# The Statistical Sleuth in R: Chapter 13

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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:

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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 **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 = 4, show.signif.stars = FALSE)
```

The specific goal of this document is to demonstrate how to calculate the quantities described in Chapter 13: The Analysis of Variance for Two-Way Classifications using R.

# 2 Intertidal seaweed grazers

This wicked complicated trial is a subset of a factorial design (6 of the possible 2 by 2 by 2 combination of factors) plus blocking. This randomized block design is analyzed in case study 13.1 in the *Sleuth*.

## 2.1 Data coding, summary statistics and graphical display

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

```
> # logit transformation
> case1301 = transform(case1301, logitcover = log(Cover/(100 - Cover)))
```

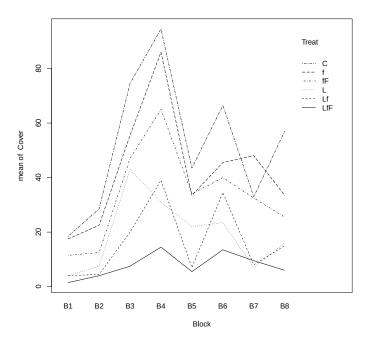
```
> summary(case1301)
     Cover
                    Block
                             Treat
                                         logitcover
 Min. : 1.0
                       :12
                             C
                                :16
                                       Min.
                                              :-4.595
                В1
 1st Qu.: 9.0
                                       1st Qu.:-2.314
                B2
                       :12
                             f :16
Median:22.5
                       :12
                                       Median :-1.237
                ВЗ
                             fF :16
 Mean :28.6
                В4
                       :12
                             L :16
                                       Mean
                                             :-1.233
 3rd Qu.:42.2
                             Lf :16
                                       3rd Qu.:-0.313
                B5
                       :12
 Max.
        :95.0
                В6
                       :12
                             LfF:16
                                              : 2.944
                                       Max.
                (Other):24
```

```
> favstats(logitcover ~ Treat, data = case1301)
       min
                Q1
                    median
                                  Q3
                                                          sd
                                                              n missing
                                         max
                                                 mean
C
    -1.815 - 0.7995
                    0.1201
                             0.80579
                                      2.9444
                                               0.1805 1.3990 16
f
    -2.091 -0.8119 -0.4898
                             0.09007
                                      2.0907 -0.3137 1.0748 16
                                                                       0
   -2.197 -1.7762 -0.5325 -0.30237
                                                                       0
fF
                                      0.9946 -0.8214 0.9599 16
    -3.178 -2.4784 -1.6964 -0.90838
                                      0.3228 -1.7120 1.0215 16
                                                                       0
L
    -3.476 -2.9444 -2.1530 -1.25519
                                      0.2819 -2.0044 1.1399 16
                                                                       0
LfF -4.595 -2.9444 -2.7515 -2.28453 -1.2657 -2.7247 0.8310 16
```

There were a total of 96 rock plots free of seaweed. These plots where split into 8 blocks based on location. Each block contained 12 plots. Then 6 treatments were randomly assigned to plots within each block. Therefore there were two plots per treatment within each block, as shown in Display 13.2 (page 377 of the *Sleuth*).

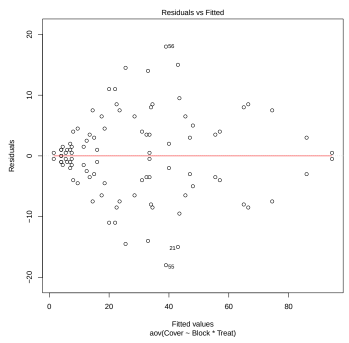
We can check for evidence of nonadditivity using interaction plots. For a figure akin to Display 13.7 on page 383 we can use the following code:

```
> with(case1301, interaction.plot(Block, Treat, Cover))
```



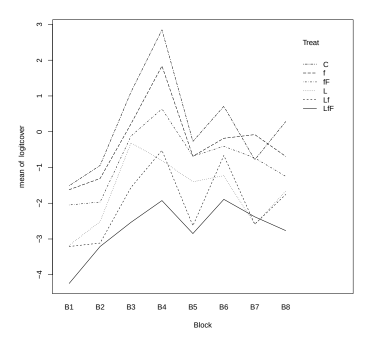
This figure shows evidence of nonadditivity. However as the authors note the type of nonadditivity seen in this figure may be removed by transformations. In addition, the residual plot from the saturated model (shown below and is akin to Display 13.8 on page 384) has a distinct funnel shape, also indicating a need for transformation.

```
> plot(aov(Cover ~ Block * Treat, data = case1301), which = 1)
```



After the log transformation, we can then observe an interaction plot on the log transformed data akin to Display 13.9 on page 385.

```
> with(case1301, interaction.plot(Block, Treat, logitcover))
```



## 2.2 Models

Then we can create an ANOVA for the nonadditive model estimating the log of the seaweed regeneration ratio as summarized on page 385 (Display 13.10).

```
> anova(lm(logitcover ~ Block * Treat, data = case1301))
Analysis of Variance Table
Response: logitcover
            Df Sum Sq Mean Sq F value Pr(>F)
Block
             7
                  76.2
                         10.89
                                  35.96 <2e-16
Treat
             5
                  97.0
                         19.40
                                  64.06 <2e-16
Block:Treat 35
                  15.2
                          0.44
                                   1.44
                                          0.12
            48
Residuals
                  14.5
                          0.30
```

This model has an  $R^2$  of 92.84%, an adjusted  $R^2$  of 85.83%, and an estimated SD of 0.5503. Notice that the interaction term has a large p-value, 0.1209, suggesting that the data may be more consistent with an additive model.

We can then compare these results to an ANOVA for the additive model estimating the log of the seaweed regeneration ratio as shown in Display 13.11 on page 387.

```
> anova(lm(logitcover ~ Block + Treat, data = case1301))
Analysis of Variance Table
```

```
Response: logitcover

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

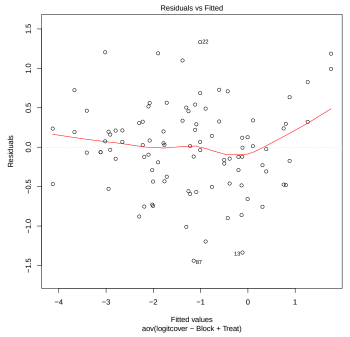
Block 7 76.2 10.89 30.4 <2e-16

Treat 5 97.0 19.40 54.1 <2e-16

Residuals 83 29.8 0.36
```

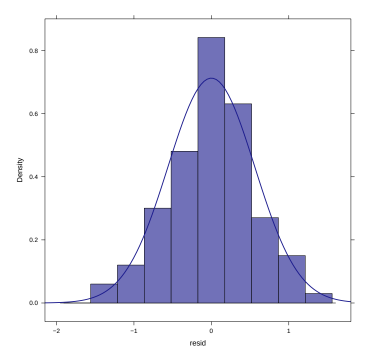
This model has an  $R^2$  of 85.34%, an adjusted  $R^2$  of 83.22%, and an estimated SD of 0.5989. Next we can assess the fit of the additive model through diagnostic plots. First we can check the linearity assumption.

```
> plot(aov(logitcover ~ Block + Treat, data = case1301), which = 1)
```

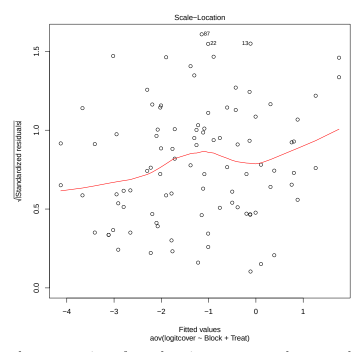


From this plot is appears that the linearity assumption seems reasonable. We will need to assume independence based on the information given. Next we will assess the normality assumption for the additive model.

