STATISTICS FOR CLINICAL TRIALS

Applications using R
Foreword

Statistical analysis were done using R version 3.4.3 (2017-11-30), which is freely available from CRAN. You may find convenient to run R through RStudio. Indeed, RStudio offers really great support for editing and running R scripts. You can even organize your work into a project, with version control and automatic reporting built on the fly. I personally choose to work like in the 80s–although I was just a kid at that time–with a simple text editor and an interactive shell available within few key presses. This is possible thanks to Emacs and the brilliant ESS mode.

I replicated all SAS code using SAS University Edition. It can run locally on your computer (using, e.g., Virtual Box) or directly in the cloud. I do not hold a personal licence for SAS (although I could get one from several universities where I am teaching) and so I found this solution particularly handy to compare SAS and R output.

In addition, I provide Stata code to replicate most if not all analyses described in this document. The code has been tested with Stata 13 but should work on any version > 10.

1 Analysis of Clinical Trials using SAS

The following analyses are based on Dmitrienko et al. [2005], with data available online at Analysis of Clinical Trials Using SAS: A Practical Guide.

1.1 The HAMD17 study

Context. This is a multicenter clinical trial comparing experimental drug vs. placebo in patients with major depression disorder. The outcome is the change from baseline after 9 weeks of acute treatment, and efficacy is measured using the total score of the Hamilton depression rating scale (17 items).

This is a classical application of unbalanced design and potential heterogeneity between clinical centres, where there is an unequal number of observations per treatment (here, drug by center).

Here is one of many ways to get data right into R:

```r
raw <- textConnection("100 P 18 100 P 14 100 D 23 100 D 18 100 P 10 100 P 17 100 D 18 100 D 22
100 P 13 100 P 12 100 D 28 100 D 21 100 P 11 100 P 6 100 D 11 100 D 25
100 P 7 100 P 10 100 D 29 100 P 12 100 P 12 100 P 10 100 D 18 100 D 14
101 P 18 101 D 19 101 D 11 101 D 9 101 P 12 101 D 11 102 P 18 102 P 15
102 P 12 102 P 18 102 D 20 102 D 18 102 P 14 102 P 12 102 D 23 102 D 19
102 P 11 102 P 10 102 D 22 102 D 22 102 P 19 102 P 13 102 D 18 102 D 24
102 P 13 102 P 6 102 D 18 102 D 26 102 P 11 102 P 16 102 D 16 102 D 17
102 D 7 102 D 19 102 D 23 102 D 12 103 P 16 103 P 11 103 D 11 103 D 25
103 P 8 103 P 15 103 D 28 103 D 22 103 P 16 103 P 17 103 D 23 103 D 18
103 P 11 103 P -2 103 D 15 103 D 28 103 P 19 103 D 21 103 D 17 104 D 13
104 P 12 104 P 6 104 D 19 104 D 23 104 P 11 104 P 20 104 D 21 104 D 25
104 P 9 104 P 4 104 D 25 104 D 19"
)
d <- scan(raw, what = "character")
rm(raw)
d <- as.data.frame(matrix(d, ncol = 3, byrow = TRUE))
names(d) <- c("center", "drug", "change")
d$change <- as.numeric(as.character(d$change))
d$drug <- relevel(d$drug, ref = "P")
```
Briefly, the idea is to copy and paste the SAS DATALINES instruction as raw text and to scan the flow of characters. The next bit of code uses matrix to arrange the data into a tabular dataset with 3 columns corresponding to center, drug and change score. When transforming this table to a data frame, center and drug will be converted to factors but we need to handle the proper conversion of change to numerical values. Also, note that we set the reference category to the Placebo group to simplify things a bit.

Some basic exploratory graphical analysis follows. In the next chunk, we display the raw data for each centre and highlight the difference between drug and placebo using a trend line (Figure 1). Note the use of aes(group = 1) when calling geom_smooth as there is no real grouping variable in the data structure other than the ones that are already used (drug on the x-axis and center for facetting).

```
p <- ggplot(data = d, aes(x = drug, y = change))
p <- p + geom_jitter(width = .2)
p <- p + geom_smooth(aes(group = 1), method = "lm", se = FALSE, colour = "grey30")
p + facet_grid(~ center) + labs(x = "Drug type", y = "HAMD17 change")
```

