The Statistical Sleuth in R: Chapter 11

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1 Introduction

This document is intended to help describe how to undertake analyses introduced as examples in the Third Edition of the *Statistical Sleuth* (2013) 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.math.smith.edu/~nhorton/sleuth3.

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:

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

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

```
> 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 **Sleuth3** package.

```
> install.packages("Sleuth3")  # note the quotation marks
```

```
> require(Sleuth3)
```

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 11: Model Checking and Refinement using R.

2 Alcohol metabolism in men and women

How do men and women metabolise alcohol? This is the question addressed in case study 11.1 in the *Sleuth*.

2.1 Data coding, summary statistics and graphical display

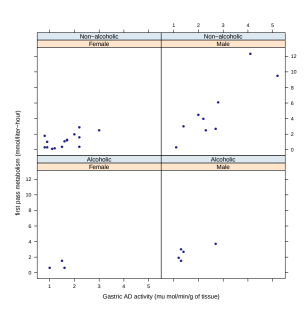
We begin by reading the data and summarizing the variables.

```
> summary(case1101)
   Subject
                 Metabol
                                Gastric
                                               Sex
Min. : 1.0
              Min. : 0.10
                                    :0.80
                                           Female:18
                             Min.
1st Qu.: 8.8
              1st Qu.: 0.60
                             1st Qu.:1.20
                                           Male :14
Median:16.5
              Median: 1.70 Median: 1.60
                   : 2.42
Mean
     :16.5
              Mean
                             Mean
                                  :1.86
              3rd Qu.: 2.92
3rd Qu.:24.2
                             3rd Qu.:2.20
              Max. :12.30
Max. :32.0
                             Max. :5.20
         Alcohol
Alcoholic : 8
Non-alcoholic:24
```

A total of 32 volunteers were included in this data. There were 18 females and 14 males, as recorded on Display 11.1 (page 311 of the *Sleuth*).

The following is a graphical display of the variables akin to Display 11.2 (page 312).

```
> xyplot(Metabol ~ Gastric | Sex + Alcohol, data = case1101, auto.key = TRUE,
+ xlab = "Gastric AD activity (mu mol/min/g of tissue)", ylab = "first pass metabolism (mm.)
```



2.2 Multiple regression

First we can fit a full model for estimating *metabolism* given a subjects *gastric AD activity*, whether they are *alcoholic* and *gender*. This first model is summarized on page 321 (Display 11.9).

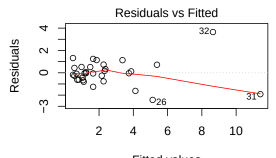
```
Coefficients:
                                    Estimate Std. Error t value Pr(>|t|)
(Intercept)
                                      -1.660
                                                  1.000
                                                         -1.66
                                                                    0.110
Gastric
                                       2.514
                                                  0.343
                                                           7.32 1.5e-07
SexFemale
                                       1.466
                                                  1.333
                                                           1.10
                                                                    0.282
                                                            1.31
AlcoholAlcoholic
                                       2.552
                                                  1.946
                                                                    0.202
Gastric:SexFemale
                                      -1.673
                                                  0.620
                                                          -2.70
                                                                    0.013
SexFemale: AlcoholAlcoholic
                                      -2.252
                                                  4.394
                                                          -0.51
                                                                    0.613
                                      -1.459
                                                          -1.39
Gastric: AlcoholAlcoholic
                                                  1.053
                                                                    0.179
Gastric:SexFemale:AlcoholAlcoholic
                                       1.199
                                                  2.998
                                                           0.40
                                                                    0.693
Residual standard error: 1.25 on 24 degrees of freedom
Multiple R-squared: 0.828, Adjusted R-squared: 0.777
F-statistic: 16.5 on 7 and 24 DF, p-value: 9.35e-08
```

Next we can calculate a number of model diagnostics, including leverage, studentized resids and Cook's distance (pages 325-327).

```
> require(MASS)
```

The following is a residual plot for the full model akin to Display 11.7 (page 319).

```
> plot(lm1, which = 1)
```



Fitted values
ol ~ Gastric + Sex + Alcohol + Gastric * Sex + Sex * Alc

From these diagnostics it appears that observations 31 and 32 may be influential points. Therefore, we next re-fit the full model excluding these two observations. The following results are found in Display 11.9 and discussed on page 321.