```
> case1301$resid = residuals(aov(logitcover ~ Block + Treat, data = case1301))
> histogram(~resid, type = "density", density = TRUE, data = case1301)
```



From this figure normality seems reasonable as well. Now we can assess equality of variance.

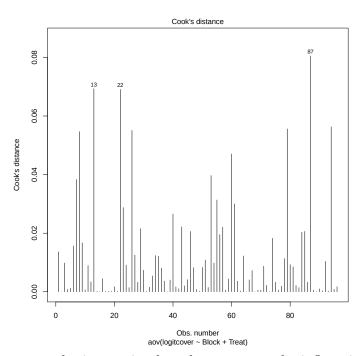


From this figure, the assumption of equal variance seems to be somewhat problematic, as seen

in the curvature of the lowess line.

Lastly we can look for influential points and/or high leverage with the additive model.

```
> plot(aov(logitcover ~ Block + Treat, data = case1301), which = 4)
```



From this figure we can obtain certain plots that appear to be influential points.

```
> case1301[c(13, 22, 87), ]
   Cover Block Treat logitcover
                                 resid
                         -1.4500 -1.336
13
      19
            B7
                    C
22
      58
            ВЗ
                    L
                          0.3228 1.333
87
            B4
                  LfF
                         -2.5867 -1.440
```

## 2.3 Linear combinations

First we can observe the Block and Treatment averages and the Block and Treatment effects from Display 13.12 (page 388).

For the effects we used:

```
> model.tables(aov(lm(logitcover ~ Block * Treat, data = case1301)), type = "effects")
Tables of effects
Block
```

```
Block
 B1
         B2 B3 B4 B5
                                     В6
                                          В7
                                                   R8
-1.4031 -0.9432 0.7015 1.5776 -0.1871 0.6220 -0.2946 -0.0731
Treat
Treat
        f fF L Lf LfF
1.4131 0.9190 0.4112 -0.4794 -0.7718 -1.4921
Block:Treat
   Treat
       f fF L Lf LfF
Block C
  B1 -0.2892 0.0951 0.1755 -0.0629 0.1972 -0.1157
  B2 -0.1797 -0.0509 -0.2013 0.1406 -0.1663 0.4576
  B3 0.2303 -0.1658 -0.0007 0.6996 -0.2540 -0.5094
  B4 1.0899 0.5743 -0.1179 -0.6724 -0.0947 -0.7791
  B5 -0.2650 -0.1850 0.3241 0.4996 -0.4376 0.0638
  B6 -0.0918 -0.4920 -0.2067 -0.1392 0.7185 0.2112
  B7 -0.6709 0.5274 0.3807 -0.5903 -0.2862 0.6394
 B8 0.1763 -0.3030 -0.3536 0.1250 0.3231 0.0322
```

For the means we changed the type attribute to "means":

```
> model.tables(aov(lm(logitcover ~ Block * Treat, data = case1301)), type = "means")
Tables of means
Grand mean
-1.233
Block
Block
         B2 B3 B4 B5 B6 B7 B8
-2.6357 -2.1758 -0.5311 0.3450 -1.4197 -0.6106 -1.5272 -1.3057
Treat
Treat
          f fF L Lf LfF
0.1805 -0.3137 -0.8214 -1.7120 -2.0044 -2.7247
Block:Treat
   Treat
Block C f fF L Lf LfF
  B1 -1.512 -1.622 -2.049 -3.178 -3.210 -4.243
  B2 -0.942 -1.308 -1.966 -2.515 -3.114 -3.210
B3 1.112 0.222 -0.121 -0.311 -1.557 -2.533
```

```
B4 2.848 1.838 0.638 -0.807 -0.522 -1.926

B5 -0.272 -0.686 -0.684 -1.399 -2.629 -2.848

B6 0.711 -0.184 -0.406 -1.229 -0.664 -1.891

B7 -0.785 -0.081 -0.735 -2.597 -2.585 -2.380

B8 0.284 -0.690 -1.248 -1.660 -1.754 -2.766
```

To answer specific questions of interest regarding subgroup comparisons we can use linear combinations. The *Sleuth* proposes five questions as detailed on pages 289-390. The code for results of these questions is displayed below and these results are also interpreted on pages 389-390 and summarized in Display 13.13. For this model the reference group is *control* followed by f, fF, L, Lf, LfF.