Using Hmisc package, we can easily build a Table of summary statistics by drug and center. For simplicity, we will limit the display to the first 3 centers in Table 1.

```
fm <- change ~ drug + center
s <- summary(fm, data = subset(d, center %in% c("100","101","102")),
method = "cross", fun = smean.sd)
```

Table 1: Mean HAMD17 change by drug, center

<table>
<thead>
<tr>
<th>drug</th>
<th>100</th>
<th>101</th>
<th>102</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>Mean</td>
<td>SD</td>
<td>N</td>
</tr>
<tr>
<td>P</td>
<td>13</td>
<td>12</td>
<td>3.4</td>
<td>7</td>
</tr>
<tr>
<td>D</td>
<td>11</td>
<td>21</td>
<td>5.6</td>
<td>7</td>
</tr>
<tr>
<td>Total</td>
<td>24</td>
<td>16</td>
<td>6.3</td>
<td>14</td>
</tr>
</tbody>
</table>

Only 3 out of 5 centres are shown.

Now, let’s consider average change scores by center, which are displayed in Figure 2. First, we need to compute the average score in each group, and then compute the difference between the
two (called delta). This could be done with Hmisc summarize command, but we will rely on the plyr package and its ddply command. What is important is that the results are returned as a data frame to facilitate the use of ggplot data structure in turn.

```r
r <- ddply(d, "center", summarize,
    delta = mean(change[drug == "D"])) - mean(change[drug == "P"]))
p <- ggplot(data = r, aes(x = center, y = delta))
p <- p + geom_point() + geom_hline(yintercept = 0, linetype = 2, colour = "grey30")
p + labs(x = "Center", y = "Difference D-P")
```

Now comes the modeling stage. First, we will analyse the primary endpoint using fixed-effect models. Dmitrienko et al. [2005] provide all the maths that are necessary to understand how to derive various types of sum of squares, and this is further addressed in, e.g., REF., or on Stack Exchange.

Let us first update the formula we used for producing Table 1 to incorporate an interaction term, drug:center (in R, drug * center will expand to drug + center + drug:center):

```r
fm <- change ~ drug * center
replications(change ~ drug:center, data = d)
## $'drug:center'
## center
## drug 100 101 102 103 104
## P 13 7 14 10 6
## D 11 7 16 9 7
```

As can be seen, data are slightly imbalanced for all but centre 101.

By default, R computes so-called “sequential” Type I sum of squares (SS), and here is what we get when using a standard combination of lm (to compute parameter estimates) and anova (to build the ANOVA table for the regression model):

```r
options(contrasts = c("contr.sum", "contr.poly"))
m <- lm(fm, data = d)
anova(m)
```

### Analysis of Variance Table

<table>
<thead>
<tr>
<th></th>
<th>Df</th>
<th>Sum Sq</th>
<th>Mean Sq</th>
<th>F value</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>drug</td>
<td>1</td>
<td>105.3</td>
<td>105.3</td>
<td>4.54</td>
<td>0.042</td>
</tr>
<tr>
<td>center</td>
<td>4</td>
<td>41.0</td>
<td>10.25</td>
<td>0.4</td>
<td>0.004</td>
</tr>
<tr>
<td>drug:center</td>
<td>4</td>
<td>41.0</td>
<td>10.25</td>
<td>0.4</td>
<td>0.004</td>
</tr>
</tbody>
</table>
```
The `car` package allows to work with Type II and Type III SS. Type III SSs, also called partial or Yates’ weighted squares of means are the default in Stata, SPSS or SAS. Stata does not even offer Type II SS. So, if we are interested in computing Type II sum of squares in R, we could call `Anova` like this:

```r
car::Anova(m, type = "II")
```

## Anova Table (Type II tests)

<table>
<thead>
<tr>
<th>Sum Sq</th>
<th>Df</th>
<th>F value</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>drug</td>
<td>889.78</td>
<td>1</td>
<td>40.1528</td>
</tr>
<tr>
<td>center</td>
<td>87.14</td>
<td>4</td>
<td>0.9831</td>
</tr>
<tr>
<td>drug:center</td>
<td>507.45</td>
<td>4</td>
<td>5.7249</td>
</tr>
<tr>
<td>Residuals</td>
<td>1994.38</td>
<td>90</td>
<td></td>
</tr>
</tbody>
</table>

---

## Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

Type III analysis is readily obtained by replacing `type = "II"` with `type = "III"` as shown in the next code block. It should be noted that without altering the default contrast treatment that are used by R, as we did in the above chunk, we would not get the correct results for the Type III analysis.

