Ch-03 R Codes

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Textbook: Montgomery, D. C. (2012). Design and analysis of experiments, 8th Edition. John Wiley & Sons. Online handouts: https://github.com/PingYangChen/ANOVA_Course_R_Code

Chapter 3 One-way ANOVA: Plasma Etching Experiment

Create an csv file to store the data and then read it in R.

А	В	С
i	Power	EtchRate
1	160	575
2	160	542
3	160	530
4	160	539
5	160	570
6	180	565
7	180	593
8	180	590
9	180	579
10	180	610
11	200	600
12	200	651
13	200	610
14	200	637
15	200	629
16	220	725
17	220	700
18	220	715
19	220	685
20	220	710

Figure 1: Plasma Etching Experiment Data

Read the csv file 3_PlasmaEtching.csv in R. Make sure that in the data.frame the variable Power is a factor. If not sure, apply as.factor() function to set the property of the variable Power after reading the dataset.

```
df1 <- read.csv(file.path("data", "3_PlasmaEtching.csv"))
df1$Power <- as.factor(df1$Power)</pre>
```

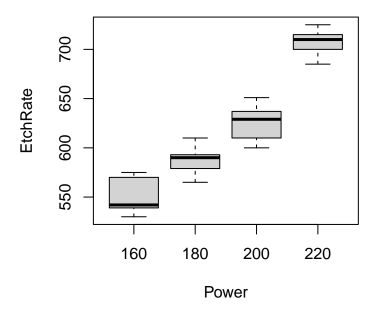
To compute descriptive statistics of the data in each subgroup of a dataset in R, we use tapply().

```
tapply(df1$EtchRate, df1$Power, summary)
```

```
## $'160'
##
      Min. 1st Qu.
                      Median
                                 Mean 3rd Qu.
                                                   Max.
##
     530.0
              539.0
                       542.0
                                551.2
                                         570.0
                                                  575.0
##
##
   $'180'
##
      Min. 1st Qu.
                      Median
                                 Mean 3rd Qu.
                                                   Max.
     565.0
                                         593.0
##
              579.0
                       590.0
                                587.4
                                                  610.0
##
## $'200'
##
      Min. 1st Qu.
                      Median
                                 Mean 3rd Qu.
                                                   Max.
##
     600.0
              610.0
                       629.0
                                625.4
                                         637.0
                                                  651.0
##
## $'220'
##
      Min. 1st Qu.
                      Median
                                 Mean 3rd Qu.
                                                   Max.
                         710
                                  707
##
       685
                700
                                           715
                                                    725
```

Alternatively, boxplots provide a quick and direct means of observing the differences among the responses of the four treatments (groups or levels of a factor).

```
# Draw the grouped boxplot
boxplot(EtchRate ~ Power, data = df1)
```



The boxplot shows that: as Power increases, the median EtchRate increases steadily. This finding would imply that higher power levels produce higher etching rates.

To analysis this plasma etching experiment data, the ANOVA model is

$$y_{ij} = \mu + \tau_i + \varepsilon_{ij}, \begin{cases} i = 1, 2, 3, 4 \\ j = 1, 2, 3, 4, 5 \end{cases}$$

where μ is an overall mean, τ_i is the *i*th treatment (Power) effect and ε_{ij} is the random experiment error that is assumed following $N(0, \sigma^2)$ identically and independently.

Let the treatment means to be $\mu_i = \mu + \tau_i$, we test the statistical hypotheses:

 $H_0: \quad \mu_1 = \mu_2 = \mu_3 = \mu_4$ $H_1: \quad \text{at least one mean is different}$

