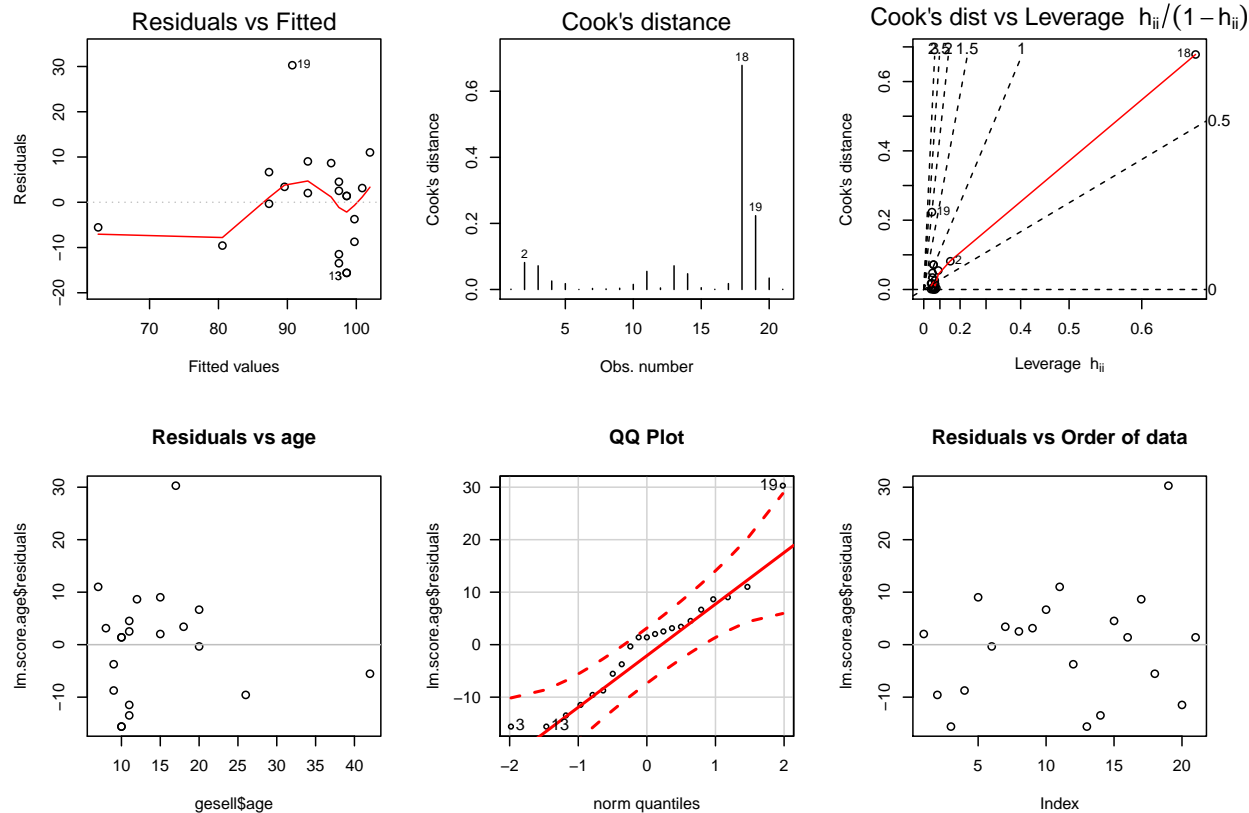
```
# plot diagnostics
par(mfrow=c(2,3))
plot(lm.score.age, which = c(1,4,6))

# residuals vs weight
plot(gesell$age, lm.score.age$residuals, main="Residuals vs age")
# horizontal line at zero
abline(h = 0, col = "gray75")

# Normality of Residuals
library(car)
qqPlot(lm.score.age$residuals, las = 1, id.n = 3, main="QQ Plot")

## 19 3 13
## 21 1 2

# residuals vs order of data
plot(lm.score.age$residuals, main="Residuals vs Order of data")
# horizontal line at zero
abline(h = 0, col = "gray75")
```



5. Observations 18 and 19 stand out with relatively high Cook's D. The cutoff line is only a rough guideline. Those two were flagged with high influence and standardized residual, respectively, also. Be sure to examine the scatter plot carefully to see why 18 and 19 stand out.

6. Consider doing two additional analyses: Analyze the data after omitting case 18 only and analyze the data after omitting case 19 only. Refit the regression model for each of these two scenarios. Provide a summary table such as the following, giving the relevant summary statistics for the three analyses. Discuss the impact that observations 18 and 19 have individually on the fit of the model.

When observation 18 is omitted, the estimated slope is not significantly different from zero ($p\text{-value} = 0.1489$), indicating that age is not an important predictor of Gesell score. This suggests that the significance of age as a predictor in the original analysis was due solely to the presence of observation 18. Note the dramatic decrease in R^2 after deleting observation 18.

The fit of the model appears to improve when observation 19 is omitted. For example, R^2 increases noticeably and the p-value for testing the significance of the slope decreases dramatically (in a relative sense). These tendencies would be expected based on the original plot. However, this improvement is misleading. Once observation 19 is omitted, observation 18 is much more influential. Again the significance of the slope is due to the presence of observation 18.

Feature	Full data	Omit 18	Omit 19
b_0	109.87	105.63	109.30
b_1	-1.13	-0.78	-1.19
$SE(b_0)$	5.07	7.16	3.97
$SE(b_1)$	0.31	0.52	0.24
R^2	0.41	0.11	0.57
p-val for $H_0 : \beta_1 = 0$	0.002	0.149	0.000

Can you think of any reasons to justify doing the analysis without observation 18?

If you include observation 18 in the analysis, you are assuming that the mean Gesell score is linearly related to age over the entire range of observed ages. Observation 18 is far from the other observations on age (age for observation 18 is 42; the second highest age is 26; the lowest age is 7). There are no

children with ages between 27 and 41, so we have no information on whether the relationship is roughly linear over a significant portion of the range of ages. I am comfortable deleting observation 18 from the analysis because it's inclusion forces me to make an assumption that I can not check using these data. I am only willing to make predictions of Gesell score for children with ages roughly between 7 and 26. However, once this point is omitted, age does not appear to be an important predictor.

A more complete analysis would delete observation 18 and 19 together. What would you expect to see if you did this?

8.10.11 Weighted Least Squares

Earlier I indicated that nonconstant error variance can be addressed (sometimes) with weighted least squares. The *scedastic function* is the conditional variance of Y given X . Y is said to be **heteroscedastic** if its variance depends on the value of X (variance changes), and **homoscedastic** if its variance does not depend on X (constant variance).

Recall the LS (OLS or ordinary LS) line chooses the values of β_0 and β_1 that minimize

$$\sum_{i=1}^n \{Y_i - (\beta_0 + \beta_1 X_i)\}^2$$

over all possible choices of β_0 and β_1 . The weighted LS (WLS) line chooses the values of β_0 and β_1 that minimize

$$\sum_{i=1}^n w_i \{Y_i - (\beta_0 + \beta_1 X_i)\}^2$$

over all possible choices of β_0 and β_1 . If $\sigma_{Y|X}$ depends up X , then the correct choice of weights is inversely proportional to *variance*, $w_i \propto \sigma_{Y|X}^{-2}$.

Consider the following data and plot of y vs. x and standardized OLS residuals vs x . It is very clear that variability increases with x .

```
#### Weighted Least Squares
# R code to generate data
set.seed(7)
n <- 100
# 1s, Xs uniform 0 to 100
X <- matrix(c(rep(1,n),runif(n,0,100)), ncol=2)
# intercept and slope (5, 5)
```

```

beta <- matrix(c(5,5),ncol=1)
# errors are X*norm(0,1), so variance increases with X
e <- X[,2]*rnorm(n,0,1)
# response variables
y <- X %*% beta + e

# put data into data.frame
wlsdat <- data.frame(y, x = X[,2])

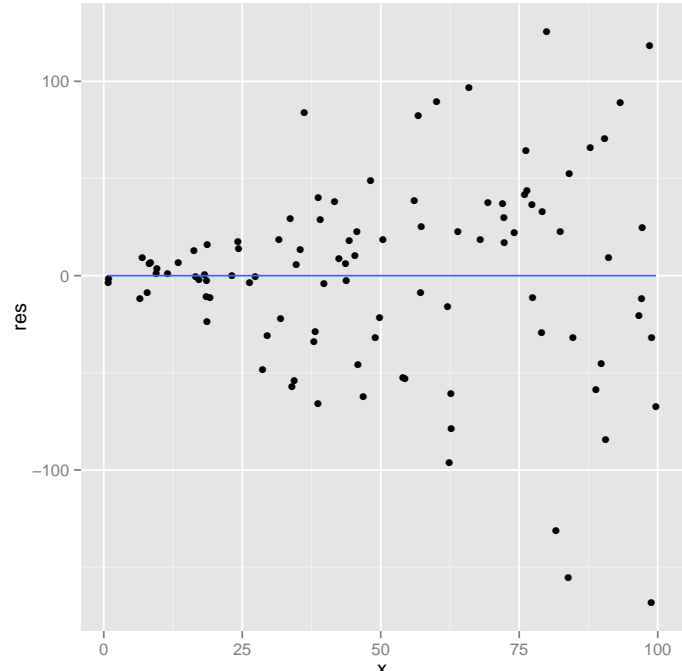
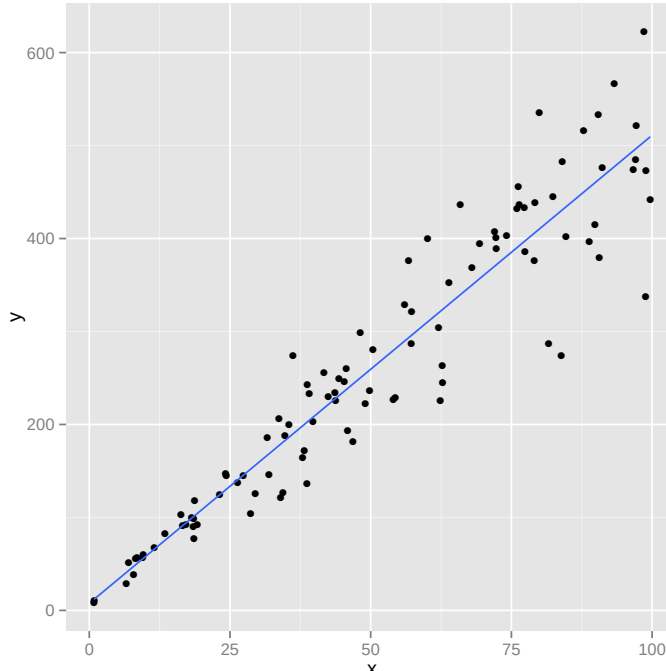
# fit regression
lm.y.x <- lm(y ~ x, data = wlsdat)

# put residuals in data.frame
wlsdat$res <- lm.y.x$residuals

# ggplot: Plot the data with linear regression fit
library(ggplot2)
p <- ggplot(wlsdat, aes(x = x, y = y))
p <- p + geom_point()
p <- p + geom_smooth(method = lm, se = FALSE)
print(p)

# ggplot: Plot the residuals
library(ggplot2)
p <- ggplot(wlsdat, aes(x = x, y = res))
p <- p + geom_point()
p <- p + geom_smooth(method = lm, se = FALSE)
print(p)