```r
car::Anova(m, type = "III")
```

## Anova Table (Type III tests)

<table>
<thead>
<tr>
<th>Sum Sq</th>
<th>Df</th>
<th>F value</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Intercept)</td>
<td>22344.6</td>
<td>1</td>
<td>1008.3442</td>
</tr>
<tr>
<td>drug</td>
<td>709.8</td>
<td>1</td>
<td>32.0320</td>
</tr>
<tr>
<td>center</td>
<td>91.5</td>
<td>4</td>
<td>1.0318</td>
</tr>
<tr>
<td>drug:center</td>
<td>507.4</td>
<td>4</td>
<td>5.7249</td>
</tr>
<tr>
<td>Residuals</td>
<td>1994.4</td>
<td>90</td>
<td></td>
</tr>
</tbody>
</table>

---

## Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

Note that in the case of Type III SS, we can also use the base command `drop1` and we will get similar results:

```r
drop1(m, scope = ~ ., test = "F")
```

## Single term deletions

<table>
<thead>
<tr>
<th>Change</th>
<th>Df</th>
<th>Sum of Sq</th>
<th>RSS</th>
<th>AIC</th>
<th>F value</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;none&gt;</td>
<td>1994.4</td>
<td>319.29</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>drug</td>
<td>1</td>
<td>709.82</td>
<td>2764.2</td>
<td>347.74</td>
<td>32.0320</td>
<td>1.783e-07 ***</td>
</tr>
<tr>
<td>center</td>
<td>4</td>
<td>91.46</td>
<td>2085.8</td>
<td>315.78</td>
<td>1.0318</td>
<td>0.3953130</td>
</tr>
<tr>
<td>drug:center</td>
<td>4</td>
<td>507.45</td>
<td>2580.1</td>
<td>333.96</td>
<td>5.7249</td>
<td>0.0003761 ***</td>
</tr>
</tbody>
</table>

---

## Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

To sum up, the results from the different approaches are exposed in Table 2.
Sidenote. Here is how we could compute the parameter estimates and the SS corresponding to the drug effect in the case of a Type III analysis. The code follows that posted on Stack Exchange, with minor adaptation. How this works is quite simple: We first get the design matrix stored in our model \( m \) and then solve the “normal equations” \((X'X)\hat{\beta} = X'y\) in order to get \( \hat{\beta} = (X'X)^{-1}X'y\).

\[
\begin{align*}
D & \leftarrow \text{model.matrix}(m) \quad \text{## design matrix} \\
\text{bhat} & \leftarrow \text{solve}(t(D) \times D) \times t(D) \times d$change \quad \text{## beta parameters} \\
\text{get.ss} & \leftarrow \text{function}(C) \{
\text{require(MASS)} \\
\text{teta} & \leftarrow C \times \text{bhat} \\
M & \leftarrow C \times \text{ginv}(t(D) \times D) \times t(C) \\
\text{SSH} & \leftarrow t(teta) \times \text{ginv}(M) \times teta \\
\text{return(as.numeric(SSH))}
\}
\end{align*}
\]

\[
\text{## SS(drug|center,drug:center)}
\text{get.ss(matrix(c(0,1,0,0,0,0,0,0,0,0), nrow = 1, ncol = 10))}
\]

\[
\text{## [1] 709.8196}
\]

Table 2: Overview of fixed-effects analysis for the HAMD17 study

<table>
<thead>
<tr>
<th></th>
<th>(a) Type I SS</th>
<th>(b) Type II SS</th>
<th>(c) Type III SS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sum Sq</td>
<td>F value</td>
<td>Pr(&gt;F)</td>
<td>Sum Sq</td>
</tr>
<tr>
<td>drug</td>
<td>888.04</td>
<td>40.07</td>
<td>0.0000</td>
</tr>
<tr>
<td>center</td>
<td>87.14</td>
<td>0.98</td>
<td>0.4209</td>
</tr>
<tr>
<td>drug:center</td>
<td>507.45</td>
<td>5.72</td>
<td>0.0004</td>
</tr>
</tbody>
</table>

The authors later used the Gail-Simon test[Gail and Simon, 1985] to test for qualitative interaction between treatment and strata. The corresponding two-tailed Likelihood ratio test is implemented in the QualInt package.

\[
\begin{align*}
\text{library}(\text{QualInt}) \\
\text{with(d, qualint(change, drug, center, test = "LRT"))}
\end{align*}
\]

\[
\begin{align*}
\text{## Call:} \\
\text{## qualint(y = change, treatment = drug, subgrp = center, test = "LRT")} \\
\text{## Type:} \\
\text{## continuous} \\
\text{## Estimating Results for Mean Difference:} \\
\text{## Estimate Std. Error Lower CI Upper CI} \\
\text{## 100} & -8.944 & 1.922 & -12.711 & -5.177 \\
\text{## 101} & 4.429 & 1.601 & 1.290 & 7.567 \\
\text{## 102} & -5.571 & 1.517 & -8.544 & -2.599 \\
\text{## 103} & -7.578 & 2.877 & -13.217 & -1.939 \\
\text{## 104} & -10.381 & 2.793 & -15.856 & -4.906 \\
\text{## Test:} \\
\text{## LRT} \\
\text{## p-value:} \\
\text{## 0.02968} \\
\text{## Power:} \\
\text{## 0.58} \\
\text{## Alpha:} \\
\text{## 0.05}
\end{align*}
\]
This R package even provides a graphical method when specifying options test = "IBGA" and plotout = TRUE (Figure 3). The IBGA method relies on simultaneous 95% confidence intervals as described in Pan and Wolfe [1997].

1.2 The Urinary incontinence trial

Context. This is a subset of data collected in an RCT on urinary incontinence where the primary endpoint was the percent change from baseline of number of incontinence episodes per week over an 8-week period. Patients were initially randomized into one of three strata depending on the baseline frequency of incontinence episodes.

This is an example of the use of stratified non-parametric analysis.

This time, we managed to get data in the right format using this little R script: urininc.R. Assuming it is located in the current working directory, we can source it into R and we will get a data frame named d.