In R, the function aov() fits the ANOVA model. For one-way ANOVA, the command is as follows. Then, we call summary() to examine the ANOVA table.

```
fit <- aov(EtchRate ~ Power, data = df1)
summary(fit)</pre>
```

```
## Df Sum Sq Mean Sq F value Pr(>F)
## Power    3 66871 22290 66.8 2.88e-09 ***
## Residuals 16 5339 334
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

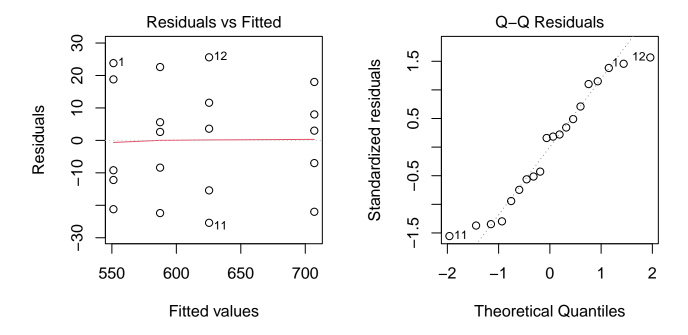
Model Adequacy Checking

The adequacy of an ANOVA model can be studied from residual plots. The basic approach is to use the plot() function with the fitted ANOVA model object as its input argument. There are four residual plots and only only need the first two. The following R commands are used to view both residual plots simultaneously.

The left plot is the residual plot against the fitted values. This plot is used to check the consistency of the variance with changes in the fitted value. A lack of any visually obvious pattern in the dots on the plot is desired.

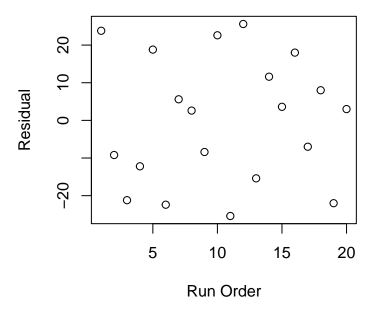
The right plot is the residuals' Normal Quantile-Quantile (QQ) plot. The QQ plot provides information about whether the residuals (or data) follow a normal distribution by comparing their quantiles to those of a theoretical normal distribution. Ideally, the dots form a straight line.

```
par(mfrow = c(1, 2))
plot(fit, which = 1:2)
```



To check the independence of the residual, we draw the scatter plot of residuals against the experiment order. A lack of any visually obvious pattern in the dots on the plot is desired.

```
resid <- fit$residuals
plot(1:length(resid), resid, xlab = "Run Order", ylab = "Residual")</pre>
```



There are also statistical tests for model adequacy checking.

For checking normality, one commonly used statistical test is the Shapiro–Wilk test. A p-value larger than the significance level indicates that there is no evidence against the assumption that the residuals are normally distributed.

shapiro.test(fit\$residuals)

```
##
## Shapiro-Wilk normality test
##
## data: fit$residuals
## W = 0.93752, p-value = 0.2152
```

For checking the equality of variance among treatment groups, a recommended statistical test is the modified Levene's test. A p-value larger than the significance level indicates that there is no evidence against the assumption that the residuals are homoscedastic (i.e., have equal variances across groups).

```
# Install the asbio package
# install.packages("asbio")
# Load the asbio package, which includes the modified Levene's test
# for homogeneity of variance
library(asbio)
# Perform Levene's test for homogeneity of variance
# It checks if the variances across the different 'Power' groups are equal
modlevene.test(fit$residuals, df1$Power)
```

```
##
## Modified Levene's test of homogeneity of variances
##
## df1 = 3, df2 = 16, F = 0.19587, p-value = 0.89767
```

Post-ANOVA Comparison of Means

The estimate of the overall mean μ and the Power's treatment effects τ_1 to τ_4 are

$$\hat{\mu} = \frac{1}{an} \sum_{i=1}^{a} \sum_{j=1}^{n} y_{ij} = \bar{y}_{..};$$

$$\hat{\tau}_{1} = \frac{1}{n} \sum_{j=1}^{n} y_{1j} - \hat{\mu} = \bar{y}_{1.} - \bar{y}_{..}; \hat{\tau}_{2} = \frac{1}{n} \sum_{j=1}^{n} y_{2j} - \hat{\mu} = \bar{y}_{2.} - \bar{y}_{..};$$

$$\hat{\tau}_{3} = \frac{1}{n} \sum_{j=1}^{n} y_{3j} - \hat{\mu} = \bar{y}_{3.} - \bar{y}_{..}; \hat{\tau}_{4} = \frac{1}{n} \sum_{j=1}^{n} y_{4j} - \hat{\mu} = \bar{y}_{4.} - \bar{y}_{..}.$$

The R codes are as follows.

```
mean(df1$EtchRate) # Overall
mean(df1$EtchRate[df1$Power == 160]) - mean(df1$EtchRate) # tau_1
mean(df1$EtchRate[df1$Power == 180]) - mean(df1$EtchRate) # tau_2
mean(df1$EtchRate[df1$Power == 200]) - mean(df1$EtchRate) # tau_3
mean(df1$EtchRate[df1$Power == 220]) - mean(df1$EtchRate) # tau_4
```

Following an ANOVA in which we have rejected the null hypothesis of equal treatment means, we wish to test all pairwise mean comparisons:

$$H_0: \ \mu_i = \mu_j$$

$$H_1: \ \mu_i \neq \mu_j$$

for all $i \neq j$. Here, we introduce three approaches.

Pairwise t-tests

The straightforward approach to test for all pairs of the hypotheses is to conduct the Pairwise t-tests simultaneously. The following codes give the results under Bonferroni adjustment on the p-value.

