A Handbook of Statistical Analyses Using R — 3rd Edition

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CHAPTER 13

Analyzing Longitudinal Data I: Computerized Delivery of Cognitive Behavioral Therapy – Beat the Blues

- 13.1 Introduction
- 13.2 Analyzing Longitudinal Data

13.3 Analysis Using R

We shall fit both random intercept and random intercept and slope models to the data including the baseline BDI values (pre.bdi), treatment group, drug, and length as fixed effect covariates. Linear mixed effects models are fitted in R by using the lmer function contained in the lme4 package (Bates and Sarkar, 2014, Pinheiro and Bates, 2000, Bates, 2005), but an essential first step is to rearrange the data from the 'wide form' in which they appear in the BtheB data frame into the 'long form' in which each separate repeated measurement and associated covariate values appear as a separate row in a data.frame. This rearrangement can be made using the following code:

```
R> data("BtheB", package = "HSAUR3")
R> BtheB$subject <- factor(rownames(BtheB))
R> nobs <- nrow(BtheB)
R> BtheB_long <- reshape(BtheB, idvar = "subject",
+ varying = c("bdi.2m", "bdi.3m", "bdi.5m", "bdi.8m"),
+ direction = "long")
R> BtheB_long$time <- rep(c(2, 3, 5, 8), rep(nobs, 4))
such that the data are now in the form (here shown for the first three subjects)</pre>
```

R> subset(BtheB_long, subject %in% c("1", "2", "3"))

	drug	length	treatment	bdi.pre	subject	time	bdi
1.2m	No	>6m	TAU	29	1	2	2
2.2m	Yes	>6m	BtheB	32	2	2	16
3.2m	Yes	<6m	TAU	25	3	2	20
1.3m	No	>6m	TAU	29	1	3	2
2.3m	Yes	>6m	BtheB	32	2	3	24
3.3m	Yes	<6m	TAU	25	3	3	NA
1.5m	No	>6m	TAU	29	1	5	NA
2.5m	Yes	>6m	BtheB	32	2	5	17
3.5m	Yes	<6m	TAU	25	3	5	NA
1.8m	No	>6m	TAU	29	1	8	NA
2.8m	Yes	>6m	BtheB	32	2	8	20
3.8m	Yes	<6m	TAU	25	3	8	NA

The resulting data frame BtheB_long contains a number of missing values

```
R> data("BtheB", package = "HSAUR3")
R> layout(matrix(1:2, nrow = 1))
R> ylim <- range(BtheB[,grep("bdi", names(BtheB))],</pre>
                  na.rm = TRUE)
   tau <- subset(BtheB, treatment == "TAU")[,</pre>
R>
       grep("bdi", names(BtheB))]
  boxplot(tau, main = "Treated as Usual", ylab = "BDI",
R>
           xlab = "Time (in months)", names = c(0, 2, 3, 5, 8),
           ylim = ylim)
R>
  btheb <- subset(BtheB, treatment == "BtheB")[,</pre>
       grep("bdi", names(BtheB))]
R> boxplot(btheb, main = "Beat the Blues", ylab = "BDI",
           xlab = "Time (in months)", names = c(0, 2, 3, 5, 8),
           ylim = ylim)
```

Treated as Usual

Beat the Blues

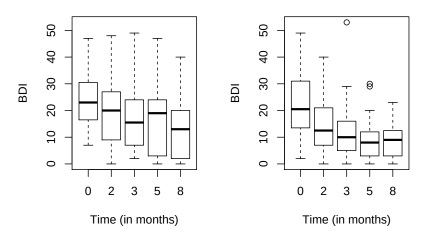


Figure 13.1 Boxplots for the repeated measures by treatment group for the BtheB data.

and in applying the lmer function these will be dropped. But notice it is only the missing values that are removed, not participants that have at least one missing value. All the available data is used in the model fitting process. The lmer function is used in a similar way to the lm function met in Chapter 6 with the addition of a random term to identify the source of the repeated measurements, here subject. We can fit the two models (??) and (??) and test which is most appropriate using

```
R> library("lme4")
```

```
R> BtheB_lmer1 <- lmer(bdi ~ bdi.pre + time + treatment + drug +
        length + (1 | subject), data = BtheB_long,
        REML = FALSE, na.action = na.omit)
R> BtheB_lmer2 <- lmer(bdi ~ bdi.pre + time + treatment + drug +
        length + (time | subject), data = BtheB_long,
        REML = FALSE, na.action = na.omit)
R> anova(BtheB_lmer1, BtheB_lmer2)
Data: BtheB_long
Models:
BtheB_lmer1: bdi ~ bdi.pre + time + treatment + drug + length + (1 | subject)
BtheB_lmer2: bdi ~ bdi.pre + time + treatment + drug + length + (time | subject)
                      BIC logLik deviance Chisq Chi Df
          Df AIC
BtheB_lmer1 8 1887.5 1916.6 -935.75 1871.5
BtheB_lmer2 10 1891.0 1927.4 -935.52 1871.0
                                     1871.0 0.4542
           Pr(>Chisq)
BtheB_lmer1
               0.7969
BtheB_lmer2
R> summary(BtheB_lmer1)
Linear mixed model fit by maximum likelihood ['lmerMod']
Formula:
bdi ~ bdi.pre + time + treatment + drug + length + (1 | subject)
   Data: BtheB long
             BIC logLik deviance df.resid
  1887.5 1916.6 -935.7 1871.5
Scaled residuals:
   Min 1Q Median
                           30
-2.6975 -0.5026 -0.0638 0.4124 3.8203
Random effects:
                    Variance Std.Dev.
 Groups Name
                            6.984
 subject (Intercept) 48.78
 Residual
                    25.14
                             5.014
Number of obs: 280, groups: subject, 97
Fixed effects:
              Estimate Std. Error t value
(Intercept)
               5.59239
                         2.24244 2.494
bdi.pre
               0.63968
                          0.07789
                                   8.212
              -0.70476
                         0.14639 -4.814
time
treatmentBtheB -2.32908
                         1.67036 -1.394
drugYes
              -2.82495
                         1.72684 -1.636
length>6m
               0.19708
                         1.63832
                                  0.120
Correlation of Fixed Effects:
           (Intr) bdi.pr time
                              trtmBB drugYs
bdi.pre
           -0.682
time
           -0.238 0.020
tretmntBthB -0.390 0.121 0.018
drugYes
           -0.073 -0.237 -0.022 -0.323
length>6m
           -0.243 -0.242 -0.036 0.002
                                       0.157
```