```



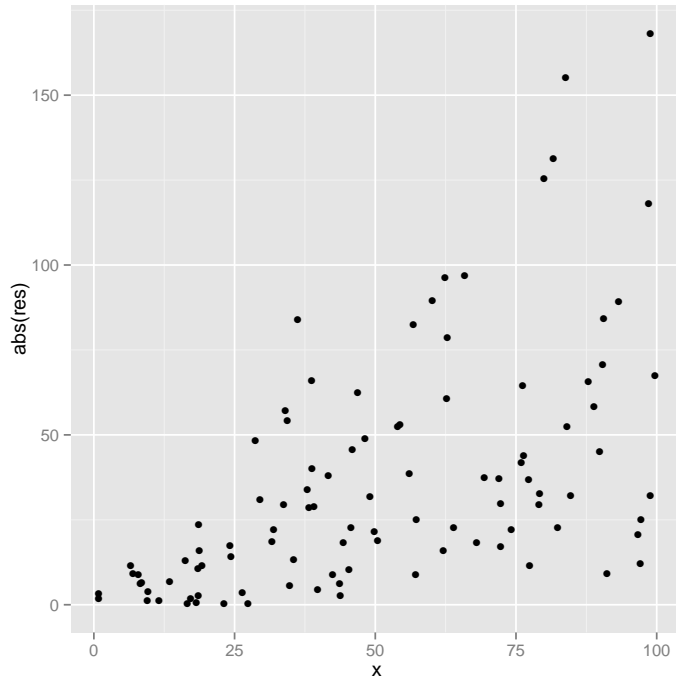
In order to use WLS to solve this problem, we need some form for $\sigma_{Y|X}^2$. Finding that form is a real problem with WLS. It can be useful to plot the absolute value of the standardized residual vs. x to see if the top boundary seems to follow a general pattern.

```

# ggplot: Plot the absolute value of the residuals
library(ggplot2)
p <- ggplot(wlsdat, aes(x = x, y = abs(res)))

```

```
p <- p + geom_point()
print(p)
```

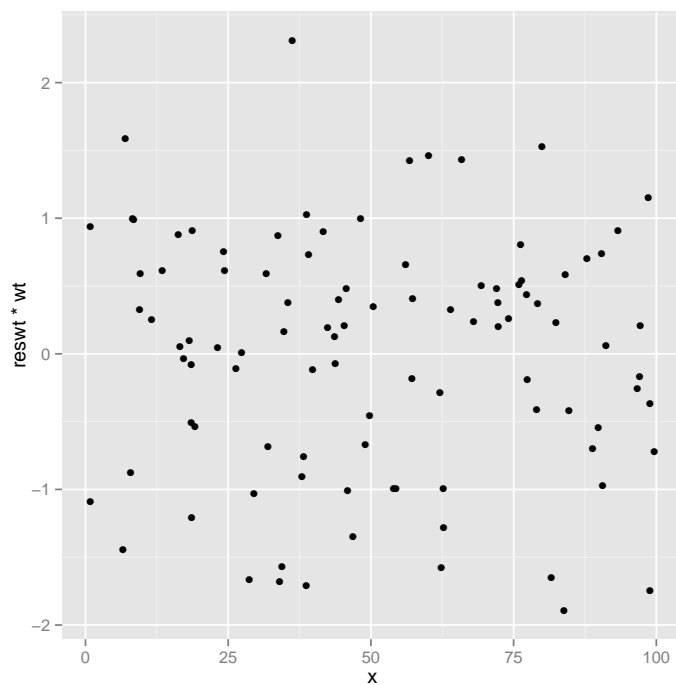


It is plausible the upper boundary is linear, so let us try $w_i = \frac{1}{x^2}$. Standardized residuals from this WLS fit look very good. Note that raw (non-standardized) residuals will still have the same pattern — it is essential to use standardized residuals here.

```
# fit regression
lm.y.x.wt <- lm(y ~ x, data = wlsdat, weights = x^(-2))

# put residuals in data.frame
wlsdat$reswt <- lm.y.x.wt$residuals
wlsdat$wt <- lm.y.x.wt$weights^(1/2)

# ggplot: Plot the absolute value of the residuals
library(ggplot2)
p <- ggplot(wlsdat, aes(x = x, y = reswt*wt))
p <- p + geom_point()
print(p)
```



Compare also the OLS fitted equation:

```
summary(lm.y.x)

##
## Call:
## lm(formula = y ~ x, data = wlsdat)
##
## Residuals:
##     Min      1Q  Median      3Q     Max 
## -168.18   -24.94    2.54   24.97  125.37 
## 
## Coefficients:
##             Estimate Std. Error t value Pr(>|t|)    
## (Intercept)  7.631     10.426   0.73   0.47    
## x          5.035      0.179   28.12  <2e-16 *** 
## ---        
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## 
## Residual standard error: 50.5 on 98 degrees of freedom
## Multiple R-squared:  0.89, Adjusted R-squared:  0.889 
## F-statistic: 790 on 1 and 98 DF,  p-value: <2e-16
```

to the WLS fitted equation:

```
summary(lm.y.x.wt)

##
## Call:
## lm(formula = y ~ x, data = wlsdat, weights = x^(-2))
##
## Weighted Residuals:
##     Min      1Q  Median      3Q     Max 
## -1.894  -0.671   0.178   0.596  2.313 
## 
## Coefficients:
```

```
##             Estimate Std. Error t value Pr(>|t|)  
## (Intercept) 5.0931     0.5493    9.27 4.6e-15 ***  
## x           5.1069     0.0939   54.40 < 2e-16 ***  
## ---  
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1  
##  
## Residual standard error: 0.89 on 98 degrees of freedom  
## Multiple R-squared: 0.968, Adjusted R-squared: 0.968  
## F-statistic: 2.96e+03 on 1 and 98 DF, p-value: <2e-16
```

Clearly the weighted fit looks better, although note that everything is based on the weighted SS. In practice it can be pretty difficult to determine the correct set of weights, but WLS works much better than OLS if appropriate. I actually simulated this data set using $\beta_0 = \beta_1 = 5$. Which fit actually did better?

Chapter 9

Introduction to the Bootstrap

Learning objectives

After completing this topic, you should be able to:

- explain** the bootstrap principle for hypothesis tests and inference.
- decide** (for simple problems) how to construct a bootstrap procedure.

Achieving these goals contributes to mastery in these course learning outcomes:

- 1. organize knowledge.
- 5. define parameters of interest and hypotheses in words and notation.
- 8. use statistical software.
- 12. make evidence-based decisions.

9.1 Introduction

Statistical theory attempts to answer three basic questions:

- 1. How should I collect my data?
- 2. How should I analyze and summarize the data that I've collected?
- 3. How accurate are my data summaries?

Question 3 constitutes part of the process known as statistical inference. The bootstrap makes certain kinds of statistical inference¹. Let's look at an example.

¹Efron (1979), “Bootstrap methods: another look at the jackknife.” Ann. Statist. 7, 1–26

Example: Aspirin and heart attacks, large-sample theory Does aspirin prevent heart attacks in healthy middle-aged men? A controlled, randomized, double-blind study was conducted and gathered the following data.
(fatal plus non-fatal)

	heart attacks	subjects
aspirin group:	104	11037
placebo group:	189	11034

A good experimental design, such as this one, simplifies the results! The ratio of the two rates (the risk ratio) is

$$\hat{\theta} = \frac{104/11037}{189/11034} = 0.55.$$

Because of the solid experimental design, we can believe that the aspirin-takers only have 55% as many heart attacks as the placebo-takers.

We are not really interested in the estimated ratio $\hat{\theta}$, but the true ratio, θ . That is the ratio if we could treat all possible subjects, not just a sample of them. Large sample theory tells us that the log risk ratio has an approximate Normal distribution. The standard error of the log risk ratio is estimated simply by the square root of the sum of the reciprocals of the four frequencies:

$$\text{SE}(\log(RR)) = \sqrt{\frac{1}{104} + \frac{1}{189} + \frac{1}{11037} + \frac{1}{11034}} = 0.1228$$

The 95% CI for $\log(\theta)$ is

$$\log(\hat{\theta}) \pm 1.96 \times \text{SE}(\log(RR)), \quad (-0.839, -0.357),$$

and exponentiating gives the CI on the ratio scale,

$$\exp\{\log(\hat{\theta}) \pm 1.96 \times \text{SE}(\log(RR))\}, \quad (0.432, 0.700).$$

The same data that allowed us to estimate the ratio θ with $\hat{\theta} = 0.55$ also allowed us to get an idea of the estimate's accuracy.

Example: Aspirin and strokes, large-sample theory The aspirin study tracked strokes as well as heart attacks.

	strokes	subjects
aspirin group:	119	11037
placebo group:	98	11034

The ratio of the two rates (the risk ratio) is

$$\hat{\theta} = \frac{119/11037}{98/11034} = 1.21.$$

It looks like aspirin is actually harmful, now, however the 95% interval for the true stroke ratio θ is $(0.925, 1.583)$. This includes the neutral value $\theta = 1$, at which aspirin would be no better or worse than placebo for strokes.

9.2 Bootstrap

The bootstrap is a data-based simulation method for statistical inference, which can be used to produce inferences like those in the previous slides. The term “bootstrap” comes from literature. In “The Adventures of Baron Munchausen”, by Rudolph Erich Raspe, the Baron had fallen to the bottom of a deep lake, and he thought to get out by *pulling himself up by his own bootstraps*.

9.2.1 Ideal versus Bootstrap world, sampling distributions

Ideal world

1. Population of interest
2. Obtain many simple random samples (SRSs) of size n
3. For each SRS, calculate statistic of interest (θ)
4. Sampling distribution is the distribution of the calculated statistic

Bootstrap world

1. Population of interest; One empirical distribution based on a sample of size n
2. Obtain many bootstrap resamples of size n
3. For each resample, calculate statistic of interest (θ^*)
4. Bootstrap distribution is the distribution of the calculated statistic

5. Bootstrap distribution estimates the sampling distribution centered at the statistic (not the parameter).

Example: Aspirin and strokes, bootstrap Here's how the bootstrap works in the stroke example. We create two populations:

- the first consisting of 119 ones and $11037 - 119 = 10918$ zeros,
- the second consisting of 98 ones and $11034 - 98 = 10936$ zeros.

We draw with replacement a sample of 11037 items from the first population, and a sample of 11034 items from the second population. Each is called a *bootstrap sample*. From these we derive the bootstrap replicate of $\hat{\theta}$:

$$\hat{\theta}^* = \frac{\text{Proportion of ones in bootstrap sample 1}}{\text{Proportion of ones in bootstrap sample 2}}.$$