```
pairwise.t.test(df1$EtchRate, df1$Power, p.adjust = "bonferroni")
```

```
##
## Pairwise comparisons using t tests with pooled SD
##
## data: df1$EtchRate and df1$Power
##
## 160 180 200
## 180 0.038 - -
## 200 5.1e-05 0.028 -
## 220 2.2e-09 1.0e-07 1.6e-05
##
## P value adjustment method: bonferroni
```

Tukey's Test

Tukey's procedure makes use of the distribution of the studentized range statistic

$$q = \frac{\bar{y}_{max} - \bar{y}_{min}}{\sqrt{MS_E/n}}$$

where \bar{y}_{max} and \bar{y}_{min} are the largest and smallest sample means respectively, out of a group of p sample means. For equal sample sizes, Tukey's test declares two means significantly different if the absolute value of their sample differences exceeds

$$T_{\alpha} = q_{\alpha}(a, f) \sqrt{\frac{MS_E}{n}}$$

where $q_{\alpha}(a, f)$ is the upper α percentage points of q and f is the number of degrees of freedom associated with the MS_E . For more insights on the distribution of q, please refer to the textbook. Tukey's method is performed by the function TukeyHSD().

TukeyHSD(fit)

```
##
     Tukey multiple comparisons of means
       95% family-wise confidence level
##
##
## Fit: aov(formula = EtchRate ~ Power, data = df1)
##
## $Power
                                           p adj
##
            diff
                        lwr
                                  upr
## 180-160
            36.2
                   3.145624
                             69.25438 0.0294279
## 200-160
            74.2
                  41.145624 107.25438 0.0000455
## 220-160 155.8 122.745624 188.85438 0.0000000
## 200-180
            38.0
                   4.945624
                             71.05438 0.0215995
## 220-180 119.6
                  86.545624 152.65438 0.0000001
## 220-200 81.6
                  48.545624 114.65438 0.0000146
```

Fisher's LSD Method

The R package agricolae provides the function LSD.test() to perform Fisher's LSD test. Adjustment for the P-value is necessary. Typically, we set p.adj = "bonferroni" for the Bonferroni method.

```
if (!("agricolae" %in% rownames(installed.packages()))) {
  install.packages("agricolae")
}
library(agricolae)
out <- LSD.test(fit, "Power", p.adj = "bonferroni")</pre>
print(out)
## $statistics
##
     MSerror Df
                  Mean
                              CV t.value
                                                MSD
##
       333.7 16 617.75 2.957095 3.008334 34.75635
##
##
   $parameters
##
           test
                 p.ajusted name.t ntr alpha
     Fisher-LSD bonferroni Power
                                        0.05
##
##
## $means
##
       EtchRate
                     std r
                                          LCL
                                                    UCL Min Max Q25 Q50 Q75
                                  se
## 160
          551.2 20.01749 5 8.169455 533.8815 568.5185 530 575 539 542 570
          587.4 16.74216 5 8.169455 570.0815 604.7185 565 610 579 590 593
## 180
          625.4 20.52559 5 8.169455 608.0815 642.7185 600 651 610 629 637
## 200
          707.0 15.24795 5 8.169455 689.6815 724.3185 685 725 700 710 715
## 220
##
## $comparison
## NULL
##
## $groups
##
       EtchRate groups
## 220
          707.0
## 200
          625.4
                     b
## 180
          587.4
                     С
## 160
          551.2
                      d
##
## attr(,"class")
## [1] "group"
```

The most important parts of the outputs are shown below:

- \$means displays the estimated mean of the etching rate at each level of power.
- \$groups indicates the significance of the difference in the etching rate at each level of power. The column groups in \$groups encodes the treatment levels with no significant difference in the etching rate by the same alphabet letter.

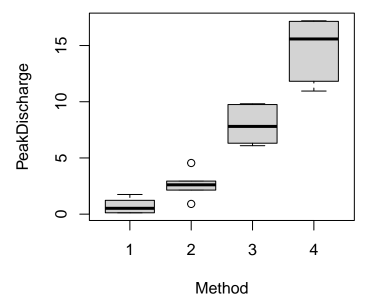
Variance-Stabilizing Transformations: Peak Discharge Experiments

Read the csv file 3_PeakDischarge.csv in R. Make sure that in the data.frame the variable Method is a factor because it represents categorical data (i.e., different methods). If not sure, apply as.factor() function to set the property of the variable Method after reading the dataset.