```r
source("./urininc.R")
str(d)
```

To summarize the data, we can again make use of Hmisc summary for "crossed" data.

```r
s <- summary(change ~ group + strata, data = d, method = "cross", overall = FALSE)
```

<table>
<thead>
<tr>
<th>group</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>Missing</td>
<td>N</td>
</tr>
<tr>
<td>Placebo</td>
<td>40</td>
<td>0</td>
<td>−29.0</td>
</tr>
<tr>
<td>Drug</td>
<td>39</td>
<td>1</td>
<td>−24.2</td>
</tr>
</tbody>
</table>
As can be seen, there is a higher number of missing values in strata 2 (around 20% in both groups) and larger variations on average between the two group in the third strata. Next, we displayed the distribution of the percent change in frequency of incontinence episodes as density curves in Figure 4. Instead of relying on `geom_density`, we use the rather generic `geom_lines` with an extra `stat=` parameter.

```r
p <- ggplot(data = d, aes(x = change, colour = group))
p <- p + geom_line(stat = "density", adjust = 1.2) + facet_grid(~ strata)
p + scale_x_continuous(limits = c(-100, 150)) + labs(x = "Percent change", y = "Density")
```

The authors used the van Elteren test [van Elteren, 1960], which can be regarded as an extension of the Wilcoxon rank sum test for stratified data where larger weights are assigned to rank sums from smaller strata. An alternative is the “aligned rank test” proposed by Hodges and Lehman [1962] as discussed by Mehrotra et al. [2010]. In R, there is an old version that is mentioned on the R listserv (August 2005), but for now we will use the `coin` package as shown below:

```r
library(coin)
dc <- subset(d, complete.cases(d))
independence_test(change ~ group | strata, data = dc,
     ytrafo = function(data) trafo(data, numeric_trafo = rank,
                        block = dc$strata),
     teststat = "quad")
```

Although we get different results from the authors, we would reach the same conclusion, namely that there is an effect of the treatment on the outcome after adjusting for the centre effect. We will get, however, closer results ($p = 0.02369$ for the row mean squares test statistic) if we simply remove the `scores=` option when calling SAS PROC FREQ [Stokes et al., 2012]:

```r
PROC FREQ;
   TABLES strata*group*change / noprint cmh2;
RUN;
```

In comparison, as noted by the authors, a Type III ANOVA would yield non-significant result about the effect of drug on change scores.