```
> case11012 = case1101[-c(31, 32), ]
> lm2 = lm(Metabol ~ Gastric + Sex + Alcohol + Gastric * Sex + Sex * Alcohol +
     Gastric * Alcohol + Gastric * Sex * Alcohol, data = case11012)
> summary(lm2)
Call:
lm(formula = Metabol ~ Gastric + Sex + Alcohol + Gastric * Sex +
   Sex * Alcohol + Gastric * Alcohol + Gastric * Sex * Alcohol,
    data = case11012)
Residuals:
   Min
            1Q Median
                            3Q
                                   Max
-1.8076 -0.5701 -0.0466 0.4976 1.4002
Coefficients:
                                  Estimate Std. Error t value Pr(>|t|)
(Intercept)
                                     -0.680
                                                1.309 -0.52
                                                                0.6088
                                                 0.608
                                                         3.16
Gastric
                                      1.921
                                                                0.0046
SexFemale
                                      0.486
                                                1.467
                                                         0.33
                                                                0.7436
AlcoholAlcoholic
                                     1.572
                                                1.812
                                                        0.87
                                                                0.3949
Gastric:SexFemale
                                    -1.081
                                                0.721
                                                       -1.50
                                                               0.1483
SexFemale: AlcoholAlcoholic
                                    -1.272
                                                3.467
                                                       -0.37
                                                                0.7172
Gastric:AlcoholAlcoholic
                                    -0.866
                                                 0.963
                                                       -0.90
                                                                0.3784
Gastric:SexFemale:AlcoholAlcoholic
                                     0.606
                                                 2.316
                                                         0.26
                                                                0.7961
Residual standard error: 0.941 on 22 degrees of freedom
Multiple R-squared: 0.685, Adjusted R-squared: 0.585
F-statistic: 6.83 on 7 and 22 DF, p-value: 0.000226
```

2.3 Refining the Model

This section addresses the process of refining the model. We first tested the lack of fit for the removal of Alcohol as shown in Display 11.13 (page 329).

```
> lm3 = lm(Metabol ~ Gastric + Sex + Gastric * Sex, data = case11012)
> summary(lm3)

Call:
lm(formula = Metabol ~ Gastric + Sex + Gastric * Sex, data = case11012)
```

```
Residuals:
   Min
            1Q Median
                           3Q
-1.5962 -0.6025 -0.0408 0.4759 1.6473
Coefficients:
                Estimate Std. Error t value Pr(>|t|)
                 0.0695
                           0.8019 0.09 0.9316
(Intercept)
                 1.5654
                           0.4074 3.84 0.0007
Gastric
                         0.9932 -0.27 0.7904
SexFemale
                 -0.2668
Gastric:SexFemale -0.7285
                           0.5394 -1.35 0.1885
Residual standard error: 0.882 on 26 degrees of freedom
Multiple R-squared: 0.673, Adjusted R-squared: 0.635
F-statistic: 17.8 on 3 and 26 DF, p-value: 1.71e-06
> anova(lm3, lm2) # page 322
Analysis of Variance Table
Model 1: Metabol ~ Gastric + Sex + Gastric * Sex
Model 2: Metabol ~ Gastric + Sex + Alcohol + Gastric * Sex + Sex * Alcohol +
   Gastric * Alcohol + Gastric * Sex * Alcohol
 Res.Df RSS Df Sum of Sq F Pr(>F)
     26 20.2
2 22 19.5 4 0.74 0.21 0.93
```

Next we assessed a model without an intercept which is scientifically plausible as summarized in Display 11.14 (page 329).

```
> lm4 = lm(Metabol ~ Gastric + Gastric:Sex - 1, data = case11012)
> summary(lm4)
Call:
lm(formula = Metabol ~ Gastric + Gastric:Sex - 1, data = case11012)
Residuals:
   Min
            1Q Median
                            30
-1.6171 -0.6075 -0.0262 0.4772 1.6230
Coefficients: (1 not defined because of singularities)
                 Estimate Std. Error t value Pr(>|t|)
Gastric
                    0.726
                               0.121
                                        5.99 1.9e-06
Gastric:SexMale
                    0.873
                               0.174
                                        5.02 2.6e-05
Gastric:SexFemale NA
                                  NA
                                          NA
```

```
Residual standard error: 0.852 on 28 degrees of freedom
Multiple R-squared: 0.877, Adjusted R-squared: 0.868
F-statistic: 99.9 on 2 and 28 DF, p-value: 1.8e-13

> anova(lm4, lm3)

Analysis of Variance Table

Model 1: Metabol ~ Gastric + Gastric:Sex - 1

Model 2: Metabol ~ Gastric + Sex + Gastric * Sex

Res.Df RSS Df Sum of Sq F Pr(>F)