```
# Load the dataset from the 'data' directory
df2 <- read.csv(file.path("data", "3_PeakDischarge.csv"))
# Convert 'Method' column to a factor variable since it's categorical
df2$Method <- as.factor(df2$Method)</pre>
```

Boxplot provides an initial insight into whether the variance across groups looks consistent and whether there might be any outliers.

```
# Boxplot to visualize the distribution of 'PeakDischarge' for each 'Method'
boxplot(PeakDischarge ~ Method, data = df2)
```



In the boxplot of the PeakDischarge data, we observe that the data variation for Methods 3 and 4 appears greater than that for Methods 1 and 2. To confirm whether the variances across the different 'Method' groups are statistically equal, we can perform modified Levene's test for homogeneity of variance. R users can implement the modified Levene's test by installing the package **asbio**. R codes are shown below.

```
# Install the asbio package
# install.packages("asbio")
# Load the asbio package, which includes the modified Levene's test
# for homogeneity of variance
library(asbio)
# Perform Levene's test for homogeneity of variance
# It checks if the variances across the different 'Method' groups are equal
modlevene.test(df2$PeakDischarge, df2$Method)
```

```
##
## Modified Levene's test of homogeneity of variances
##
## df1 = 3, df2 = 20, F = 4.56844, p-value = 0.01357
```

The p-value of the modified Levene's test is 0.0136, which is lower than the pre-specified significance level of 0.05, indicating that the variances across the different 'Method' groups are significantly different.

Now, suppose we did not perform the modified Levene's test and directly fit an one-way ANOVA instead.

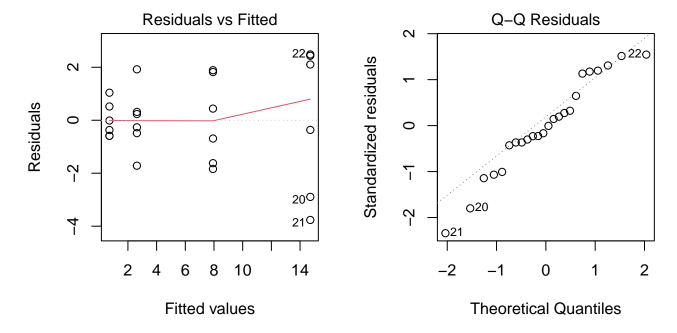
```
# Perform a one-way ANOVA to test if there is a significant difference
# in 'PeakDischarge' across methods
fit2 <- aov(PeakDischarge ~ Method, data = df2)
# Summary of the ANOVA results, including F-statistic and p-value
summary(fit2)</pre>
```

```
## Df Sum Sq Mean Sq F value Pr(>F)
## Method    3 708.3 236.1 76.07 4.11e-11 ***
## Residuals    20 62.1    3.1
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

The p-value of the F-test in the ANOVA table is extremely small, providing significant evidence that the mean Peak Discharge differs across the Methods. But is this conclusion valid?

If we examine the residual plots, particularly the scatter of fitted values versus residuals, we notice an increasing trend in the variation of residuals across the Method groups. This suggests that the assumption of constant variance in the ANOVA model may be violated.

```
# Diagnostic plots for the ANOVA model: Residuals and Q-Q plot
par(mfrow = c(1, 2))
plot(fit2, which = 1:2)
```



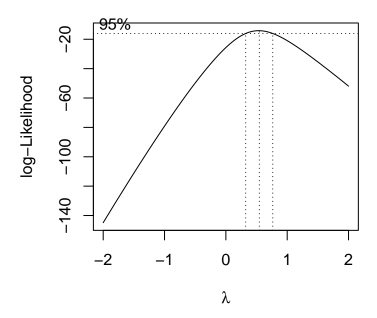
Referring back to Section 3.4.3, to address the issue of unequal variances among groups, a common approach is to apply the Box-Cox transformation. This technique helps stabilize variances and make the data more normally distributed, aligning with the assumptions of ANOVA. The transformation identifies an optimal parameter, λ (lambda), which suggests the most suitable transformation. The value of λ will guide the appropriate transformation to apply, as outlined in Table 3.9 of the textbook, shown below.