```r
m <- lm(change ~ group + strata, data = d)
car::Anova(m, type = "III")
```
## Anova Table (Type III tests)

<table>
<thead>
<tr>
<th></th>
<th>Sum Sq</th>
<th>Df</th>
<th>F value</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Intercept)</td>
<td>176982</td>
<td>1</td>
<td>25.7904</td>
<td>9.499e-07 ***</td>
</tr>
<tr>
<td>group</td>
<td>9602</td>
<td>1</td>
<td>1.3993</td>
<td>0.2384</td>
</tr>
<tr>
<td>strata</td>
<td>8094</td>
<td>2</td>
<td>0.5898</td>
<td>0.5555</td>
</tr>
<tr>
<td>Residuals</td>
<td>1228358</td>
<td>179</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

---

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

### 1.3 The Severe sepsis trial

**Context.** This is a placebo-controlled RCT examining the effect of an experimental drug on 28-day all-cause mortality in patients with severe sepsis. Patients were allocated to one of four strata depending on their APACHE II score [Knaus et al., 1985].

This is a classical application of stratified analysis of a binary outcome (dead/alive).

To enter the data in R, we will input individual values of the three-way Table of events as an array. Note that it would also be possible to create two matrix objects and then bind into a 3-dimensional table. In what follows, we write data for the treated group first. Note that when using array, data should be entered column-wise (there is no byrow = option as in matrix).

```r
varnames <- list(strata = 1:4,
                 status = c("Dead", "Alive", "Total"),
                 group = c("Experimental", "Placebo"))

d <- array(c(33,49,48,80,185,169,156,130,218,218,204,210,
             26,57,58,118,189,165,104,123,215,222,162,241),
           dim = c(4,3,2), dimnames = varnames)
```

Note also that the third column ("Total") can be safely omitted as margins can be computed automatically with R, e.g.:

```r
addmargins(d[,,-3], c(1,2))

d <- d[,,-3]

dim(d)
## [1] 4 2 2
```

An alternative representation of this array-based Table is provided by R’s flat tables (ftable), in long or wide format; see Table 4 for the wide format using ftable(d, row.vars = 1, col.vars = c(3,2)):

```r
ftable(d)
## group Experimental Placebo
## strata status
## 1 Dead 33 26
## 2 Alive 185 189
## 3 Dead 49 57
## 4 Alive 169 165
## 4 Dead 48 58
## 4 Alive 156 104
## 4 Alive 80 118
## 4 Alive 130 123
```

The following code is used to depict the situation in graphical terms:

```r
dd <- as.data.frame(ftable(d))
r <- ddply(dd, c("strata", "group"), mutate, prop = Freq/sum(Freq))
```
<table>
<thead>
<tr>
<th>group:</th>
<th>Experimental</th>
<th>Placebo</th>
</tr>
</thead>
<tbody>
<tr>
<td>strata status:</td>
<td>Dead</td>
<td>Alive</td>
</tr>
<tr>
<td>1</td>
<td>33</td>
<td>185</td>
</tr>
<tr>
<td>2</td>
<td>49</td>
<td>169</td>
</tr>
<tr>
<td>3</td>
<td>48</td>
<td>156</td>
</tr>
<tr>
<td>4</td>
<td>80</td>
<td>130</td>
</tr>
</tbody>
</table>

Table 4: 28-day mortality data from the 1690-patient sepsis study

Figure 5: Proportion of patients who died by the end of the study

```r
p <- ggplot(subset(r, status == "Dead"), aes(x = prop, y = group))
p <- p + geom_point() + facet_wrap(~ strata, nrow = 2)
p + scale_x_continuous(limits = c(0,0.5)) + labs(x = "Proportion deads", y = "")
library(vcd)
cotabplot(d, 1)
```

Based on a logistic regression model, the authors presented a summary of a Type III analysis of effects. Here is what can be done in R. First, we will slightly rearrange the data table so that we have a working data frame with total counts for success (here, dead patients) and failure (here, patients still alive) in separate columns, together with columns describing strata and treatment levels.

```r
n <- rbind(d[,1:2,1], d[,1:2,2])
rownames(n) <- NULL
n <- as.data.frame(n)
n$strata <- gl(4, 1)
n$group <- gl(2, 4, labels = c("Experimental", "Placebo"))
n$group <- relevel(n$group, ref = "Placebo")
```

Then, since we are working with grouped or aggregated data, we will use the `cbind()` option to R’s `glm`, as shown below. Note that we also ask to use SAS treatment contrast for the `strata` factor, in order to ensure that the fourth level is used as the reference category. Type III analysis is readily available within the `car` package.

