The Statistical Sleuth in R: Chapter 2

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1 Introduction

This document is intended to help describe how to undertake analyses introduced as examples in the Second Edition of the *Statistical Sleuth* (2002) by Fred Ramsey and Dan Schafer. More information about the book can be found at http://www.proaxis.com/~panorama/home.htm. This file as well as the associated knitr reproducible analysis source file can be found at http://www.amherst.edu/~nhorton/sleuth.

This work leverages initiatives undertaken by Project MOSAIC (http://www.mosaic-web. org), an NSF-funded effort to improve the teaching of statistics, calculus, science and computing in the undergraduate curriculum. In particular, we utilize the mosaic package, which was written to simplify the use of R for introductory statistics courses. A short summary of the R needed to teach introductory statistics can be found in the mosaic package vignette (http://cran.r-project. org/web/packages/mosaic/vignettes/MinimalR.pdf).

To use a package within R, it must be installed (one time), and loaded (each session). The package can be installed using the following command:

> install.packages('mosaic') # note the quotation marks

Once this is installed, it can be loaded by running the command:

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> require(mosaic)

This needs to be done once per session.

In addition the data files for the *Sleuth* case studies can be accessed by installing the *Sleuth2* package.

```
> install.packages('Sleuth2')  # note the quotation marks
```

```
> require(Sleuth2)
```

We also set some options to improve legibility of graphs and output.

```
> trellis.par.set(theme=col.mosaic()) # get a better color scheme for lattice
> options(digits=3, show.signif.stars=FALSE)
```

The specific goal of this document is to demonstrate how to calculate the quantities described in Chapter 2: Inference Using t-Distributions using R.

2 Bumpus's Data on Natural Selection

Is humerus length related to whether the bird would survive or perish? That's the question being addressed by Case Study 2.1 in the *Sleuth*.

2.1 Statistical summary and graphical display

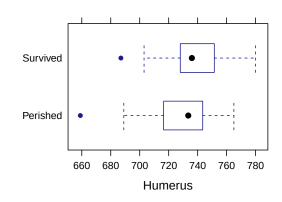
We begin by reading the data and summarizing the variables.

```
> summary(case0201)
```

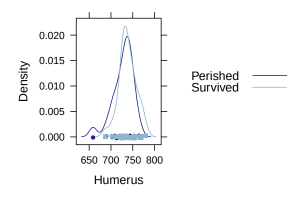
```
Humerus
                    Status
 Min.
        :659
               Perished:24
 1st Qu.:724
               Survived:35
Median :736
 Mean
        :734
3rd Qu.:747
Max.
       :780
> fav=favstats(Humerus ~ Status, data=case0201); fav
    Status min Q1 median Q3 max mean
                                         sd n missing
                      734 743 765 728 23.5 24
1 Perished 659 718
                                                     0
2 Survived 687 728 736 752 780 738 19.8 35
                                                     0
```

A total of 59 subjects are included in the data: 35 are adult male sparrows that survived and 24 that perished. The following figure replicates Display 2.1 on page 29.

> bwplot(Status ~ Humerus, data=case0201)



> densityplot(~ Humerus, groups=Status, auto.key=TRUE, data=case0201)



Both distributions are approximately normally distributed.

2.2 Inferential procedures (two-sample t-test)

First, we calculate the pooled SD and the standard error between these two different sample average (page 40, Display 2.8).

```
> # Calculate Pooled SD
> n1 = fav["Perished", "n"]; n1
[1] NA
> n2 = fav["Survived", "n"]; n2
[1] NA
> s1 = fav["Perished", "sd"]; s1
```

```
[1] NA
> s2 = fav["Survived", "sd"]; s2
[1] NA
> Sp = sqrt(((n1-1)*(s1)^2+(n2-1)*(s2)^2)/(n1+n2-2)); Sp
[1] NA
> # Calculate standard error
> SE = Sp*sqrt(1/n1+1/n2); SE
[1] NA
```