Repeat this process a large number of times, say 10000 times, and obtain 10000 *bootstrap replicates* $\hat{\theta}^*$. The summaries are in the code, followed by a histogram of bootstrap replicates, $\hat{\theta}^*$.

```
#### Example: Aspirin and strokes, bootstrap
# sample size (n) and successes (s) for sample 1 (aspirin) and 2 (placebo)
n <- c(11037, 11034)
s <- c( 119,     98)
# data for samples 1 and 2, where 1 = success (stroke), 0 = failure (no stroke)
dat1 <- c(rep(1, s[1]), rep(0, n[1] - s[1]))
dat2 <- c(rep(1, s[2]), rep(0, n[2] - s[2]))
# draw R bootstrap replicates
R <- 10000
# init location for bootstrap samples
bs1 <- rep(NA, R)
bs2 <- rep(NA, R)
# draw R bootstrap resamples of proportions
for (i in 1:R) {
  # proportion of successes in bootstrap samples 1 and 2
  # (as individual steps for group 1:)
  resam1 <- sample(dat1, n[1], replace = TRUE)
  success1 <- sum(resam1)
  bs1[i] <- success1 / n[1]
  # (as one line for group 2:)
  bs2[i] <- sum(sample(dat2, n[2], replace = TRUE)) / n[2]
}
# bootstrap replicates of ratio estimates
rat <- bs1 / bs2
# sort the ratio estimates to obtain bootstrap CI
rat.sorted <- sort(rat)
# 0.025th and 0.975th quantile gives equal-tail bootstrap CI
```

```

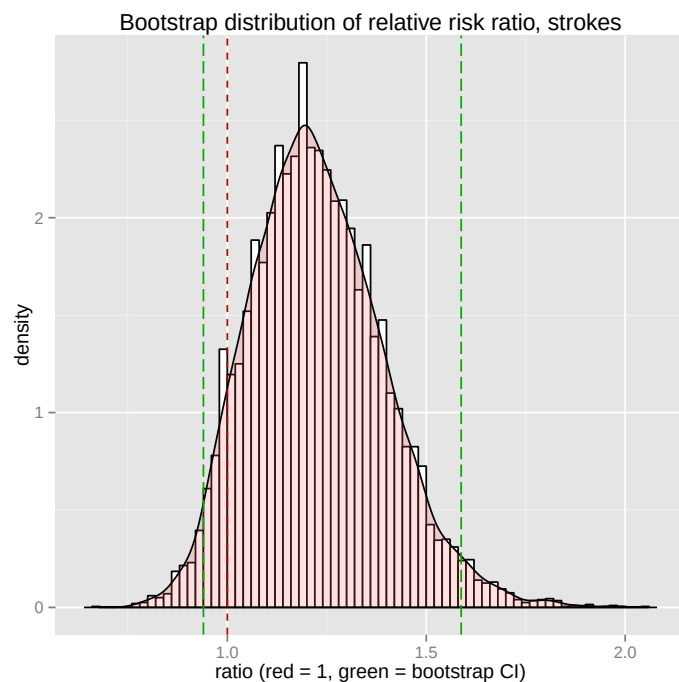
CI.bs <- c(rat.sorted[round(0.025*R)], rat.sorted[round(0.975*R+1)])
CI.bs
## [1] 0.9399 1.5878

## Plot the bootstrap distribution with CI
# First put data in data.frame for ggplot()
dat.rat <- data.frame(rat)

library(ggplot2)
p <- ggplot(dat.rat, aes(x = rat))
p <- p + geom_histogram(aes(y=..density..),
                        , binwidth=0.02
                        , colour="black", fill="white")
# Overlay with transparent density plot
p <- p + geom_density(alpha=0.2, fill="#FF6666")
# vertical line at 1 and CI
p <- p + geom_vline(xintercept=1, colour="#BB0000", linetype="dashed")
p <- p + geom_vline(xintercept=CI.bs[1], colour="#00AA00", linetype="longdash")
p <- p + geom_vline(xintercept=CI.bs[2], colour="#00AA00", linetype="longdash")
p <- p + labs(title = "Bootstrap distribution of relative risk ratio, strokes")
p <- p + xlab("ratio (red = 1, green = bootstrap CI)")
print(p)

## Warning: position_stack requires constant width: output may be incorrect

```



In this simple case, the confidence interval derived from the bootstrap (0.94, 1.588) agrees very closely with the one derived from statistical theory (0.925, 1.583). Bootstrap methods are intended to simplify the calculation of inferences like those using large-sample theory, producing them in an automatic way even in situations much more complicated than the risk ratio in the aspirin

example.

9.2.2 The accuracy of the sample mean

For sample means, and essentially *only* for sample means, an accuracy formula (for the standard error of the parameter) is easy to obtain (using the delta method). We'll see how to use the bootstrap for the sample mean, then for the more complicated situation of assessing the accuracy of the median.

Bootstrap Principle The **plug-in principle** is used when the underlying distribution is unknown and you substitute your best guess for what that distribution is. What to substitute?

Empirical distribution ordinary bootstrap

Smoothed distribution (kernel) smoothed bootstrap

Parametric distribution parametric bootstrap

Satisfy assumptions such as the null hypothesis

This substitution works in many cases, but not always. Keep in mind that the bootstrap distribution is centered at the statistic, not the parameter. Implementation is done by Monte Carlo sampling.

The bootstrap is commonly implemented in one of two ways, nonparametrically or parametrically. An *exact* **nonparametric bootstrap** requires n^n samples! That's one for every possible combination of each of n observation positions taking the value of each of n observations. This is sensibly approximated by using the Monte Carlo strategy of drawing a large number (1000 or 10000) of random resamples. On the other hand, a **parametric bootstrap** first assumes a distribution for the population (such as a normal distribution) and estimates the distributional parameters (such as the mean and variance) from the observed sample. Then, the Monte Carlo strategy is used to draw a large number (1000 or 10000) of samples from the estimated parametric distribution.

Example: Mouse survival, two-sample t-test, mean Sixteen mice were randomly assigned to a treatment group or a control group. Shown are their survival times, in days, following a test surgery. Did the treatment prolong survival?

Group	Data	n	Mean	SE
Control:	52, 104, 146, 10, 51, 30, 40, 27, 46	9	56.22	14.14
Treatment:	94, 197, 16, 38, 99, 141, 23	7	86.86	25.24
Difference:			30.63	28.93

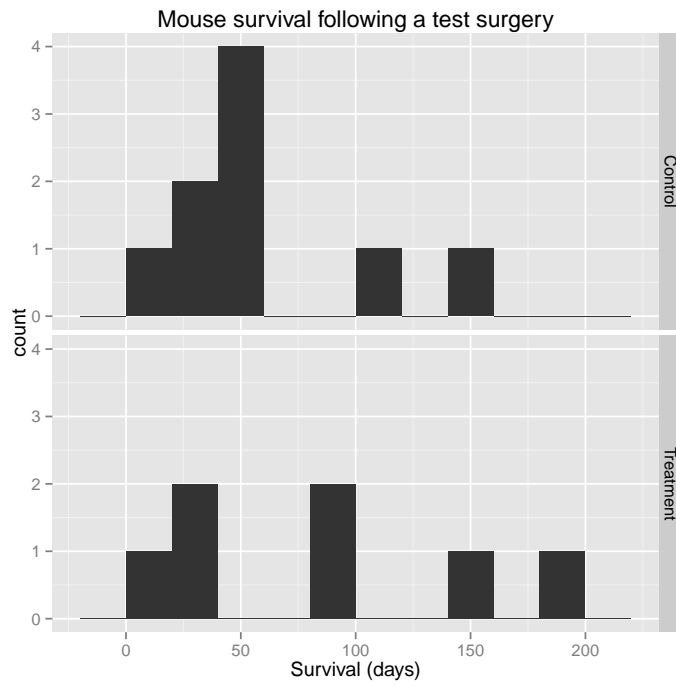
Numerical and graphical summaries of the data are below. There seems to be a slight difference in variability between the two treatment groups.

```
#### Example: Mouse survival, two-sample t-test, mean
treatment <- c(94, 197, 16, 38, 99, 141, 23)
control   <- c(52, 104, 146, 10, 51, 30, 40, 27, 46)
survive <- c(treatment, control)
group   <- c(rep("Treatment", length(treatment)), rep("Control", length(control)))
mice <- data.frame(survive, group)

library(plyr)
# ddply "dd" means the input and output are both data.frames
mice.summary <- ddply(mice,
  "group",
  function(X) {
    data.frame( m = mean(X$survive),
                s = sd(X$survive),
                n = length(X$survive)
    )
  }
)
# standard errors
mice.summary$se   <- mice.summary$s/sqrt(mice.summary$n)
# individual confidence limits
mice.summary$ci.l <- mice.summary$m - qt(1-0.05/2, df=mice.summary$n-1) * mice.summary$se
mice.summary$ci.u <- mice.summary$m + qt(1-0.05/2, df=mice.summary$n-1) * mice.summary$se

mice.summary
##      group     m     s     n     se   ci.l   ci.u
## 1 Control 56.22 42.48 9 14.16 23.57 88.87
## 2 Treatment 86.86 66.77 7 25.24 25.11 148.61
diff(mice.summary$m)                                ##
## [1] 30.63

# histogram using ggplot
p <- ggplot(mice, aes(x = survive))
p <- p + geom_histogram(binwidth = 20)
p <- p + facet_grid(group ~ .)
p <- p + labs(title = "Mouse survival following a test surgery") + xlab("Survival (days)")
print(p)
```



The standard error for the difference is $28.93 = \sqrt{25.24^2 + 14.14^2}$, so the observed difference of 30.63 is only $30.63/28.93=1.05$ estimated standard errors greater than zero, an *insignificant* result.

The two-sample t -test of the difference in means confirms the lack of statistically significant difference between these two treatment groups with a p-value=0.3155.

```
t.test(survive ~ group, data = mice)
##
##  Welch Two Sample t-test
##
## data:  survive by group
## t = -1.059, df = 9.654, p-value = 0.3155
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -95.42 34.15
## sample estimates:
## mean in group Control mean in group Treatment
## 56.22 86.86
```

But these are small samples, and the control sample does not look normal. We could do a nonparametric two-sample test of difference of medians. Or, we could use the bootstrap to make our inference.

Example: Mouse survival, two-sample bootstrap, mean Here's how the bootstrap works in the two-sample mouse example. We draw with replacement from each sample, calculate the mean for each sample, then take

the difference in means. Each is called a *bootstrap sample* of the difference in means. From these we derive the bootstrap replicate of $\hat{\mu}$:

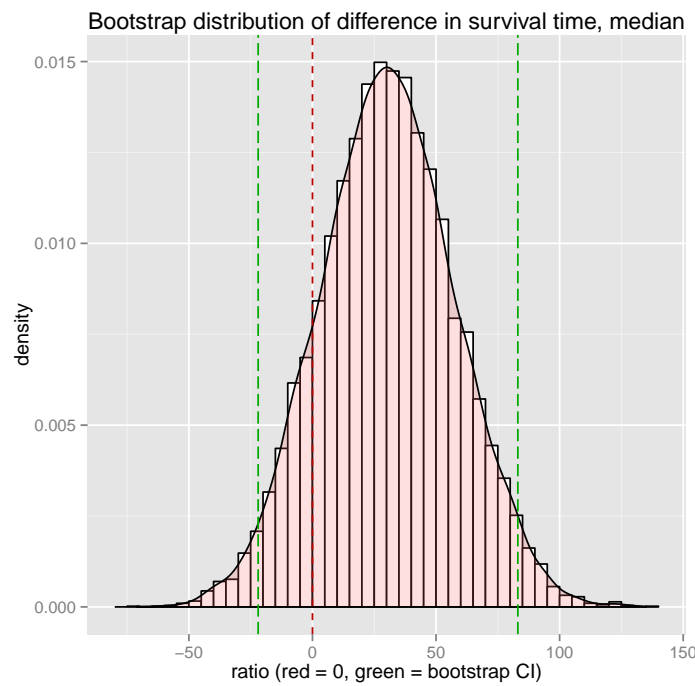
$$\hat{\mu}^* = \bar{x}^* - \bar{y}^*.$$

Repeat this process a large number of times, say 10000 times, and obtain 10000 *bootstrap replicates* $\hat{\mu}^*$. The summaries are in the code, followed by a histogram of bootstrap replicates, $\hat{\mu}^*$.

```
#### Example: Mouse survival, two-sample bootstrap, mean
# draw R bootstrap replicates
R <- 10000
# init location for bootstrap samples
bs1 <- rep(NA, R)
bs2 <- rep(NA, R)
# draw R bootstrap resamples of means
for (i in 1:R) {
  bs2[i] <- mean(sample(control, replace = TRUE))
  bs1[i] <- mean(sample(treatment, replace = TRUE))
}
# bootstrap replicates of difference estimates
bs.diff <- bs1 - bs2
sd(bs.diff)
## [1] 27
# sort the difference estimates to obtain bootstrap CI
diff.sorted <- sort(bs.diff)
# 0.025th and 0.975th quantile gives equal-tail bootstrap CI
CI.bs <- c(diff.sorted[round(0.025*R)], diff.sorted[round(0.975*R+1)])
CI.bs
## [1] -21.97  83.10

## Plot the bootstrap distribution with CI
# First put data in data.frame for ggplot()
dat.diff <- data.frame(bs.diff)

library(ggplot2)
p <- ggplot(dat.diff, aes(x = bs.diff))
p <- p + geom_histogram(aes(y=..density..),
                        , binwidth=5
                        , colour="black", fill="white")
# Overlay with transparent density plot
p <- p + geom_density(alpha=0.2, fill="#FF6666")
# vertical line at 0 and CI
p <- p + geom_vline(xintercept=0, colour="#BB0000", linetype="dashed")
p <- p + geom_vline(xintercept=CI.bs[1], colour="#00AA00", linetype="longdash")
p <- p + geom_vline(xintercept=CI.bs[2], colour="#00AA00", linetype="longdash")
p <- p + labs(title = "Bootstrap distribution of difference in survival time, median")
p <- p + xlab("ratio (red = 0, green = bootstrap CI)")
print(p)
```



Example: Mouse survival, two-sample bootstrap, median For most statistics (such as the median) we don't have a formula for the limiting value of the standard error, but in fact no formula is needed. Instead, we use the numerical output of the bootstrap program. The summaries are in the code, followed by a histogram of bootstrap replicates, $\hat{\eta}^*$.