```r
m <- glm(cbind(Dead, Alive) ~ group + strata, data = n, family = binomial,
        contrasts = list(strata = "contr.SAS"))
car::Anova(m, type = "III")
```
## Analysis of Deviance Table (Type III tests)

## Response: cbind(Dead, Alive)

<table>
<thead>
<tr>
<th></th>
<th>LR Chisq Df Pr(&gt;Chisq)</th>
</tr>
</thead>
<tbody>
<tr>
<td>group</td>
<td>6.989 1 0.008201 **</td>
</tr>
<tr>
<td>strata</td>
<td>105.609 3 &lt; 2.2e-16 ***</td>
</tr>
</tbody>
</table>

---

# Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Finally, profile likelihood 95% confidence intervals are simply obtained using `confint()` which will call the appropriate profile method depending on the kind of model at hand.

```r
exp(confint(m))
```

```r
# 2.5 % 97.5 %
# (Intercept) 0.6412481 0.9318192
# group1 1.0392135 1.2965299
# strata1 0.1443251 0.2807120
# strata2 0.3048534 0.5420573
# strata3 0.3970567 0.7146984
```

### 1.4 The dose-finding hypertension trial

**Context.** This trial aimed to compare low, medium and high doses of a new antihypertensive drug to a placebo. The primary efficacy variable that is being considered in this study is diastolic blood pressure.

This example is used to illustrate various methods to deal with multiple testing issues. In what follows we will work with p-values (raw data are not available) estimated when comparing all four groups (P, placebo vs. L, M, and H, the low, medium and high dose groups).

The `p.adjust()` command can be used to compute various “adjusted” p-values, the default being the step-down method proposed by Holm [1979].

```r
pv <- c(0.047, 0.0167, 0.015)
p.adjust(pv, method = "bonferroni")
```

```r
# [1] 0.1410 0.0501 0.0450
```
The Šidák method is not available in \texttt{p.adjust()} but it is not difficult to implement a custom function to perform this correction which amounts to update the nominal \( \alpha \) level with \( 1 - (1 - \alpha)^{1/n} \), that is:

\[
f <- \text{function}(x) \ (1-(1-x)^{\text{length}(x)})
\]

\[
f(pvals)
\]

\[
\# [1] 0.13447682 0.04926799 0.04432838
\]

Alternatively, one can dig into the \texttt{multtest} package by Dudoit and van der Laan [2008], available on \url{http://www.bioconductor.org} (see the \texttt{mt.rawp2adjp()} command).

Contrary to the preceding results, Holm’s adjusted \( p \)-values will all be < 0.05 as illustrated below:

\[
p.adjust(pvals, \text{method} = "holm")
\]

\[
\# [1] 0.047 0.045 0.045
\]

And here is a comparison of Holm and Hommel’s adjusted \( p \)-values for the second scenario (Table 5):

\[
pvals <- \text{c}(0.047, 0.027, 0.015) \# \text{scenario 2}
\]

\[
p.adjust(pvals, \text{method} = "holm")
\]

\[
\# [1] 0.054 0.054 0.045
\]

\[
p.adjust(pvals, \text{method} = "hommel")
\]

\[
\# [1] 0.0470 0.0470 0.0405
\]

Finally, Hommel’s method is compared to Hochberg’s approach for the third scenario:

\[
pvals <- \text{c}(0.053, 0.026, 0.017) \# \text{scenario 3}
\]

\[
p.adjust(pvals, \text{method} = "hochberg")
\]

\[
\# [1] 0.053 0.052 0.051
\]

\[
p.adjust(pvals, \text{method} = "hommel")
\]

\[
\# [1] 0.053 0.052 0.039
\]

One can also look into the \texttt{cherry} package [Goeman and Solari, 2011] whose vignette includes a comparison of Simes vs. Hommel or Fisher approach to multiple testing, as well as examples of closed testing methods.

### 1.5 The allergen-induced asthma trial

**Context.** Data comes from a trial designed to assess the efficacy profile of a bronchodilator in allergen-induced asthma. There are 20 patients that were randomly assigned to receive either an experimental drug or a placebo [Taylor et al., 1991]. The forced expiratory volume in one second (FEV) was used to measure how the drug attenuated bronchoconstriction, and FEV curves were averaged at each time point in both groups (Table 6). The therapeutic effect was the time to the onset of action—that is, the first time point at which clinically and statistically significant separation between the FEV curves is observed.

<table>
<thead>
<tr>
<th>Scenario</th>
<th>L vs. P</th>
<th>M vs. P</th>
<th>H vs. P</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.047</td>
<td>0.0167</td>
<td>0.015</td>
</tr>
<tr>
<td>2</td>
<td>0.047</td>
<td>0.027</td>
<td>0.015</td>
</tr>
<tr>
<td>3</td>
<td>0.053</td>
<td>0.026</td>
<td>0.017</td>
</tr>
</tbody>
</table>

Table 5: \( p \)-values obtained from different approaches
Beside stepwise approaches relying on data-driven ordering of p-values—this is also known as closed testing—fixed-sequence testing methods are used when we are interested in prespecified sequences of hypotheses. This is illustrated in the next example.

<table>
<thead>
<tr>
<th></th>
<th>Experimental drug</th>
<th>Placebo</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time (hours)</td>
<td>n</td>
<td>Mean</td>
</tr>
<tr>
<td>0.25</td>
<td>10</td>
<td>0.58</td>
</tr>
<tr>
<td>0.5</td>
<td>10</td>
<td>0.62</td>
</tr>
<tr>
<td>0.75</td>
<td>10</td>
<td>0.51</td>
</tr>
<tr>
<td>1</td>
<td>10</td>
<td>0.34</td>
</tr>
<tr>
<td>2</td>
<td>10</td>
<td>-0.06</td>
</tr>
<tr>
<td>3</td>
<td>10</td>
<td>0.05</td>
</tr>
</tbody>
</table>

Table 6: Reduction in FEV measurements from baseline by time after the allergen challenge

To load the data, we will use a simple matrix to store the values displayed in Table 6 and then reshape the matrix to a “tidy” data frame using `melt`, although this last step is not really necessary.