1 28 20.3
2 26 20.2 2 0.094 0.06 0.94
```

Note that the "Summary of Statistical Findings" section (page 312) is based on this final model.

3 Blood brain barrier

Neuroscientists working to better understand the blood brain barrier have infused rats with cells to induce brain tumors. This is the topic addressed in case study 11.2 in the *Sleuth*.

3.1 Data coding and summary statistics

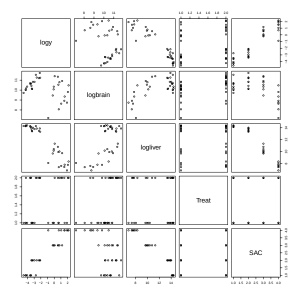
We begin by reading the data, performing transformations where needed and summarizing the variables.

```
> names(case1102)
                                       "Treatment" "Days"
[1] "Brain"
               "Liver"
                           "Time"
                                                               "Sex"
[7] "Weight"
               "Loss"
                           "Tumor"
> case1102 = transform(case1102, Y = Brain/Liver)
> case1102 = transform(case1102, logliver = log(Liver))
> case1102 = transform(case1102, logbrain = log(Brain))
> case1102 = transform(case1102, SAC = as.factor(Time))
> case1102 = transform(case1102, logy = log(Brain/Liver))
> case1102 = transform(case1102, logtime = log(Time))
> case1102 = transform(case1102, Treat = relevel(Treatment, ref = "NS"))
> summary(case1102)
                                        Time
                                                                Days
    Brain
                     Liver
                                                  Treatment
Min. : 1334 Min. :
                             928
                                   Min. : 0.5
                                                  BD:17
                                                           Min. : 9
                                                            1st Qu.:10
 1st Qu.: 19281 1st Qu.: 16210
                                   1st Qu.: 1.1
                                                  NS:17
Median: 32572 Median: 643965
                                   Median: 3.0
                                                            Median:10
Mean : 39965 Mean : 668776
                                   Mean :23.5
                                                            Mean :10
```

```
3rd Qu.: 50654
                  3rd Qu.:1318557
                                     3rd Qu.:24.0
                                                               3rd Qu.:10
       :123730
                         :1790863
                                            :72.0
                                                               Max.
Max.
                 Max.
                                     Max.
                                                                      :11
                                                                 Y
    Sex
                 Weight
                                Loss
                                                Tumor
                                   :-4.90
                                                    : 25
Female:26
            Min.
                    :184
                           Min.
                                            Min.
                                                           Min.
                                                                   :0.01
Male : 8
            1st Qu.:225
                           1st Qu.: 1.20
                                                           1st Qu.:0.03
                                            1st Qu.:136
            Median:240
                           Median: 3.95
                                            Median:166
                                                           Median:0.12
            Mean
                    :242
                           Mean
                                   : 3.64
                                            Mean
                                                    :183
                                                           Mean
                                                                   :1.50
            3rd Qu.:259
                           3rd Qu.: 5.97
                                            3rd Qu.:223
                                                           3rd Qu.:1.95
                                                           Max.
                    :298
                                   :12.80
                                            Max.
                                                    :484
            Max.
                           Max.
                                                                   :8.55
                                   SAC
   logliver
                    logbrain
                                                              logtime
                                               logy
       : 6.83
                        : 7.20
                                  0.5:9
                                                  :-4.58
                                                                   :-0.69
Min.
                Min.
                                          Min.
                                                           Min.
1st Qu.: 9.69
                 1st Qu.: 9.86
                                  3 :9
                                          1st Qu.:-3.39
                                                           1st Qu.:-0.25
Median :13.37
                Median :10.39
                                  24:8
                                          Median :-2.13
                                                           Median: 1.10
Mean
       :11.61
                Mean
                        :10.23
                                  72:8
                                          Mean
                                                  :-1.39
                                                           Mean
                                                                   : 1.86
3rd Qu.:14.09
                 3rd Qu.:10.83
                                          3rd Qu.: 0.67
                                                           3rd Qu.: 3.18
Max.
       :14.40
                Max.
                        :11.73
                                          Max.
                                                  : 2.15
                                                           Max.
                                                                   : 4.28
Treat
NS:17
BD:17
```

A total of 34 rats were included in this experiment. Each rat was given either the barrier solution (n = 17) or a normal saline solution (n = 17). Then variables of interest were calculated and are displayed in Display 11.4 (page 314 of the *Sleuth*).

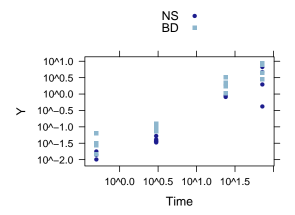
We can graphically relationships between the variables using a pairs plot.



3.2 Graphical presentation

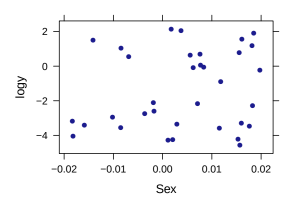
The following displays a scatterplot of log ratio (Y) as a function of log time, akin to Display 11.5 on page 315.