Group	Data	(n)	Median	est.	SE
Control:	52, 104, 146, 10, 51, 30, 40, 27, 46	(9)	46	?	
Treatment:	94, 197, 16, 38, 99, 141, 23	(7)	94	?	
Difference:			48	?	

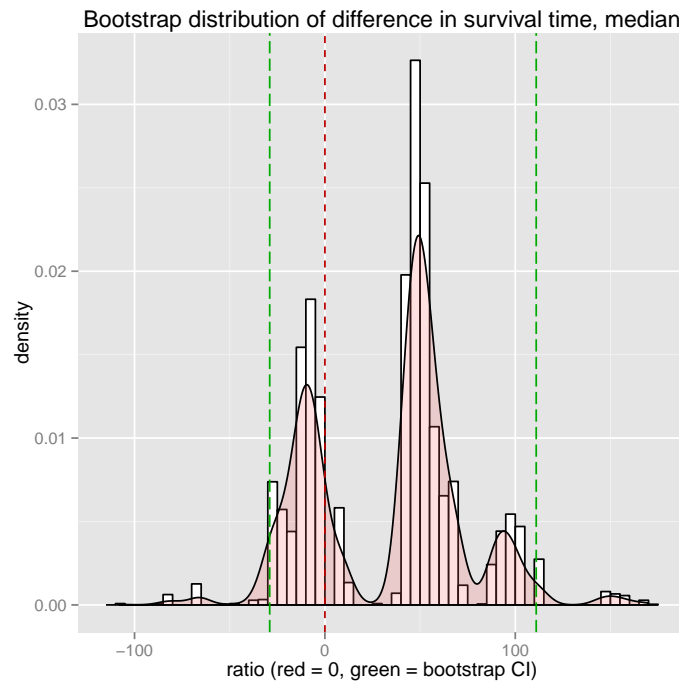
```
#### Example: Mouse survival, two-sample bootstrap, median
sort(control)
## [1] 10 27 30 40 46 51 52 104 146
sort(treatment)
## [1] 16 23 38 94 99 141 197
# draw R bootstrap replicates
R <- 10000
# init location for bootstrap samples
bs1 <- rep(NA, R)
bs2 <- rep(NA, R)
# draw R bootstrap resamples of medians
for (i in 1:R) {
```

```
bs2[i] <- median(sample(control,    replace = TRUE))
bs1[i] <- median(sample(treatment, replace = TRUE))
}
# bootstrap replicates of difference estimates
bs.diff <- bs1 - bs2
sd(bs.diff)
## [1] 40.43

# sort the difference estimates to obtain bootstrap CI
diff.sorted <- sort(bs.diff)
# 0.025th and 0.975th quantile gives equal-tail bootstrap CI
CI.bs <- c(diff.sorted[round(0.025*R)], diff.sorted[round(0.975*R+1)])
CI.bs
## [1] -29 111

## Plot the bootstrap distribution with CI
# First put data in data.frame for ggplot()
dat.diff <- data.frame(bs.diff)

library(ggplot2)
p <- ggplot(dat.diff, aes(x = bs.diff))
p <- p + geom_histogram(aes(y=..density..),
                        , binwidth=5
                        , colour="black", fill="white")
# Overlay with transparent density plot
p <- p + geom_density(alpha=0.2, fill="#FF6666")
# vertical line at 0 and CI
p <- p + geom_vline(xintercept=0, colour="#BB0000", linetype="dashed")
p <- p + geom_vline(xintercept=CI.bs[1], colour="#00AA00", linetype="longdash")
p <- p + geom_vline(xintercept=CI.bs[2], colour="#00AA00", linetype="longdash")
p <- p + labs(title = "Bootstrap distribution of difference in survival time, median")
p <- p + xlab("ratio (red = 0, green = bootstrap CI)")
print(p)
```



9.2.3 Comparing bootstrap sampling distribution from population and sample

Example: Law School, correlation of (LSAT, GPA) The population of average student measurements of (LSAT, GPA) for the universe of 82 law schools are in the table below. Imagine that we don't have all 82 schools worth of data. Consider taking a random sample of 15 schools, indicated by the +'s.

School	LSAT	GPA	School	LSAT	GPA	School	LSAT	GPA
1	622	3.23	28	632	3.29	56	641	3.28
2	542	2.83	29	587	3.16	57	512	3.01
3	579	3.24	30	581	3.17	58	631	3.21
4+	653	3.12	31+	605	3.13	59	597	3.32
5	606	3.09	32	704	3.36	60	621	3.24
6+	576	3.39	33	477	2.57	61	617	3.03
7	620	3.10	34	591	3.02	62	637	3.33
8	615	3.40	35+	578	3.03	62	572	3.08
9	553	2.97	36+	572	2.88	64	610	3.13
10	607	2.91	37	615	3.37	65	562	3.01
11	558	3.11	38	606	3.20	66	635	3.30
12	596	3.24	39	603	3.23	67	614	3.15
13+	635	3.30	40	535	2.98	68	546	2.82
14	581	3.22	41	595	3.11	69	598	3.20
15+	661	3.43	42	575	2.92	70+	666	3.44
16	547	2.91	43	573	2.85	71	570	3.01
17	599	3.23	44	644	3.38	72	570	2.92
18	646	3.47	45+	545	2.76	73	605	3.45
19	622	3.15	46	645	3.27	74	565	3.15
20	611	3.33	47+	651	3.36	75	686	3.50
21	546	2.99	48	562	3.19	76	608	3.16
22	614	3.19	49	609	3.17	77	595	3.19
23	628	3.03	50+	555	3.00	78	590	3.15
24	575	3.01	51	586	3.11	79+	558	2.81
25	662	3.39	52+	580	3.07	80	611	3.16
26	627	3.41	53+	594	2.96	81	564	3.02
27	608	3.04	54	594	3.05	82+	575	2.74
			55	560	2.93			

```
#### Example: Law School, correlation of (LSAT, GPA)
```

```
School <- 1:82
```

```
LSAT <- c(622, 542, 579, 653, 606, 576, 620, 615, 553, 607, 558, 596, 635,
      581, 661, 547, 599, 646, 622, 611, 546, 614, 628, 575, 662, 627,
      608, 632, 587, 581, 605, 704, 477, 591, 578, 572, 615, 606, 603,
      535, 595, 575, 573, 644, 545, 645, 651, 562, 609, 555, 586, 580,
      594, 594, 560, 641, 512, 631, 597, 621, 617, 637, 572, 610, 562,
      635, 614, 546, 598, 666, 570, 570, 605, 565, 686, 608, 595, 590,
      558, 611, 564, 575)
```

```
GPA <- c(3.23, 2.83, 3.24, 3.12, 3.09, 3.39, 3.10, 3.40, 2.97, 2.91, 3.11,
       3.24, 3.30, 3.22, 3.43, 2.91, 3.23, 3.47, 3.15, 3.33, 2.99, 3.19,
       3.03, 3.01, 3.39, 3.41, 3.04, 3.29, 3.16, 3.17, 3.13, 3.36, 2.57,
       3.02, 3.03, 2.88, 3.37, 3.20, 3.23, 2.98, 3.11, 2.92, 2.85, 3.38,
       2.76, 3.27, 3.36, 3.19, 3.17, 3.00, 3.11, 3.07, 2.96, 3.05, 2.93,
```

```

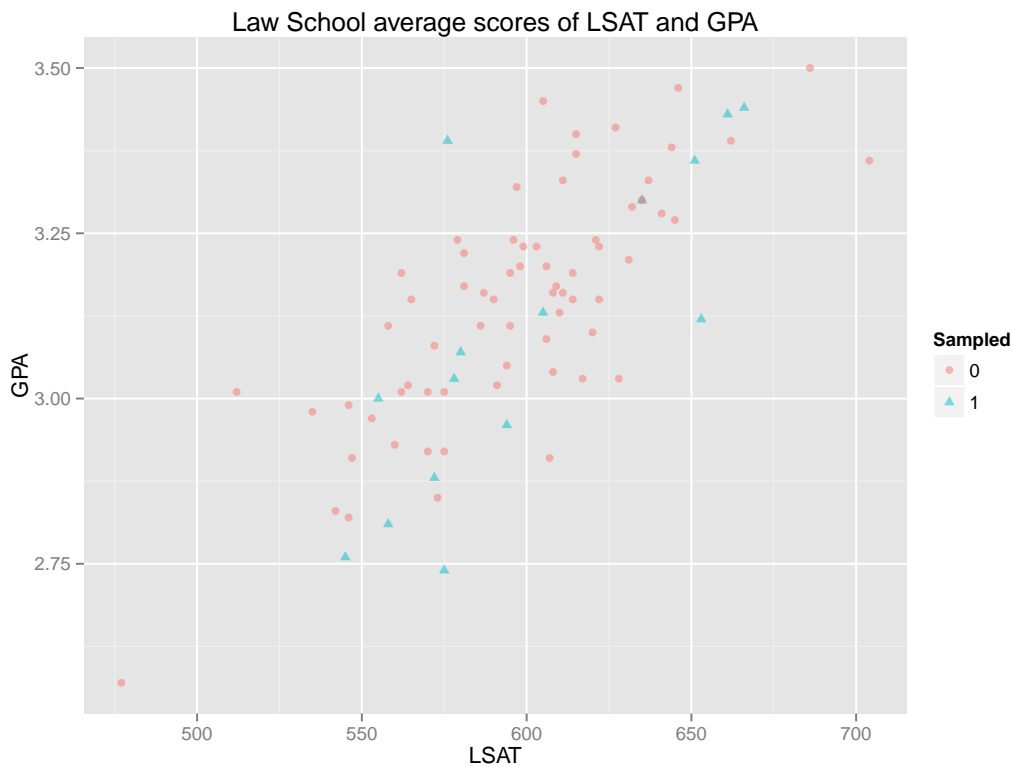
3.28, 3.01, 3.21, 3.32, 3.24, 3.03, 3.33, 3.08, 3.13, 3.01, 3.30,
3.15, 2.82, 3.20, 3.44, 3.01, 2.92, 3.45, 3.15, 3.50, 3.16, 3.19,
3.15, 2.81, 3.16, 3.02, 2.74)

Sampled <- c(0, 0, 0, 1, 0, 0, 0, 0, 0, 0, 1, 0, 1, 0, 0, 0, 0, 0, 0,
0, 0, 0, 0, 0, 0, 1, 0, 0, 0, 1, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0,
1, 0, 1, 0, 0, 1, 0, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0,
0, 0, 0, 1, 0, 0, 0, 0, 0, 0, 0, 1, 0, 0, 1)

# law = population
law <- data.frame(School, LSAT, GPA, Sampled)
law$Sampled <- factor(law$Sampled)
# law.sam = sample
law.sam <- subset(law, Sampled == 1)

library(ggplot2)
p <- ggplot(law, aes(x = LSAT, y = GPA))
p <- p + geom_point(aes(colour = Sampled, shape = Sampled), alpha = 0.5, size = 2)
p <- p + labs(title = "Law School average scores of LSAT and GPA")
print(p)

