A Handbook of Statistical Analyses Using R —2nd Edition

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CHAPTER 13

Analysing Longitudinal Data II – Generalised Estimation Equations and Linear Mixed Effect Models: Treating Respiratory Illness and Epileptic Seizures

- 13.1 Introduction
- 13.2 Methods for Non-normal Distributions
- 13.3 Analysis Using R: GEE

13.3.1 Beat the Blues Revisited

To use the gee function, package gee (Carey et al., 2012) has to be installed and attached:

```
R> library("gee")
```

The gee function is used in a similar way to the lme function met in Chapter 12 with the addition of the features of the glm function that specify the appropriate error distribution for the response and the implied link function, and an argument to specify the structure of the working correlation matrix. Here we will fit an independence structure and then an exchangeable structure. The R code for fitting generalised estimation equations to the BtheB_long data (as constructed in Chapter 12) with identity working correlation matrix is as follows (note that the gee function assumes the rows of the data.frame BtheB_long to be ordered with respect to subjects):

```
R> osub <- order(as.integer(BtheB_long$subject))
R> BtheB_long <- BtheB_long[osub,]
R> btb_gee <- gee(bdi ~ bdi.pre + trt + length + drug,
+ data = BtheB_long, id = subject, family = gaussian,
+ corstr = "independence")
and with exchangeable correlation matrix:
R> btb_gee1 <- gee(bdi ~ bdi.pre + trt + length + drug,
+ data = BtheB_long, id = subject, family = gaussian,
+ corstr = "exchangeable")</pre>
```

The summary method can be used to inspect the fitted models; the results are shown in Figures 13.1 and 13.2.

```
R> summary(btb_gee)
Model:
                          Identity
 Variance to Mean Relation: Gaussian
 Correlation Structure:
                          Independent
Coefficients:
          Estimate Naive S.E. Naive z Robust S.E. Robust z
(Intercept) 3.569 1.4833 2.41 2.2695
                                                  1.572
                              10.32
             0.582
                       0.0564
                                          0.0916
                                                   6.355
bdi.pre
            -3.237
                       1.1296
                                          1.7746
trtBtheB
                              -2.87
                                                  -1.824
                       1.1380
length>6m
             1.458
                                1.28
                                          1.4826
                                                    0.983
drugYes
             -3.741
                       1.1766
                                -3.18
                                          1.7827
                                                   -2.099
Estimated Scale Parameter: 79.3
```

Figure 13.1 R output of the summary method for the btb_gee model (slightly abbreviated).

13.3.2 Respiratory Illness

The baseline status, i.e., the status for month == 0, will enter the models as an explanatory variable and thus we have to rearrange the *data.frame* respiratory in order to create a new variable baseline:

The new variable nstat is simply a dummy coding for a poor respiratory status. Now we can use the data resp to fit a logistic regression model and GEE models with an independent and an exchangeable correlation structure as follows.

```
R> summary(btb_gee1)
Model:
Tink:
                          Identity
 Variance to Mean Relation: Gaussian
 Correlation Structure:
                          Exchangeable
Coefficients:
          Estimate Naive S.E. Naive z Robust S.E. Robust z
(Intercept) 3.023 2.3039 1.3122 2.2320 1.3544
                                                  7.7583
             0.648
                       0.0823 7.8741
                                          0.0835
bdi.pre
             -2.169
                      1.7664 -1.2281
                                          1.7361 -1.2495
trtBtheB
            -0.111
                      1.7309 -0.0643
                                          1.5509 -0.0718
length>6m
                       1.8257 -1.6430
drugYes
             -3.000
                                          1.7316 -1.7323
Estimated Scale Parameter: 81.7
```

Figure 13.2 R output of the summary method for the btb_gee1 model (slightly abbreviated).

```
+ + age, data = resp, family = "binomial", id = subject,
+ corstr = "exchangeable", scale.fix = TRUE,
+ scale.value = 1)
```

The estimated treatment effect taken from the exchangeable structure GEE model is 1.299 which, using the robust standard errors, has an associated 95% confidence interval

These values reflect effects on the log-odds scale. Interpretation becomes simpler if we exponentiate the values to get the effects in terms of odds. This gives a treatment effect of 3.666 and a 95% confidence interval of