```
> xyplot(Y ~ Time, group = Treat, scales = list(y = list(log = TRUE), x = list(log = TRUE)),
+ auto.key = TRUE, data = case1102)
```

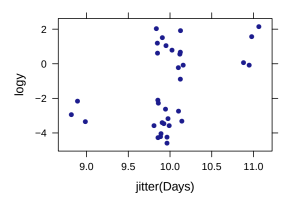


The following graphs are akin to the second and third plots in Display 11.16 on page 333.

```
> case1102 = transform(case1102, female = ifelse(Sex == "F", 1, 0))
> xyplot(logy ~ jitter(female), xlab = "Sex", data = case1102)
```



```
> xyplot(logy ~ jitter(Days), data = case1102)
```



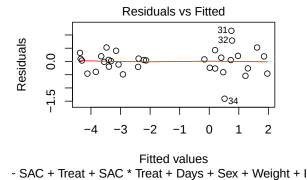
3.3 Multiple regression

We first fit a model that reflects the initial investigation. This is the proposed model from page 317.

```
Coefficients:
               Estimate Std. Error t value Pr(>|t|)
                                      -1.13
(Intercept)
              -3.836741
                           3.391046
                                                0.271
SAC3
               1.015463
                           0.399578
                                        2.54
                                                0.019
SAC24
               4.337135
                           0.477836
                                        9.08
                                              1.0e-08
SAC72
               5.010605
                           0.454953
                                      11.01
                                              3.5e-10
TreatBD
               0.795999
                           0.378970
                                        2.10
                                                0.048
Days
              -0.036987
                           0.295645
                                      -0.13
                                                0.902
                                        0.00
SexMale
               0.001295
                           0.373368
                                                0.997
Weight
              -0.000558
                           0.005330
                                       -0.10
                                                0.918
Loss
              -0.059544
                           0.030422
                                      -1.96
                                                0.064
Tumor
               0.001551
                           0.001226
                                       1.26
                                                0.220
SAC3:TreatBD
                           0.551964
                                        0.33
                                                0.748
               0.179831
SAC24:TreatBD -0.386047
                           0.585450
                                      -0.66
                                                0.517
SAC72:TreatBD
               0.379104
                           0.569242
                                        0.67
                                                0.513
Residual standard error: 0.564 on 21 degrees of freedom
Multiple R-squared: 0.96, Adjusted R-squared: 0.937
F-statistic: 41.9 on 12 and 21 DF, p-value: 6.45e-12
```

We can then display a residual plot to assess the fit of the above model. This is provided in Display 11.6 (page 318).

```
> plot(lm1, which = 1)
```



3.4 Refining the model

Lastly, we fit a refined model. These results can be found in Display 11.17 (page 334).

```
> lm2 = lm(logy ~ SAC + Treat, data = case1102)
> summary(lm2)
```

```
Call:
lm(formula = logy ~ SAC + Treat, data = case1102)
Residuals:
   Min
         1Q Median 3Q
-1.7402 -0.1755 -0.0178 0.2477 1.0551
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) -4.302 0.205 -21.01 < 2e-16
            1.134
                      0.252 4.50 0.00010
SAC3
SAC24
             4.257
                      0.259 16.43 3.1e-16
                      0.259 19.89 < 2e-16
SAC72
            5.154
TreatBD
             0.797
                      0.183 4.35 0.00016
Residual standard error: 0.533 on 29 degrees of freedom
Multiple R-squared: 0.951, Adjusted R-squared: 0.944
F-statistic: 140 on 4 and 29 DF, p-value: <2e-16
> anova(lm2, lm1)
Analysis of Variance Table
Model 1: logy ~ SAC + Treat
Model 2: logy ~ SAC + Treat + SAC * Treat + Days + Sex + Weight + Loss +
   Tumor
 Res.Df RSS Df Sum of Sq F Pr(>F)
    29 8.23
2 21 6.68 8 1.55 0.61 0.76
```