```



Let's bootstrap the sample of 15 observations to get the bootstrap sampling distribution of correlation (for sampling 15 from the population). From the bootstrap sampling distribution we'll calculate a bootstrap confidence interval for the true population correlation, as well as a bootstrap standard deviation for the correlation. But how well does this work? Let's compare it against the *true* sampling distribution by drawing 15 random schools from the population of 82

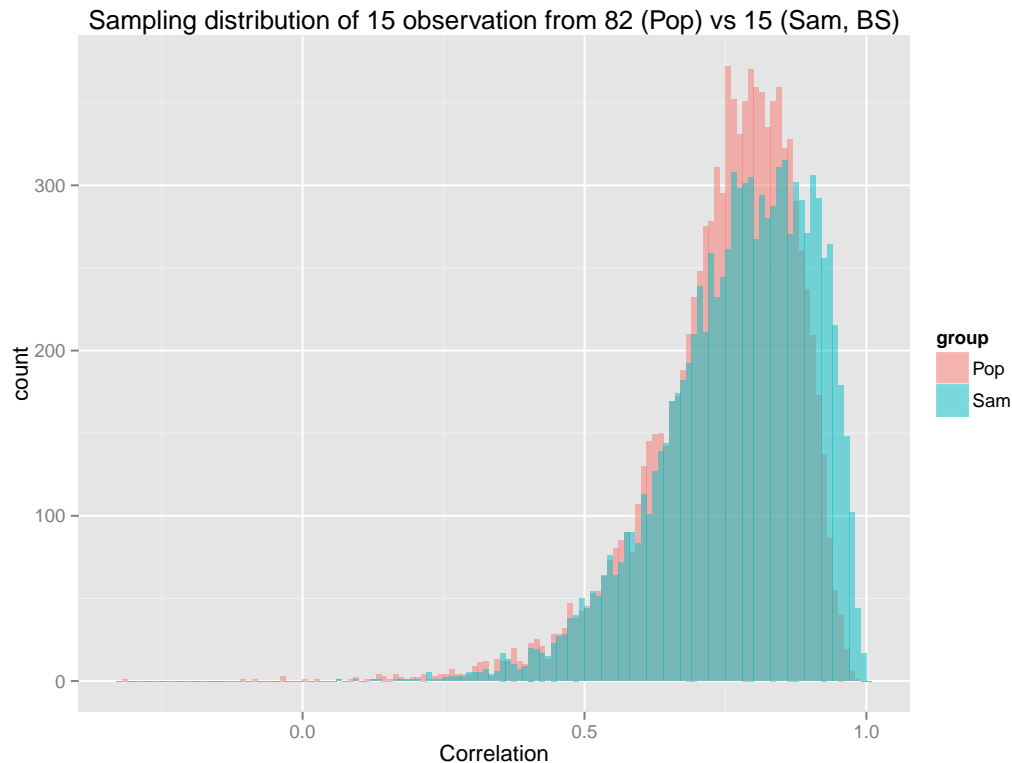
schools and calculating the correlation. If the bootstrap works well (from our hopefully representative sample of 15), then the bootstrap sampling distribution from the 15 schools will be close to the true sampling distribution.

The code below does that, followed by two histograms. In this case, the histograms are noticeably non-normal, having a long tail toward the left. Inferences based on the normal curve are suspect when the bootstrap histogram is markedly non-normal. The histogram on the left is the nonparametric bootstrap sampling distribution using only the $n = 15$ sampled schools with 10000 bootstrap replicates of $\widehat{\text{corr}}(x^*)$. The histogram on the right is the true sampling distribution using 10000 replicates of $\widehat{\text{corr}}(x^*)$ from the population of law school data, repeatedly drawing $n = 15$ without replacement from the $N = 82$ points. Impressively, the bootstrap histogram on the left strongly resembles the population histogram on the right. Remember, in a real problem we would only have the information on the left, from which we would be trying to infer the situation on the right.

```
# draw R bootstrap replicates
R <- 10000
# init location for bootstrap samples
bs.pop <- rep(NA, R)
bs.sam <- rep(NA, R)
# draw R bootstrap resamples of medians
for (i in 1:R) {
  # sample() draws indices then bootstrap correlation of LSAT and GPA
  # population
  bs.pop[i] = cor(law      [sample(seq(1,nrow(law      )), nrow(law.sam)
                                    , replace = TRUE), 2:3])[1, 2]
  # sample
  bs.sam[i] = cor(law.sam[sample(seq(1,nrow(law.sam)), nrow(law.sam)
                                    , replace = TRUE), 2:3])[1, 2]
}
# sort the difference estimates to obtain bootstrap CI
diff.sorted <- sort(bs.pop)
# 0.025th and 0.975th quantile gives equal-tail bootstrap CI
CI.bs.pop <- c(diff.sorted[round(0.025*R)], diff.sorted[round(0.975*R+1)])
# population correlation
cor(law      [, c(2,3)])[1,2]
## [1] 0.76
CI.bs.pop
## [1] 0.4297 0.9271
sd(bs.pop)
## [1] 0.1295
```

```
# sort the difference estimates to obtain bootstrap CI
diff.sorted <- sort(bs.sam)
# 0.025th and 0.975th quantile gives equal-tail bootstrap CI
CI.bs.sam <- c(diff.sorted[round(0.025*R)], diff.sorted[round(0.975*R+1)])
# sample correlation
cor(law.sam[, c(2,3)][1,2]
## [1] 0.7764
CI.bs.sam
## [1] 0.4638 0.9638
sd(bs.sam)
## [1] 0.1335

law.bs.df <- data.frame(corr = c(bs.pop, bs.sam), group = c(rep("Pop",R),rep("Sam",R)))
# histogram using ggplot
library(ggplot2)
p <- ggplot(law.bs.df, aes(x = corr, fill=group))
p <- p + geom_histogram(binwidth = .01, alpha = 0.5, position="identity")
p <- p + labs(title = "Sampling distribution of 15 observation from 82 (Pop) vs 15 (Sam, BS)",
               xlab("Correlation"))
print(p)
```



Chapter 10

Power and Sample size

Learning objectives

After completing this topic, you should be able to:

- assess** the power of a test or
- determin** the required sample size for a study.

Achieving these goals contributes to mastery in these course learning outcomes:

- 7. Distinguish between statistical significance and scientific relevance.
- 10. Identify and explain the statistical methods, assumptions, and limitations.
- 12. Make evidence-based decisions by constructing and deciding between testable hypotheses using appropriate data and methods.

10.1 Power Analysis

The meaning of statistical power *Power is the probability $(1 - \beta)$ of detecting an effect, given that the effect is really there.* In other words, it is the probability of correctly rejecting the null hypothesis when it is in fact false. For example, let's say that we have a simple study with drug A and a placebo group, and that the drug truly is effective; the power is the probability of finding a difference between the two groups. So, imagine that we had a power of $1 - \beta = 0.8$ and that this simple study was conducted many times. Having power of 0.8 means that 80% of the time, we would get a statistically significant difference between the drug A and placebo groups. This also means that 20%

of the times that we run this experiment, we will not obtain a statistically significant effect between the two groups, even though there really is an effect in reality. That is, the probability of a Type-II error is $\beta = 0.2$.

One-sample power figure Consider the plot below for a one-sample one-tailed greater-than t -test. If the null hypothesis, $H_0 : \mu = \mu_0$, is true, then the test statistic t follows the null distribution indicated by the hashed area. Under a specific alternative hypothesis, $H_1 : \mu = \mu_1$, the test statistic t follows the distribution indicated by the solid area. If α is the probability of making a Type-I error (rejecting H_0 when it is true), then “crit. val.” indicates the location of the t_{crit} value associated with H_0 on the scale of the data. The rejection region is the area under H_0 that is at least as far as “crit. val.” is from μ_0 . The power ($1 - \beta$) of the test is the green area, the area under H_1 in the rejection region. A Type-II error is made when H_1 is true, but we fail to reject H_0 in the red region. (Note, for a two-tailed test the rejection region for both tails under the H_1 curve contribute to the power.)

```
#### One-sample power
# Power plot with two normal distributions
# http://stats.stackexchange.com/questions/14140/how-to-best-display-graphically-type-ii-beta-1

x <- seq(-4, 4, length=1000)
hx <- dnorm(x, mean=0, sd=1)

plot(x, hx, type="n", xlim=c(-4, 8), ylim=c(0, 0.5),
ylab = "",
xlab = "",
main= expression(paste("Type-II Error (", beta, ") and Power (", 1-beta, ")")), axes=FALSE)

#shift = qnorm(1-0.025, mean=0, sd=1)*1.7
shift = qnorm(1-0.05, mean=0, sd=1)*1.7 # one-tailed
xfit2 <- x + shift
yfit2 <- dnorm(xfit2, mean=shift, sd=1)

#axis(1, at = c(-qnorm(.025), 0, shift, -4),
#      labels = expression("p-value", 0, mu, -infinity ))
#axis(1, at = c(-qnorm(.025), 0, shift),
#      labels = expression((t[alpha/2]), mu[0], mu[1]))
axis(1, at = c(-qnorm(.05), 0, shift),
      labels = expression("crit. val.", mu[0], mu[1]))
axis(1, at = c(-4, 4+shift),
      labels = expression(-infinity, infinity ), lwd=1, lwd.tick=FALSE)

## Print null hypothesis area
```

```
#col_null = "#DDDDDD"
#polygon(c(min(x), x,max(x)), c(0,hx,0), col=col_null)
#lines(x, hx, lwd=2)
col_null = "#AAAAAA"
polygon(c(min(x), x,max(x)), c(0,hx,0), col=col_null, lwd=2, density=c(10, 40), angle=-45, border=1)
lines(x, hx, lwd=2, lty="dashed", col=col_null)

# The alternative hypothesis area

## The red - underpowered area
lb <- min(xfit2)
#ub <- round(qnorm(.975),2)
ub <- round(qnorm(.95),2)
col1 = "#CC2222"

i <- xfit2 >= lb & xfit2 <= ub
polygon(c(lb,xfit2[i],ub), c(0,yfit2[i],0), col=col1)

## The green area where the power is
col2 = "#22CC22"
i <- xfit2 >= ub
polygon(c(ub,xfit2[i],max(xfit2)), c(0,yfit2[i],0), col=col2)

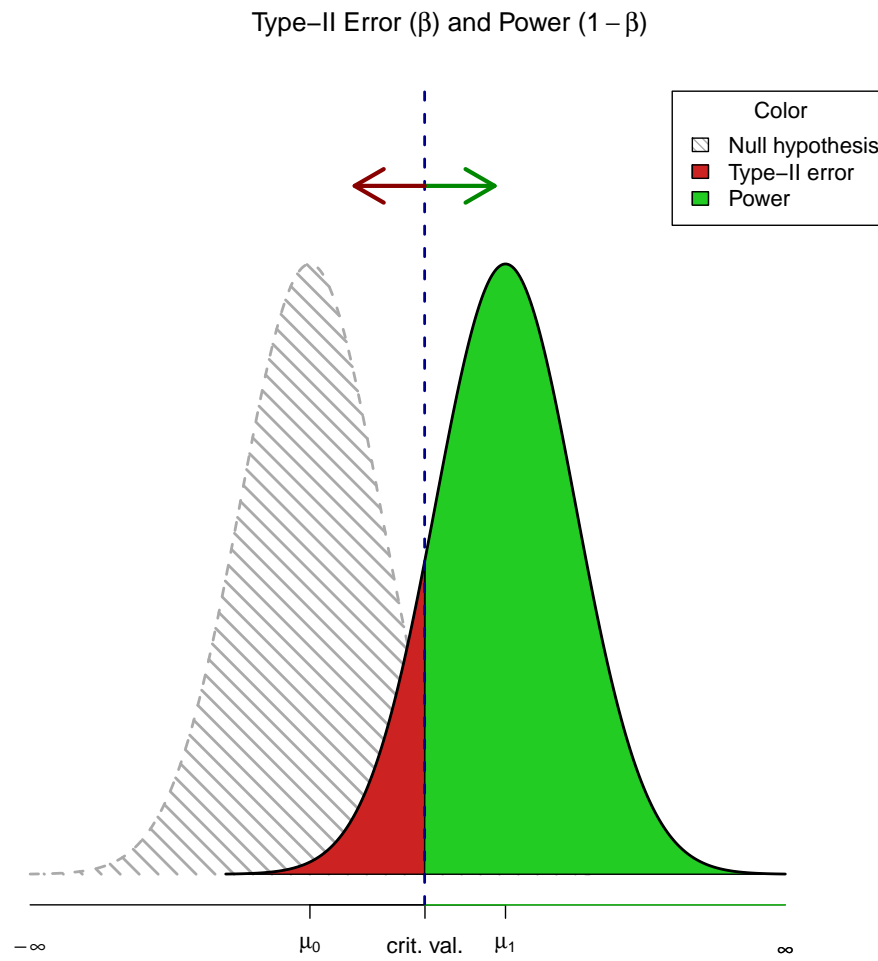
# Outline the alternative hypothesis
lines(xfit2, yfit2, lwd=2)

axis(1, at = (c(ub, max(xfit2))), labels=c("", expression(infinity)),
      col=col2, lwd=1, lwd.tick=FALSE)

#legend("topright", inset=.05, title="Color",
#       c("Null hypotheses","Type II error", "True"), fill=c(col_null, col1, col2), horiz=FALSE)
legend("topright", inset=.015, title="Color",
       c("Null hypothesis","Type-II error", "Power"), fill=c(col_null, col1, col2),
       angle=-45,
       density=c(20, 1000, 1000), horiz=FALSE)

abline(v=ub, lwd=2, col="#000088", lty="dashed")

arrows(ub, 0.45, ub+1, 0.45, lwd=3, col="#008800")
arrows(ub, 0.45, ub-1, 0.45, lwd=3, col="#880000")
```