The odds of achieving a 'good' respiratory status with the active treatment is between about twice and seven times the corresponding odds for the placebo.

```
R> summary(resp_glm)
Call:
glm(formula = status ~ centre + trt + gender + baseline + age,
    family = "binomial", data = resp)
Deviance Residuals:
  Min 1Q Median
                           30
                                  Max
-2.315 -0.855
               0.434
                                1.925
                         0.895
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
(Intercept) -0.90017
                        0.33765
                                  -2.67
                                          0.0077
             0.67160
                        0.23957
                                   2.80
                                          0.0051
centre2
             1.29922
                        0.23684
                                   5.49 4.1e-08
trttrt
                        0.29467
             0.11924
                                   0.40
                                          0.6857
gendermale
                         0.24129
baselinegood 1.88203
                                   7.80 6.2e-15
             -0.01817
                        0.00886
                                   -2.05
                                          0.0404
(Dispersion parameter for binomial family taken to be 1)
    Null deviance: 608.93 on 443 degrees of freedom
Residual deviance: 483.22 on 438 degrees of freedom
AIC: 495.2
Number of Fisher Scoring iterations: 4
```

Figure 13.3 R output of the summary method for the resp_glm model.

13.3.3 Epilepsy

Moving on to the count data in epilepsy from Table ??, we begin by calculating the means and variances of the number of seizures for all interactions between treatment and period:

```
R> data("epilepsy", package = "HSAUR2")
R> itp <- interaction(epilepsy$treatment, epilepsy$period)</pre>
R> tapply(epilepsy$seizure.rate, itp, mean)
  placebo.1 Progabide.1
                         placebo.2 Progabide.2
                                                  placebo.3
       9.36
              8.58
                               8.29
                                         8.42
                                                       8.79
             placebo.4 Progabide.4
Progabide.3
                   7.96
       8.13
R> tapply(epilepsy$seizure.rate, itp, var)
  placebo.1 Progabide.1
                          placebo.2 Progabide.2
                                                  placebo.3
                          66.7
      102.8
                332.7
                                      140.7
                                                     215.3
Progabide.3
             placebo.4 Progabide.4
      193.0
                   58.2
```

```
R> summary(resp_gee1)
Model:
Tink:
                          Logit
 Variance to Mean Relation: Binomial
 Correlation Structure:
                          Independent
Coefficients:
           Estimate Naive S.E. Naive z Robust S.E. Robust z
(Intercept) -0.9002 0.33765 -2.666 0.460
                                                   -1.956
                               2.803
             0.6716
                       0.23957
                                            0.357
                                                    1.882
centre2
                       0.23684
                               5.486
                                            0.351
                                                     3.704
              1.2992
trttrt
                       0.29467
                               0.405
                                            0.443
             0.1192
                                                     0.269
gendermale
                                            0.350
baselinegood 1.8820
                       0.24129
                                7.800
                                                     5.376
             -0.0182
                       0.00886 -2.049
                                            0.013
                                                    -1.397
Estimated Scale Parameter: 1
```

Figure 13.4 R output of the summary method for the resp_gee1 model (slightly abbreviated).

Some of the variances are considerably larger than the corresponding means, which for a Poisson variable may suggest that overdispersion may be a problem, see Chapter 7.

We can now fit a Poisson regression model to the data assuming independence using the ${\tt glm}$ function. We also use the GEE approach to fit an independence structure, followed by an exchangeable structure using the following R code:

```
R> per <- rep(log(2),nrow(epilepsy))
R> epilepsy$period <- as.numeric(epilepsy$period)
R> names(epilepsy) [names(epilepsy) == "treatment"] <- "trt"
R> fm <- seizure.rate ~ base + age + trt + offset(per)
R> epilepsy_glm <- glm(fm, data = epilepsy, family = "poisson")
R> epilepsy_gee1 <- gee(fm, data = epilepsy, family = "poisson", + id = subject, corstr = "independence", scale.fix = TRUE, + scale.value = 1)
R> epilepsy_gee2 <- gee(fm, data = epilepsy, family = "poisson", + id = subject, corstr = "exchangeable", scale.fix = TRUE, + scale.value = 1)
R> epilepsy_gee3 <- gee(fm, data = epilepsy, family = "poisson", + id = subject, corstr = "exchangeable", scale.fix = FALSE, + scale.value = 1)</pre>
```

```
R> summary(resp_gee2)
Model:
Link:
                           Logit
 Variance to Mean Relation: Binomial
 Correlation Structure:
                         Exchangeable
Coefficients:
            Estimate Naive S.E. Naive z Robust S.E. Robust z
(Intercept) -0.9002
                     0.4785 -1.881
                                        0.460
                                                    -1.956
              0.6716
                         0.3395 1.978
                                             0.357
                                                     1.882
centre2
              1.2992
                         0.3356 3.871
                                             0.351
trttrt
                                                      3.704
                         0.4176 0.286
             0.1192
                                             0.443
                                                      0.269
gendermale
baselinegood 1.8820
                         0.3419
                                 5.504
                                             0.350
                                                      5.376
             -0.0182
                         0.0126 -1.446
                                             0.013
                                                     -1.397
Estimated Scale Parameter: 1
```