```
> require(gmodels)
Loading required package: gmodels
> lm1 = lm(logitcover ~ Treat + Block, data = case1301)
> coef(lm1)
                          TreatfF
                                     TreatL
(Intercept)
               Treatf
                                                 TreatLf
                                                            TreatLfF
   -1.2226
               -0.4941
                          -1.0019
                                      -1.8925
                                                 -2.1849
                                                            -2.9052
   BlockB2
              BlockB3
                          BlockB4
                                     BlockB5
                                                 BlockB6
                                                             BlockB7
              2.1046
                          2.9807
                                  1.2160
    0.4600
                                                2.0251
                                                             1.1085
   BlockB8
    1.3300
> large = rbind(`Large fish` = c(0, -1/2, 1/2, 0, -1/2, 1/2))
> small = rbind(`Small fish` = c(-1/2, 1/2, 0, -1/2, 1/2, 0))
> limpets = rbind(Limpets = c(-1/3, -1/3, -1/3, 1/3, 1/3, 1/3))
> limpetsSmall = rbind(`Limpets X Small` = c(1, -1/2, -1/2, -1, 1/2, 1/2))
> limpetsLarge = rbind(`Limpets X Large` = c(0, 1, -1, 0, -1, 1))
> fit.contrast(lm1, "Treat", large, conf.int = 0.95)
               Estimate Std. Error t value Pr(>|t|) lower CI upper CI
                          0.1497 -4.101 9.54e-05 -0.9118 -0.3162
TreatLarge fish -0.614
> fit.contrast(lm1, "Treat", small, conf.int = 0.95)
               Estimate Std. Error t value Pr(>|t|) lower CI upper CI
                        0.1497 -2.627 0.01026 -0.691 -0.09549
TreatSmall fish -0.3933
> fit.contrast(lm1, "Treat", limpets, conf.int = 0.95)
            Estimate Std. Error t value Pr(>|t|) lower CI upper CI
TreatLimpets
            -1.829
                       0.1222 -14.96 2.778e-25 -2.072 -1.586
> fit.contrast(lm1, "Treat", limpetsSmall, conf.int = 0.95)
```