Example: IQ drug Imagine that we are evaluating the effect of a putative memory enhancing drug. We have randomly sampled 25 people from a population known to be normally distributed with a μ of 100 and a σ of 15. We administer the drug, wait a reasonable time for it to take effect, and then test our subjects' IQ. Assume that we were so confident in our belief that the drug would either increase IQ or have no effect that we entertained one-sided (directional) hypotheses. Our null hypothesis is that after administering the drug $\mu \leq 100$ and our alternative hypothesis is $\mu > 100$.

These hypotheses must first be converted to exact hypotheses. Converting the null is easy: it becomes $\mu = 100$. The alternative is more troublesome. If we knew that the effect of the drug was to increase IQ by 15 points, our exact alternative hypothesis would be $\mu = 115$, and we could compute power, the probability of correctly rejecting the false null hypothesis given that μ is really equal to 115 after drug treatment, not 100 (normal IQ). But if we already

knew how large the effect of the drug was, we would not need to do inferential statistics...

One solution is to decide on a **minimum nontrivial effect size**. What is the smallest effect that you would consider to be nontrivial? Suppose that you decide that if the drug increases μ_{IQ} by 2 or more points, then that is a nontrivial effect, but if the mean increase is less than 2 then the effect is trivial.

Now we can test the null of $\mu = 100$ versus the alternative of $\mu = 102$. Consider the previous plot. Let the left curve represent the distribution of sample means if the null hypothesis were true, $\mu = 100$. This sampling distribution has a $\mu = 100$ and a $\sigma_{\bar{Y}} = 15/\sqrt{25} = 3$. Let the right curve represent the sampling distribution if the exact alternative hypothesis is true, $\mu = 102$. Its μ is 102 and, assuming the drug has no effect on the variance in IQ scores, also has $\sigma_{\bar{Y}} = 3$.

The green area in the upper tail of the null distribution (gray hatched curve) is α . Assume we are using a one-tailed α of 0.05. How large would a sample mean need be for us to reject the null? Since the upper 5% of a normal distribution extends from 1.645σ above the μ up to positive infinity, the sample mean IQ would need be $100 + 1.645(3) = 104.935$ or more to reject the null. What are the chances of getting a sample mean of 104.935 or more if the alternative hypothesis is correct, if the drug increases IQ by 2 points? The area under the alternative curve from 104.935 up to positive infinity represents that probability, which is power. Assuming the alternative hypothesis is true, that $\mu = 102$, the probability of rejecting the null hypothesis is the probability of getting a sample mean of 104.935 or more in a normal distribution with $\mu = 102$, $\sigma = 3$. $Z = (104.935 - 102)/3 = 0.98$, and $P(Z > 0.98) = 0.1635$. That is, power is about 16%. If the drug really does increase IQ by an average of 2 points, we have a 16% chance of rejecting the null. If its effect is even larger, we have a greater than 16% chance.

Suppose we consider 5 (rather than 2) the minimum nontrivial effect size. This will separate the null and alternative distributions more, decreasing their overlap and increasing power. Now, $Z = (104.935 - 105)/3 = -0.02$, $P(Z > -0.02) = 0.5080$ or about 51%. **It is easier to detect large effects than small effects.**

Suppose we conduct a 2-tailed test, since the drug could actually decrease IQ; α is now split into both tails of the null distribution, 0.025 in each tail. We shall reject the null if the sample mean is 1.96 or more standard errors away from the μ of the null distribution. That is, if the mean is $100 + 1.96(3) = 105.88$ or more (or if it is $100 - 1.96(3) = 94.12$ or less) we reject the null. The probability of that happening if the alternative is correct ($\mu = 105$) is: $Z = (105.88 - 105)/3 = 0.29$, $P(Z > 0.29) = 0.3859$, and $P(Z < (94.12 - 105)/3) = P(Z < -3.63) = 0.00014$, for a total power = $(1 - \beta) = 0.3859 + 0.00014$, or about 39%. Note that our power is less than it was with a one-tailed test. **If you can correctly predict the direction of effect, a one-tailed test is more powerful than a two-tailed test.**

Consider what would happen if you increased sample size to 100. Now the $\sigma_{\bar{Y}} = 15/\sqrt{100} = 1.5$. With the null and alternative distributions are narrower, and should overlap less, increasing power. With $\sigma_{\bar{Y}} = 1.5$ the sample mean will need be $100 + (1.96)(1.5) = 102.94$ (rather than 105.88 from before) or more to reject the null. If the drug increases IQ by 5 points, power is: $Z = (102.94 - 105)/1.5 = -1.37$, $P(Z > -1.37) = 0.9147$, or between 91 and 92%. **Anything that decreases the standard error will increase power. This may be achieved by increasing the sample size N or by reducing the σ of the dependent variable.** The σ of the dependent variable may be reduced by reducing the influence of extraneous variables upon the dependent variable (eliminating “noise” in the dependent variable makes it easier to detect the signal).

Now consider what happens if you change the significance level, α . Let us reduce α to 0.01. Now the sample mean must be 2.58 or more standard errors from the null μ before we reject the null. That is, $100 + 2.58(1.5) = 103.87$ (rather than 102.94 with $\alpha = 0.05$). Under the alternative, $Z = (103.87 - 105)/1.5 = -0.75$, $P(Z > -0.75) = 0.7734$ or about 77%, less than it was with $\alpha = 0.05$. **Reducing α reduces power.**

Please note that all of the above analyses have assumed that we have used a normally distributed test statistic, as $Z = (\bar{Y} - \mu_0)/\sigma_{\bar{Y}}$ will be if the dependent variable is normally distributed in the population or if sample size is large enough to invoke the central limit theorem (CLT). Remember that using Z

also requires that you know the population σ rather than estimating it from the sample data. We more often estimate the population σ , using Student's t as the test statistic. If N is fairly large, Student's t is nearly normal, so this is no problem. For example, with a two-tailed $\alpha = 0.05$ and $N = 25$, we went out ± 1.96 standard errors to mark off the rejection region. With Student's t on $N - 1 = 24$ df we should have gone out ± 2.064 standard errors. But 1.96 versus 2.06 is a relatively trivial difference, so we should feel comfortable with the normal approximation. If, however, we had $N = 5$, $df = 4$, critical $t = \pm 2.776$, then the normal approximation would not do. A more complex analysis would be needed.

10.2 Effect size

For the one-sample test, the effect size in σ units is $d = (\mu_1 - \mu_0)/\sigma$. For our IQ problem with minimum nontrivial effect size at 5 IQ points, $d = (105 - 100)/15 = 1/3$. Cohen's¹ conventions for small, medium, and large effects for a two-sample difference test between two means is in the table below.

One- or two-sample difference of means		
Size of effect	d	% variance
small	0.2	1
medium	0.5	6
large	0.8	16

Cohen has conventions for other tests (correlation, contingency tables, etc.), but they should be used with caution.

What is a small or even trivial effect in one context may be a large effect in another context. For example, Rosnow and Rosenthal (1989) discussed a 1988 biomedical research study on the effects of taking a small, daily dose of aspirin. Each participant was instructed to take one pill a day. For about half of the participants the pill was aspirin, for the others it was a placebo. The dependent variable was whether or not the participant had a heart attack during the study. In terms of a correlation coefficient, the size of the observed

¹Cohen, J. (1988). Statistical power analysis for the behavior sciences. (2nd ed.). Hillsdale, NJ: Erlbaum.

effect was $r = 0.034$. In terms of percentage of variance explained, that is 0.12%. In other contexts this might be considered a trivial effect, but in this context it was so large an effect that the researchers decided it was unethical to continue the study and they contacted all of the participants who were taking the placebo and told them to start taking aspirin every day.

10.3 Sample size

Before you can answer the question "how many subjects do I need," you will have to answer several other questions, such as:

- How much power do I want?
- What is the likely size (in the population) of the effect I am trying to detect, or, what is smallest effect size that I would consider of importance?
- What criterion of statistical significance will I employ?
- What test statistic will I employ?
- What is the standard deviation (in the population) of the criterion variable?
- For correlated samples designs, what is the correlation (in the population) between groups?

If one considers Type I and Type II errors equally serious, then one should have enough power to make $\alpha = \beta$. If employing the traditional 0.05 criterion of statistical significance, that would mean you should have 95% power. However, getting 95% power usually involves expenses too great – that is, too many samples.

A common convention is to try to get at least enough data to have 80% power. So, how do you figure out how many subjects you need to have the desired amount of power. There are several methods, including:

- You could buy an expensive, professional-quality software package to do the power analysis.
- You could buy an expensive, professional-quality book on power analysis and learn to do the calculations yourself and/or to use power tables and figures to estimate power.
- You could try to find an interactive web page on the Internet that will do

the power analysis for you. This is probably fine, but be cautious.

- You could download and use the G Power program, which is free, not too difficult to use, and generally reliable (this is not to say that it is error free).
- You could use the simple guidelines provided in Jacob Cohen's "A Power Primer" (Psychological Bulletin, 1992, 112, 155-159).