■ TABLE 3.9 Variance-Stabilizing Transformations

Relationship Between σ_y and μ	α	$\lambda = 1 - \alpha$	Transformation	Comment
$\sigma_{\rm v} \propto {\rm constant}$	0	1	No transformation	
$\sigma_{\rm v} \propto \mu^{1/2}$	1/2	1/2	Square root	Poisson (count) data
$\sigma_{\rm y} \propto \mu$	1	0	Log	
$\sigma_{\rm y} \propto \mu^{3/2}$	3/2	-1/2	Reciprocal square root	
$\sigma_{\rm y} \propto \mu^2$	2	-1	Reciprocal	

For R users, there are various ways to implement the Box-Cox transformation. Here, we demostrate the use of the boxcox function in the MASS package.

```
# Load the MASS package to apply the Box-Cox transformation
library(MASS)
# Apply the Box-Cox transformation to the ANOVA model to stabilize the variance
fit2bc <- boxcox(fit2, plotit = TRUE)</pre>
```



The Box-Cox plot is to visually identify the best transformation (optimal λ) to apply to the data, which will help meet the assumptions of normality and homogeneity of variance in ANOVA model. The x-axis shows different possible values of λ , typically ranging from -2 to 2, depending on the data. The y-axis represents the log-likelihood of the data under different transformations. The goal is to find the value of λ that maximizes the log-likelihood, indicating the optimal transformation to stabilize variance and improve normality.

```
# Extract the lambda value corresponding to the maximum likelihood estimate
lambda <- fit2bc$x[which.max(fit2bc$y)]
# Print the estimated lambda for the Box-Cox transformation
sprintf("lambda = %.3f", lambda)</pre>
```

[1] "lambda = 0.545"

From the results generated by the boxcox function, we can use the following code to identify the optimal λ value. In this case, the λ value is 0.545, which is close to 0.5. According to Table 3.9 in the textbook, this suggests applying a square root transformation to the response variable PeakDischarge.

Before refitting the ANOVA model, we can first perform the modified Levene's test again to check for equal variances of the square root-transformed PeakDischarge across the Method groups.

```
# library(asbio)
# Perform modified Levene's test again after applying the
# square root transformation (common for stabilizing variance)
modlevene.test(sqrt(df2$PeakDischarge), df2$Method)
```

```
##
## Modified Levene's test of homogeneity of variances
##
## df1 = 3, df2 = 20, F = 0.23917, p-value = 0.86798
```

The p-value of the modified Levene's test is 0.868, which is large, indicating that the variances across the different 'Method' groups are likely similar.

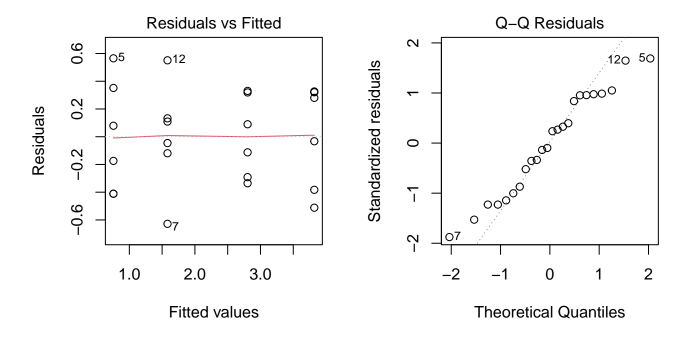
Finally, we fit the ANOVA model using the square root-transformed PeakDischarge.

```
# Fit a new ANOVA model using the square root-transformed 'PeakDischarge'
fit2s <- aov(sqrt(PeakDischarge) ~ Method, data = df2)
# Summary of the new ANOVA results after transformation
summary(fit2s)</pre>
```

```
## Df Sum Sq Mean Sq F value Pr(>F)
## Method    3    32.68    10.895    81.05    2.3e-11 ***
## Residuals    20    2.69    0.134
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

The p-value of the F-test in this new ANOVA table remains extremely small, providing strong evidence that the mean Peak Discharge differs across the Methods. The residual plots confirm that the assumptions of normality and constant variance are satisfied.

```
# Diagnostic plots for the transformed ANOVA model
par(mfrow = c(1, 2))
plot(fit2s, which = 1:2)
```



If there is concern about the normality assumption, as indicated by a light-tailed distribution in the QQ-plot, we can conduct a formal hypothesis test for normality. The goal is to check if the test shows an insignificant result, indicating that the residuals do not deviate significantly from a normal distribution. For example, using the Shapiro-Wilk test for normality, we find that the p-value is 0.414, which is greater than the prespecified significance level of 0.05. This result indicates there is no evidence to suggest that the residuals are from a non-normal distribution.

```
shapiro.test(fit2s$residuals)
```

```
##
## Shapiro-Wilk normality test
##
## data: fit2s$residuals
## W = 0.95877, p-value = 0.4141
```