So the pooled SD is NA and the standard error is NA. Based on this information, we can construct a 95% confidence interval (page 41, Display 2.9).

```
> Y1 = fav["Perished", "mean"]; Y1
[1] NA
> Y2 = fav["Survived", "mean"]; Y2
[1] NA
> Yd = Y2-Y1; Yd
[1] NA
> df = n1+n2-2; df
[1] NA
> qt = qt(0.975, df); qt
[1] NA
> hw = qt*SE; hw
[1] NA
> lower = Yd-hw; lower
[1] NA
> upper = Yd+hw; upper
[1] NA
```

So the 95% confidence interval of the difference between means is (NA, NA)Now we want to calculate the *t*-statistic and *p*-value (as shown on page 44, Display 2.10).

```
> tstats = (Yd-0)/SE; tstats  # The hypothesis difference=0
[1] NA
> onepval = 1-pt(tstats, df); onepval
[1] NA
> twopval = 2*onepval; twopval
[1] NA
```

The one-sided *p*-value is NA and the two-sided *p*-value is NA.

We can get the results of "Summary of Statistical Findings" (page 29) by using the following code:

```
> t.test(Humerus ~ Status, var.equal=TRUE, data=case0201)
Two Sample t-test
data: Humerus by Status
t = -2, df = 57, p-value = 0.08
alternative hypothesis: true difference in means between group Perished and group Survived is
95 percent confidence interval:
-21.45
         1.28
sample estimates:
mean in group Perished mean in group Survived
                  728
                                          738
> confint(lm(Humerus ~ Status, data=case0201))
                2.5 % 97.5 %
              719.17 736.7
(Intercept)
StatusSurvived -1.28 21.4
```

3 Anatomical Abnormalities Associated with Schizophrenia

Is the area of brain related to the development of schizophrenia? That's the question being addressed by case study 2.2 in the *Sleuth*.

3.1 Statistical summary and graphical display

We begin by reading the data and summarizing the variables.

> summary(case0202)

Unaf	fect	Affected		
Min.	:1.25	Min.	:1.02	
1st Qu.	:1.60	1st Qu.	:1.31	
Median	:1.77	Median	:1.59	
Mean	:1.76	Mean	:1.56	
3rd Qu.	:1.94	3rd Qu.	:1.78	
Max.	:2.08	Max.	:2.02	

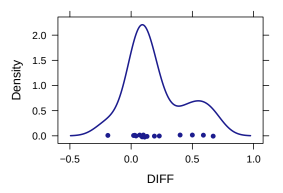
A total of 15 subjects are included in the data. There are 15 pairs of twins; one of the twins has schizophrenia, and the other does not. So there are 15 affected subjects and 15 unaffected subjects. The difference in area of left hippeermaps of these pairs of twins is:

The difference in area of left hippocampus of these pairs of twins is:

```
> DIFF = case0202[, "Unaffect"]-case0202[, "Affected"]
> favstats(DIFF)
min Q1 median Q3 max mean sd n missing
-0.19 0.055 0.11 0.315 0.67 0.199 0.238 15 0
```

This matches the results on page 30, Display 2.2.

> densityplot(DIFF)



3.2 Inferential procedures (two-sample t-test)

We want to calculate the paired t-test and 95% confidence interval.

```
> # Calculate t-statistics
> difmean = favstats(DIFF)[, "mean"]; difmean
[1] 0.199
```

```
> difsd = favstats(DIFF)[, "sd"]; difsd
[1] 0.238
> difSE = difsd/sqrt(15); difSE
[1] 0.0615
> tscore = (difmean-0)/difSE; tscore  # hypothesis difference=0
[1] 3.23
> twopvalue = 2*(1-pt(tscore, 15-1)); twopvalue
[1] 0.00606
> # Construct confidence interval
> q = qt(0.975, 15-1); q
[1] 2.14
> schizolower = difmean-q*difSE; schizolower
[1] 0.0667
> schizoupper = difmean+q*difSE; schizoupper
```

So the two-sided *p*-value is 0.006 and the 95% confidence interval is (0.07, 0.33). Or we can get the results displayed on page 31 by conducting a paired *t*-test.

[1] 0.331