The plots below indicate the amount of power for a given effect size and sample size for a one-sample t -test and ANOVA test. This graph makes clear the diminishing returns you get for adding more and more subjects if you already have moderate to high power. For example, let's say we're doing a one-sample test and we an effect size of 0.2 and have only 10 subjects. We can see that we have a power of about 0.15, which is really, really low. Going to 25 subjects increases our power to about 0.25, and to 100 subjects increases our power to about 0.6. But if we had a large effect size of 0.8, 10 subjects would already give us a power of about 0.8, and using 25 or 100 subjects would both give a power at least 0.98. So each additional subject gives you less additional power. This curve also illustrates the "cost" of increasing your desired power from 0.8 to 0.98.

```
# Power curve plot for one-sample t-test with range of sample sizes
# http://stackoverflow.com/questions/4680163/power-vs-effect-size-plot/4680786#4680786

P      <- 3                                # number of groups for ANOVA
fVals <- seq(0, 1.2, length.out=100)        # effect sizes f for ANOVA
dVals <- seq(0, 3, length.out=100)          # effect sizes d for t-Test
#nn    <- seq(10, 25, by=5)                 # group sizes
nn     <- c(5,10,25,100)                   # group sizes
alpha <- 0.05                               # test for level alpha

# function to calculate one-way ANOVA power for given group size
getFPow <- function(n) {
  critF <- qf(1-alpha, P-1, P*n - P) # critical F-value

  # probabilities of exceeding this F-value given the effect sizes f
  # P*n*fVals^2 is the non-centrality parameter
  1-pf(critF, P-1, P*n - P, P*n * fVals^2)
}

# function to calculate one-sample t-Test power for given group size
getTPow <- function(n) {
  critT <- qt(1-alpha, n-1)                # critical t-value

  # probabilities of exceeding this t-value given the effect sizes d
```

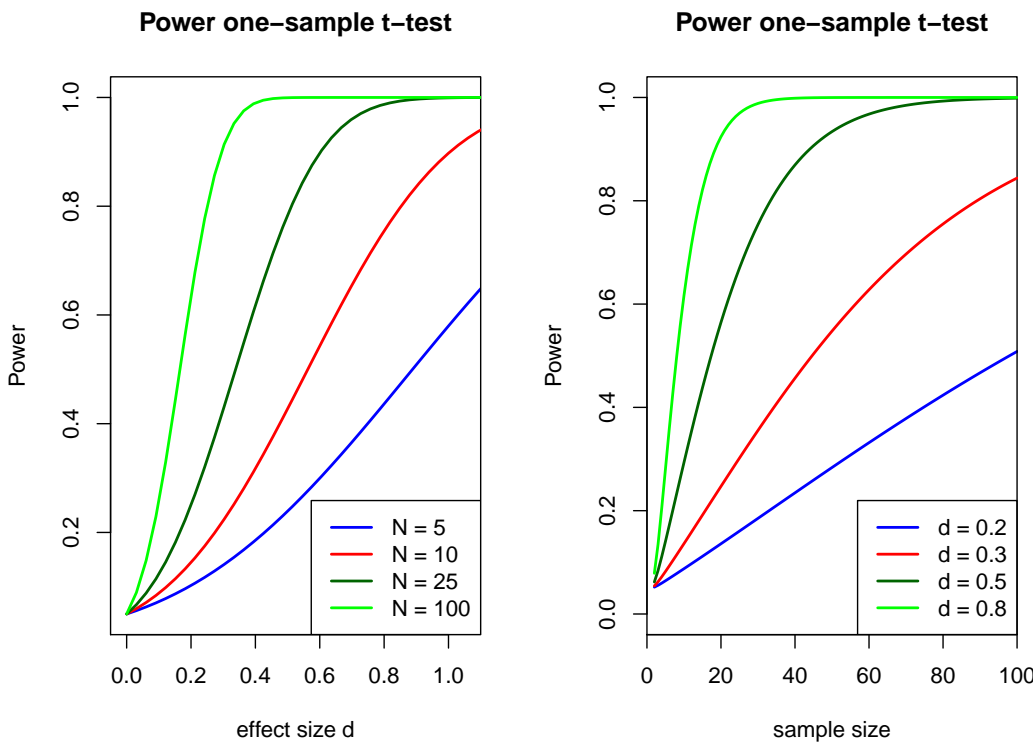
```
# sqrt(n)*d is the non-centrality parameter
1-pt(critT, n-1, sqrt(n)*dVals)
}

powsF <- sapply(nn, getFPow)      # ANOVA power for all group sizes
powsT <- sapply(nn, getTPow)      # t-Test power for all group sizes

#dev.new(width=10, fig.height=5)
par(mfrow=c(1, 2))
matplot(dVals, powsT, type="l", lty=1, lwd=2, xlab="effect size d",
         ylab="Power", main="Power one-sample t-test", xaxs="i",
         xlim=c(-0.05, 1.1), col=c("blue", "red", "darkgreen", "green"))
#legend(x="bottomright", legend=paste("N =", c(5,10,25,100)), lwd=2,
#       col=c("blue", "red", "darkgreen", "green"))
legend(x="bottomright", legend= paste("N =", nn), lwd=2,
       col=c("blue", "red", "darkgreen", "green"))

#matplot(fVals, powsF, type="l", lty=1, lwd=2, xlab="effect size f",
#         ylab="Power", main=paste("Power one-way ANOVA, ", P, " groups", sep=""), xaxs="i",
#         xlim=c(-0.05, 1.1), col=c("blue", "red", "darkgreen", "green"))
##legend(x="bottomright", legend= paste("Nj =", c(10, 15, 20, 25)), lwd=2,
##       col=c("blue", "red", "darkgreen", "green"))
#legend(x="bottomright", legend= paste("Nj =", nn), lwd=2,
#       col=c("blue", "red", "darkgreen", "green"))

library(pwr)
pwrt2 <- pwr.t.test(d=.2,n=seq(2,100,1),
                      sig.level=.05,type="one.sample", alternative="two.sided")
pwrt3 <- pwr.t.test(d=.3,n=seq(2,100,1),
                      sig.level=.05,type="one.sample", alternative="two.sided")
pwrt5 <- pwr.t.test(d=.5,n=seq(2,100,1),
                      sig.level=.05,type="one.sample", alternative="two.sided")
pwrt8 <- pwr.t.test(d=.8,n=seq(2,100,1),
                      sig.level=.05,type="one.sample", alternative="two.sided")
#plot(pwrt$n, pwrt$power, type="b", xlab="sample size", ylab="power")
matplot(matrix(c(pwrt2$n,pwrt3$n,pwrt5$n,pwrt8$n),ncol=4),
        matrix(c(pwrt2$power,pwrt3$power,pwrt5$power,pwrt8$power),ncol=4),
        type="l", lty=1, lwd=2, xlab="sample size",
        ylab="Power", main="Power one-sample t-test", xaxs="i",
        xlim=c(0, 100), ylim=c(0,1), col=c("blue", "red", "darkgreen", "green"))
legend(x="bottomright", legend= paste("d =", c(0.2, 0.3, 0.5, 0.8)), lwd=2,
       col=c("blue", "red", "darkgreen", "green"))
```



Reasons to do a power analysis There are several of reasons why one might do a power analysis. (1) Perhaps the most common use is to determine the necessary number of subjects needed to detect an effect of a given size. Note that trying to find the absolute, bare minimum number of subjects needed in the study is often not a good idea. (2) Additionally, power analysis can be used to determine power, given an effect size and the number of subjects available. You might do this when you know, for example, that only 75 subjects are available (or that you only have the budget for 75 subjects), and you want to know if you will have enough power to justify actually doing the study. In most cases, there is really no point to conducting a study that is seriously underpowered. Besides the issue of the number of necessary subjects, there are other good reasons for doing a power analysis. (3) For example, a power analysis is often required as part of a grant proposal. (4) And finally, doing a power analysis is often just part of doing good research. A power analysis is a good way of making sure that you have thought through every aspect of the study and the statistical analysis before you start collecting data.

Limitations Despite these advantages of power analyses, there are some limitations. (1) One limitation is that power analyses do not typically generalize very well. If you change the methodology used to collect the data or change the statistical procedure used to analyze the data, you will most likely have to redo the power analysis. (2) In some cases, a power analysis might suggest a number of subjects that is inadequate for the statistical procedure. For example (beyond the scope of this class), a power analysis might suggest that you need 30 subjects for your logistic regression, but logistic regression, like all maximum likelihood procedures, require much larger sample sizes. (3) Perhaps the most important limitation is that a standard power analysis gives you a “best case scenario” estimate of the necessary number of subjects needed to detect the effect. In most cases, this “best case scenario” is based on assumptions and educated guesses. If any of these assumptions or guesses are incorrect, you may have less power than you need to detect the effect. (4) Finally, because power analyses are based on assumptions and educated guesses, you often get a range of the number of subjects needed, not a precise number. For example, if you do not know what the standard deviation of your outcome measure will be, you guess at this value, run the power analysis and get X number of subjects. Then you guess a slightly larger value, rerun the power analysis and get a slightly larger number of necessary subjects. You repeat this process over the plausible range of values of the standard deviation, which gives you a range of the number of subjects that you will need.

Other considerations After all of this discussion of power analyses and the necessary number of subjects, we need to stress that power is not the only consideration when determining the necessary sample size. For example, different researchers might have different reasons for conducting a regression analysis. (1) One might want to see if the regression coefficient is different from zero, (2) while the other wants to get a very precise estimate of the regression coefficient with a very small confidence interval around it. This second purpose requires a larger sample size than does merely seeing if the regression coefficient is different from zero. (3) Another consideration when determining the necessary sample size is the assumptions of the statistical procedure that is going to be used (e.g.,

parametric vs nonparametric procedure). (4) The number of statistical tests that you intend to conduct will also influence your necessary sample size: the more tests that you want to run, the more subjects that you will need (multiple comparisons). (5) You will also want to consider the representativeness of the sample, which, of course, influences the generalizability of the results. Unless you have a really sophisticated sampling plan, the greater the desired generalizability, the larger the necessary sample size.

10.4 Power calculation via simulation

Using the principles of the bootstrap (to be covered later) we can estimate statistical power through simulation.

Example: IQ drug, revisited Recall that we sample $N = 25$ people from a population known to be normally distributed with a μ of 100 and a σ of 15. Consider the first one-sided alternative $H_0 : \mu = 100$ and $H_1 : \mu > 100$. Assume the *minimum nontrivial effect size* was that the drug increases μ_{IQ} by 2 or more points, so that the specific alternative to consider is $H_1 : \mu = 102$. What is the power of this test?

We already saw how to calculate this analytically. To solve this computationally, we need to simulate samples of $N = 25$ from the alternative distribution ($\mu = 102$ and $\sigma = 15$) and see what proportion of the time we correctly reject H_0 .

```
#### Example: IQ drug, revisited
# R code to simulate one-sample one-sided power

# Strategy:
# Do this R times:
#   draw a sample of size N from the distribution specified by the alternative hypothesis
#   That is, 25 subjects from a normal distribution with mean 102 and sigma 15
#   Calculate the mean of our sample
#   Calculate the associated z-statistic
#   See whether that z-statistic has a p-value < 0.05 under H0: mu=100
#   If we reject H0, then set reject = 1, else reject = 0.
# Finally, the proportion of rejects we observe is the approximate power

n      <- 25;          # sample size of 25
mu0   <- 100;         # null hypothesis mean of 100
```

```

mu1    <- 102;          # alternative mean of 102
#mu1    <- 105;          # alternative mean of 105
sigma <- 15;            # standard deviation of normal population

alpha <- 0.05;           # significance level

R      <- 10000;         # Repetitions to draw sample and see whether we reject H0
#   The proportion of these that reject H0 is the power

reject <- rep(NA, R);   # allocate a vector of length R with missing values (NA)
#   to fill with 0 (fail to reject H0) or 1 (reject H0)

for (i in 1:R) {
  sam <- rnorm(n, mean=mu1, sd=sigma);  # sam is a vector with 25 values

  ybar <- mean(sam); # Calculate the mean of our sample sam

  z <- (ybar - mu0) / (sigma / sqrt(n)); # z-statistic (assumes we know sigma)
#   we could also have calculated the t-statistic, here

  pval <- 1-pnorm(z); # one-sided right-tail p-value
#   pnorm gives the area to the left of z
#   therefore, the right-tail area is 1-pnorm(z)

  if (pval < 0.05) {
    reject[i] <- 1; # 1 for correctly rejecting H0
  } else {
    reject[i] <- 0; # 0 for incorrectly fail to reject H0
  }
}

power <- mean(reject); # the average reject (proportion of rejects) is the power
power
## [1] 0.166
# 0.1655 for mu1=102
# 0.5082 for mu1=105

```

Our simulation (this time) with $\mu_1 = 102$ gave a power of 0.166 (exact answer is $P(Z > 0.98) = 0.1635$). Rerunning with $\mu_1 = 105$ gave a power of 0.5082 (exact answer is $P(Z > -0.02) = 0.5080$). Our simulation well-approximates the true value, and the power can be made more precise by increasing the number of repetitions calculated. However, two to three decimal precision is quite sufficient.