```
Estimate Std. Error t value Pr(>|t|) lower CI
TreatLimpets X Small
                      0.09549
                                  0.2593 0.3682
                                                   0.7136 - 0.4203
                     upper CI
TreatLimpets X Small
                       0.6113
> fit.contrast(lm1, "Treat", limpetsLarge, conf.int = 0.95)
                     Estimate Std. Error t value Pr(>|t|) lower CI
TreatLimpets X Large
                     -0.2125
                                  0.2994 - 0.7097
                                                   0.4799
                     upper CI
                        0.383
TreatLimpets X Large
```

To attain the confidence intervals discussed in the "Summary of Statistical Findings" (page 376) we need to exponential the lower and upper bounds of the above 95% confidence intervals. Therefore, for the limpets estimation, the corresponding 95% confidence interval is (0.126, 0.205). The resulting large fish 95% confidence interval is (0.402, 0.729). Lastly for the estimation of the regeneration ratio for small fish the 95% confidence interval is (0.501, 0.909).

## 3 Pygmalion effect

Does expected excellence affect performance? More specifically, does telling a manager that some of the supervisees are "superior" affect the supervisor's perception of their performance (Pygmalion effect)? This is the question addressed in case study 13.2 in the *Sleuth*.

#### 3.1 Statistical summary

We begin by reading the data and summarizing the variables.

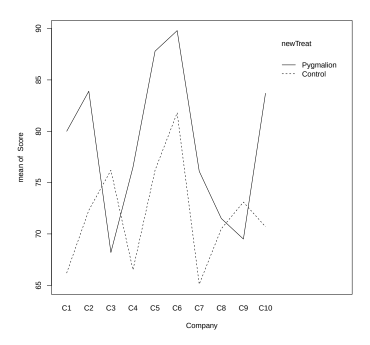
```
> summary(case1302)
                     Treat
                                  Score
    Company
              Pygmalion:10
 C1
        : 3
                                      :59.5
                              Min.
        : 3
 C2
              Control :19
                              1st Qu.:69.2
 C4
        : 3
                              Median:73.9
 C5
        : 3
                              Mean
                                    :74.1
 C6
                              3rd Qu.:78.9
        : 3
 C7
        : 3
                              Max.
                                      :89.8
 (Other):11
> case1302$newTreat = relevel(case1302$Treat, ref = "Control")
```

There were a total of 29 platoons. For each of the 10 companies, one platoon received the Pygmalion treatment and two platoons were control, with the exception of one company that only had one control platoon. Therefore, there were 10 Pygmalion platoons and 19 control platoons. As shown in Display 13.3 (page 378 of the *Sleuth*).

## 3.2 Graphical presentation

The following figure displays an interaction plot for the Pygmalion dataset, akin to Display 13.14 on page 392.

```
> with(case1302, interaction.plot(Company, newTreat, Score))
```



## 3.3 Two way ANOVA (fit using multiple linear regression model)

We can then use multiple linear regression models for the additive and nonadditive models and compare them using the two-way ANOVA.

The following is similar to Display 13.16 (page 394).

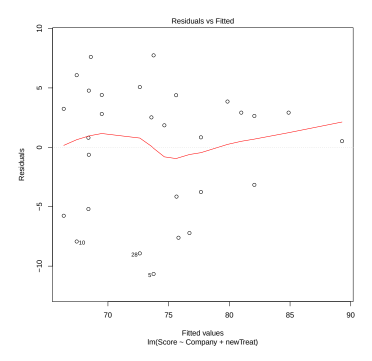
```
> lm1 = lm(Score ~ Company * newTreat, data = case1302)
> summary(lm1)
Call:
lm(formula = Score ~ Company * newTreat, data = case1302)
Residuals:
  Min
         1Q Median
                         3Q
                               Max
 -9.2
        -2.3
              0.0
                        2.3
                               9.2
Coefficients:
                             Estimate Std. Error t value Pr(>|t|)
```

(Intercept)	66.20	5.09	13.00	3.9e-07	
CompanyC2	6.10	7.20	0.85	0.419	
CompanyC3	10.00	8.82	1.13	0.286	
CompanyC4	0.30	7.20	0.04	0.968	
CompanyC5	10.00	7.20	1.39	0.198	
CompanyC6	15.60	7.20	2.17	0.059	
CompanyC7	-1.10	7.20	-0.15	0.882	
CompanyC8	4.30	7.20	0.60	0.565	
CompanyC9	6.90	7.20	0.96	0.363	
CompanyC10	4.50	7.20	0.62	0.548	
newTreatPygmalion	13.80	8.82	1.56	0.152	
CompanyC2:newTreatPygmalion	-2.20	12.48	-0.18	0.864	
CompanyC3:newTreatPygmalion	-21.80	13.48	-1.62	0.140	
CompanyC4:newTreatPygmalion	-3.80	12.48	-0.30	0.768	
CompanyC5:newTreatPygmalion	-2.20	12.48	-0.18	0.864	
CompanyC6:newTreatPygmalion	-5.80	12.48	-0.46	0.653	
CompanyC7:newTreatPygmalion	-2.80	12.48	-0.22	0.827	
CompanyC8:newTreatPygmalion	-12.80	12.48	-1.03	0.332	
CompanyC9:newTreatPygmalion	-17.40	12.48	-1.39	0.197	
CompanyC10:newTreatPygmalion	-0.80	12.48	-0.06	0.950	
<pre>&gt; lm2 = lm(Score ~ Company + &gt; summary(lm2) # Display 13.</pre>			e1302)		
Call: lm(formula = Score ~ Company	+ newTreat,	data = ca	nse1302)		
Residuals: Min 1Q Median 3Q -10.66 -4.15 1.85 3.85	Max 7.74				
Coefficients:		7 D ()	1.15		
	3.8931		> t ) 9e-13		
± '	5.3697		9e-13 0.331		
1 0					
CompanyC3 0.1966	6.0189		).974		
CompanyC4 -0.9667			0.859		
CompanyC5 9.2667	5.3697		0.102		
CompanyC6 13.6667 CompanyC7 -2.0333	5.3697 5.3697		0.020 0.709		
CompanyC7 -2.0333					