Figure 13.2 R output of the linear mixed-effects model fit for the BtheB data.

The summary method for *lmer* objects doesn't print *p*-values for Gaussian mixed models because the degrees of freedom of the *t* reference distribution are

not obvious. However, one can rely on the asymptotic normal distribution for computing univariate p-values for the fixed effects using the cftest function from package **multcomp**. The asymptotic p-values are given in Figure 13.3.

```
R> cftest(BtheB_lmer1)
         Simultaneous Tests for General Linear Hypotheses
Fit: lmer(formula = bdi ~ bdi.pre + time + treatment + drug + length +
    (1 | subject), data = BtheB_long, REML = FALSE, na.action = na.omit)
Linear Hypotheses:
                    Estimate Std. Error z value Pr(>|z|)
                                2.24244
(Intercept) == 0
                     5.59239
                                          2.494
bdi.pre == 0
                     0.63968
                                0.07789
                                          8.212 2.22e-16
                    -0.70476
time == 0
                                0.14639
                                         -4.814 1.48e-06
treatmentBtheB == 0 -2.32908
                                1.67036
                                         -1.394
drugYes == 0
                    -2.82495
                                1.72684
                                         -1.636
                                                   0.1019
length>6m == 0
                     0.19708
                                1.63832
                                           0.120
(Univariate p values reported)
```

Figure 13.3 R output of the asymptotic p-values for linear mixed-effects model fit for the BtheB data.

We can check the assumptions of the final model fitted to the BtheB data, i.e., the normality of the random effect terms and the residuals, by first using the ranef method to predict the former and the residuals method to calculate the differences between the observed data values and the fitted values, and then using normal probability plots on each. How the random effects are predicted is explained briefly in Section ??. The necessary R code to obtain the effects, residuals, and plots is shown with Figure 13.4. There appear to be no large departures from linearity in either plot.

```
ANALYSIS USING R
```

```
R> layout(matrix(1:2, ncol = 2))
R> qint <- ranef(BtheB_lmer1)$subject[["(Intercept)"]]</pre>
```

R> qres <- residuals(BtheB_lmer1)</pre> R> qqnorm(qint, ylab = "Estimated random intercepts",

xlim = c(-3, 3), ylim = c(-20, 20),

main = "Random intercepts")

R> qqline(qint)

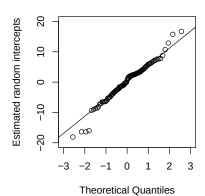
R> qqnorm(qres, xlim = c(-3, 3), ylim = c(-20, 20),

ylab = "Estimated residuals",

main = "Residuals")

R> qqline(qres)

Random intercepts



20 Estimated residuals 10 -20 2 -2 0 3 Theoretical Quantiles

Residuals

7

Quantile-quantile plots of predicted random intercepts and residuals for the random intercept model BtheB_lmer1 fitted to the BtheB data.

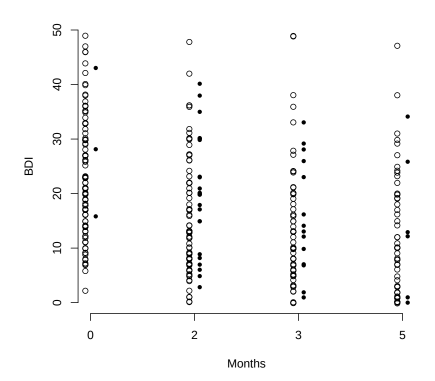


Figure 13.5 Distribution of BDI values for patients that do (circles) and do not (bullets) attend the next scheduled visit.

Bibliography

- Bates, D. (2005), "Fitting linear mixed models in R," R News, 5, 27-30, URL http://CRAN.R-project.org/doc/Rnews/.
- Bates, D. and Sarkar, D. (2014), *lme4: Linear Mixed-Effects Models Using S4 Classes*, URL http://CRAN.R-project.org/package=lme4, R package version 1.1-5.
- Pinheiro, J. C. and Bates, D. M. (2000), *Mixed-Effects Models in S and S-PLUS*, New York, USA: Springer-Verlag.