```r
tmp <- matrix(c(0.25,10,0.58,0.29,10,0.71,0.35,
                 0.5,10,0.62,0.31,10,0.88,0.33,
                 0.75,10,0.51,0.33,10,0.73,0.36,
                 1,10,0.34,0.27,10,0.68,0.29,
                 2,10,-0.06,0.22,10,0.37,0.25,
                 3,10,0.05,0.23,10,0.43,0.28),
               nrow = 6, byrow = TRUE)
colnames(tmp) <- c("time", "N0", "Mean0", "SD0", "N1", "Mean1", "SD1")
d <- melt(as.data.frame(tmp), id.vars = 1, measure.vars = c(3,4,6,7))
```

Note that it is quite easy to go back to the matrix form by using `dcast` as shown below:

```r
r <- ddply(dcast(d, time ~ variable), "time", mutate,
            diff = Mean1 - Mean0, se = sqrt((1/10+1/10)*(SD0^2+SD1^2)/2))
```

In what follows, we compute the lowest bound of a 95% confidence interval and display its value at consecutive time points. This amounts to looking at results from sequential testing, i.e. determine the first statistically significant difference. Equivalently, we could rely on simple Student t-test.

```r
p <- ggplot(r, aes(x = time, y = diff))
p <- p + geom_line() + geom_point()
p <- p + geom_line(aes(x = time, y = diff - qt(0.95, 20-2) * se), linetype = 2)
p <- p + scale_x_continuous(breaks = seq(0, 3, by = 1))
p <- p + scale_y_continuous(breaks = seq(-0.2, 0.5, by = 0.1))
p <- p + geom_hline(aes(yintercept = 0))
p + labs(x = "Time (hours)", y = "Treatment difference (95% Lower CI)"
```

Note that to display error bars instead of the lower bound for the 95% confidence interval, we would use:

```r
p <- p + geom_errorbar(aes(ymin = diff - qt(0.975, 20-2) * se, 
                           ymax = diff + qt(0.975, 20-2) * se), 
                          width = .1)
```

This approach, however, does not control the familywise error rate. Looking at treatment differences from the last measurement to the first (“step-down” approach) suggests that a significant difference at one hour, and not 30 minutes as in the previous case.
Figure 7: Treatment comparisons in the asthma study
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