```
0.0333
                           5.3697 0.01 0.995
CompanyC8
CompanyC9
                  1.1000
                           5.3697
                                    0.20
                                           0.840
                  4.2333
                           5.3697
                                    0.79
                                           0.441
CompanyC10
                7.2205
                                    2.80 0.012
newTreatPygmalion
                           2.5795
Residual standard error: 6.58 on 18 degrees of freedom
Multiple R-squared: 0.565, Adjusted R-squared: 0.323
F-statistic: 2.33 on 10 and 18 DF, p-value: 0.0564
> anova(lm1)
Analysis of Variance Table
Response: Score
               Df Sum Sq Mean Sq F value Pr(>F)
Company
               9
                    671
                          75
                                1.44 0.299
newTreat
              1
                    339
                           339 6.53 0.031
Company:newTreat 9 311
                          35 0.67 0.722
Residuals 9 467 52
> anova(lm2)
Analysis of Variance Table
Response: Score
        Df Sum Sq Mean Sq F value Pr(>F)
              671 75 1.72 0.156
Company 9
              339
                     339
                           7.84 0.012
newTreat 1
Residuals 18
              779
                     43
> anova(lm2, lm1)
Analysis of Variance Table
Model 1: Score ~ Company + newTreat
Model 2: Score ~ Company * newTreat
 Res.Df RSS Df Sum of Sq F Pr(>F)
    18 779
1
2 9 467 9
               312 0.67 0.72
```

Lastly we can observe the residual plot from the fit of the additive model, akin to Display 13.17 on page 395.

```
> plot(lm2, which = 1)
```



#### 3.4 Randomization Methods

As introduced in Chapter 4, we can construct a randomization distribution by considering the distribution of a test statistic over all possible ways the randomization could have turned out. For the Pygmalion data we can construct a randomization distribution for the t-statistic of the treatment effect as discussed on pages 397-398.

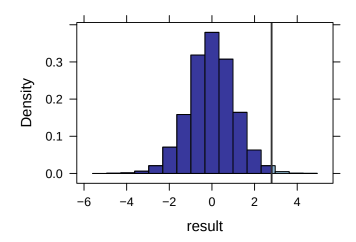
```
> mod = lm(Score ~ Company+newTreat, data=case1302)
> obs = summary(mod)$coefficients["newTreatPygmalion", "t value"]
> obs

[1] 2.799

> nulldist = do(10000) * summary(lm(Score ~ shuffle(Company)+shuffle(newTreat),
+ data=case1302))$coefficients["shuffle(newTreat)Pygmalion", "t value"]
> histogram(~ result, groups=result >= obs, v=obs, data=nulldist)
> # akin to Display 13.20 page 398
> tally(~ result >= obs, format="proportion", data=nulldist)

TRUE FALSE Total
0.0056 0.9944 1.0000
```

3 PYGMALION EFFECT



From this simulation we observed that the proportion of t-statistics that were as extreme or more extreme than our observed t-statistic (2.799) is 0.0056.