Example: Head breadth Recall the head breadth example in Chapter 3 comparing maximum head breadths (in millimeters) of modern day Englishmen with ancient Celts. The data are summarized below.

Descriptive Statistics: ENGLISH, CELTS

Variable	N	Mean	SE Mean	StDev	Minimum	Q1	Median	Q3	Maximum
ENGLISH	18	146.50	1.50	6.38	132.00	141.75	147.50	150.00	158.00
CELTS	16	130.75	1.36	5.43	120.00	126.25	131.50	135.50	138.00

Imagine that we don't have the information above. Imagine we have been invited to a UK university to take skull measurements for 18 modern day Englishmen, and 16 ancient Celts. We have some information about modern day skulls to use as prior information for measurement mean and standard deviation. What is the power to observe a difference between the populations? Let's make some reasonable assumptions that allows us to be a bit conservative. Let's assume the sampled skulls from each of our populations is a random sample with common standard deviation 7mm, and let's assume we can't get the full sample but can only measure 15 skulls from each population. At a significance level of $\alpha = 0.05$, what is the power for detecting a difference of 5, 10, 15, 20, or 25 mm?

The theoretical two-sample power result is not too hard to derive (and is available in text books), but let's simply compare the power calculated exactly and by simulation.

For the exact result we use R library **pwr**. Below is the function call as well as the result. Note that we specified multiple effect sizes (diff/SD) in one call of the function.

```
# R code to compute exact two-sample two-sided power
library(pwr) # load the power calculation library

pwr.t.test(n = 15,
            d = c(5,10,15,20,25)/7,
            sig.level = 0.05,
            power = NULL,
            type = "two.sample",
            alternative = "two.sided")

##
##      Two-sample t test power calculation
##
##              n = 15
##              d = 0.7143, 1.4286, 2.1429, 2.8571, 3.5714
##      sig.level = 0.05
##              power = 0.4717, 0.9652, 0.9999, 1.0000, 1.0000
##      alternative = two.sided
##
##      NOTE: n is number in *each* group
```

To simulate the power under the same circumstances, we follow a similar strategy as in the one-sample example.

```
# R code to simulate two-sample two-sided power

# Strategy:
# Do this R times:
#   draw a sample of size N from the two hypothesized distributions
#   That is, 15 subjects from a normal distribution with specified means and sigma=7
#   Calculate the mean of the two samples
#   Calculate the associated z-statistic
#   See whether that z-statistic has a p-value < 0.05 under H0: mu_diff=0
#   If we reject H0, then set reject = 1, else reject = 0.
# Finally, the proportion of rejects we observe is the approximate power

n      <- 15;           # sample size of 25
mu1    <- 147;          # null hypothesis English mean
mu2    <- c(142, 137, 132, 127, 122); # Celt means
sigma   <- 7;           # standard deviation of normal population

alpha   <- 0.05;         # significance level

R       <- 2e4;          # Repetitions to draw sample and see whether we reject H0
#   The proportion of these that reject H0 is the power

power <- rep(NA,length(mu2)); # allocate a vector to store the calculated power in

for (j in 1:length(mu2)) { # do for each value of mu2

  reject <- rep(NA, R);     # allocate a vector of length R with missing values (NA)
#   to fill with 0 (fail to reject H0) or 1 (reject H0)

  for (i in 1:R) {
    sam1 <- rnorm(n, mean=mu1, sd=sigma); # English sample
    sam2 <- rnorm(n, mean=mu2[j], sd=sigma); # Celt sample

    ybar1 <- mean(sam1); # Calculate the mean of our sample sam
    ybar2 <- mean(sam2); # Calculate the mean of our sample sam

    # z-statistic (assumes we know sigma)
    # we could also have calculated the t-statistic, here
    z <- (ybar2 - ybar1) / (sigma * sqrt(1/n+1/n));

    pval.Left <- pnorm(z); # area under left tail
    pval.Right <- 1-pnorm(z); # area under right tail
    # p-value is twice the smaller tail area
    pval <- 2 * min(pval.Left, pval.Right);

    if (pval < 0.05) {
      reject[i] <- 1; # 1 for correctly rejecting H0
    } else {
      reject[i] <- 0; # 0 for incorrectly fail to reject H0
    }
  }
}

# calculate the average proportion of rejections across all repetitions
power <- mean(reject)
```

```

    }
}

# the average reject (proportion of rejects) is the power
power[j] <- mean(reject);
}

power
## [1] 0.4928 0.9765 1.0000 1.0000 1.0000

```

Note the similarity between power calculated using both the exact and simulation methods. If there is a power calculator for your specific problem, it is best to use that because it is faster and there is no programming. However, using the simulation method is better if we wanted to entertain different sample sizes with different standard deviations, etc. There may not be a standard calculator for our specific problem, so knowing how to simulate the power can be valuable.

Mean				Sample size		Power	
μ_E	μ_C	diff	SD	n_E	n_C	exact	simulated
147	142	5	7	15	15	0.4717	0.4928
147	137	10	7	15	15	0.9652	0.9765
147	132	15	7	15	15	0.9999	1
147	127	20	7	15	15	1.0000	1
147	122	25	7	15	15	1.0000	1

Appendix A

Custom R functions

A.1 Ch 2. Estimation in One-Sample Problems

```
# a function to compare the bootstrap sampling distribution with a
# normal distribution with mean and SEM estimated from the data
bs.one.samp.dist <- function(dat, N = 10000) {
  n <- length(dat)
  # resample from data
  sam <- matrix(sample(dat, size = N * n, replace = TRUE), ncol = N)
  # draw a histogram of the means
  sam.mean <- colMeans(sam)
  # save par() settings
  old.par <- par(no.readonly = TRUE)
  # make smaller margins
  par(mfrow = c(2, 1), mar = c(3, 2, 2, 1), oma = c(1, 1, 1, 1))
  # Histogram overlaid with kernel density curve
  hist(dat, freq = FALSE, breaks = 6, main = "Plot of data with smoothed density curve")
  points(density(dat), type = "l")
  rug(dat)

  hist(sam.mean, freq = FALSE, breaks = 25, main = "Bootstrap sampling distribution of the m
    xlab = paste("Data: n =", n, ", mean =", signif(mean(dat), digits = 5),
                ", se =", signif(sd(dat)/sqrt(n)), digits = 5))
  # overlay a density curve for the sample means
  points(density(sam.mean), type = "l")
  # overlay a normal distribution, bold and red
  x <- seq(min(sam.mean), max(sam.mean), length = 1000)
  points(x, dnorm(x, mean = mean(dat), sd = sd(dat)/sqrt(n)), type = "l",
         lwd = 2, col = "red")
  # place a rug of points under the plot
  rug(sam.mean)
  # restore par() settings
  par(old.par)
}
```

```
# Function to plot t-distribution with shaded p-value
t.dist.pval <- function(t.summary) {
  par(mfrow = c(1, 1))
  lim.extreme <- max(4, abs(t.summary$statistic) + 0.5)
  lim.lower <- -lim.extreme
  lim.upper <- lim.extreme
  x.curve <- seq(lim.lower, lim.upper, length = 200)
  y.curve <- dt(x.curve, df = t.summary$parameter)
  plot(x.curve, y.curve, type = "n", ylab = paste("t-dist( df =", signif(t.summary$parameter
    3), ")"), xlab = paste("t-stat =", signif(t.summary$statistic,
    5), ", Shaded area is p-value =", signif(t.summary$p.value, 5)))
  if ((t.summary$alternative == "less") | (t.summary$alternative == "two.sided")) {
    x.pval.l <- seq(lim.lower, -abs(t.summary$statistic), length = 200)
    y.pval.l <- dt(x.pval.l, df = t.summary$parameter)
    polygon(c(lim.lower, x.pval.l, -abs(t.summary$statistic)), c(0,
      y.pval.l, 0), col = "gray")
  }
  if ((t.summary$alternative == "greater") | (t.summary$alternative ==
    "two.sided")) {
    x.pval.u <- seq(abs(t.summary$statistic), lim.upper, length = 200)
    y.pval.u <- dt(x.pval.u, df = t.summary$parameter)
    polygon(c(abs(t.summary$statistic), x.pval.u, lim.upper), c(0,
      y.pval.u, 0), col = "gray")
  }
  points(x.curve, y.curve, type = "l", lwd = 2, col = "blue")
}
```

A.2 Ch 3. Two-Sample Inferences

```
# a function to compare the bootstrap sampling distribution of the
# difference of means from two samples with a normal distribution
# with mean and SEM estimated from the data
bs.two.samp.diff.dist <- function(dat1, dat2, N = 10000) {
  n1 <- length(dat1)
  n2 <- length(dat2)
  # resample from data
  sam1 <- matrix(sample(dat1, size = N * n1, replace = TRUE), ncol = N)
  sam2 <- matrix(sample(dat2, size = N * n2, replace = TRUE), ncol = N)
  # calculate the means and take difference between populations
  sam1.mean <- colMeans(sam1)
  sam2.mean <- colMeans(sam2)
  diff.mean <- sam1.mean - sam2.mean
  # save par() settings
  old.par <- par(no.readonly = TRUE)
  # make smaller margins
  par(mfrow = c(3, 1), mar = c(3, 2, 2, 1), oma = c(1, 1, 1, 1))
  # Histogram overlaid with kernel density curve
  hist(dat1, freq = FALSE, breaks = 6, main = paste("Sample 1", "\n",
    "n =", n1, ", mean =", signif(mean(dat1), digits = 5), ", sd =",
```

```
signif(sd(dat1), digits = 5)), xlim = range(c(dat1, dat2)))
points(density(dat1), type = "l")
rug(dat1)

hist(dat2, freq = FALSE, breaks = 6, main = paste("Sample 2", "\n",
  "n =", n2, ", mean =", signif(mean(dat2), digits = 5), ", sd =", 
  signif(sd(dat2), digits = 5)), xlim = range(c(dat1, dat2)))
points(density(dat2), type = "l")
rug(dat2)

hist(diff.mean, freq = FALSE, breaks = 25, main = paste("Bootstrap sampling distribution of the difference in means", "\n",
  "mean =", signif(mean(diff.mean), digits = 5), ", se =", 
  signif(sd(diff.mean), digits = 5)))
# overlay a density curve for the sample means
points(density(diff.mean), type = "l")
# overlay a normal distribution, bold and red
x <- seq(min(diff.mean), max(diff.mean), length = 1000)
points(x, dnorm(x, mean = mean(diff.mean), sd = sd(diff.mean)), type = "l",
  lwd = 2, col = "red")
# place a rug of points under the plot
rug(diff.mean)
# restore par() settings
par(old.par)
}
```