Figure 13.5 R output of the summary method for the resp_gee2 model (slightly abbreviated).

As usual we inspect the fitted models using the summary method, the results are given in Figures 13.8, 13.9, 13.10, and 13.11.

13.4 Analysis Using R: Random Effects

As an example of using generalised mixed models for the analysis of longitudinal data with a non-normal response, the following logistic model will be fitted to the respiratory illness data

```
logit(P(status = good)) = \beta_0 + \beta_1treatment + \beta_2time + \beta_3gender
+ \beta_4age + \beta_5centre + \beta_6baseline + u
```

where u is a subject-specific random effect.

The necessary R code for fitting the model using the glmer function from package lme4 (Bates and Sarkar, 2012, Bates, 2005) is:

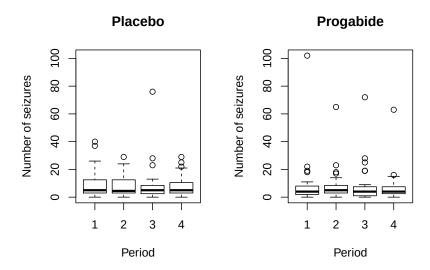


Figure 13.6 Boxplots of numbers of seizures in each two-week period post randomisation for placebo and active treatments.

month.C	trttrt	gendermale	age
0.701	8.725	1.269	0.975
centre2			
2.825			

The significance of the effects as estimated by this random effects model and by the GEE model described in Section 13.3.2 is generally similar. But as expected from our previous discussion the estimated coefficients are substantially larger. While the estimated effect of treatment on a randomly sampled individual, given the set of observed covariates, is estimated by the marginal model using GEE to increase the log-odds of being disease free by 1.299, the corresponding estimate from the random effects model is 2.166. These are not inconsistent results but reflect the fact that the models are estimating

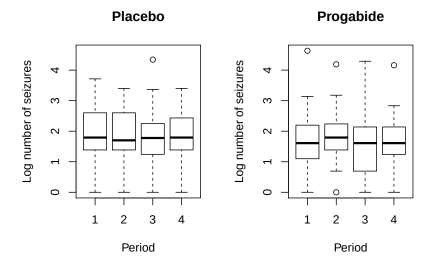


Figure 13.7 Boxplots of log of numbers of seizures in each two-week period post randomisation for placebo and active treatments.

different parameters. The random effects estimate is conditional upon the patient<U+2019>s random effect, a quantity that is rarely known in practise. Were we to examine the log-odds of the average predicted probabilities with and without treatment (averaged over the random effects) this would give an estimate comparable to that estimated within the marginal model.

```
R> summary(epilepsy_glm)
Call:
glm(formula = fm, family = "poisson", data = epilepsy)
Deviance Residuals:
  Min
          1Q Median
                            3Q
                                  Max
                        0.484
-4.436 -1.403 -0.503
                               12.322
Coefficients:
             Estimate Std. Error z value Pr(>|z|)
            -0.130616
                        0.135619
                                   -0.96
                                          0.3355
(Intercept)
              0.022652
                        0.000509
base
                                   44.48
                                          < 2e-16
             0.022740
                        0.004024
                                    5.65 1.6e-08
age
trtProgabide -0.152701
                         0.047805
                                   -3.19
                                           0.0014
(Dispersion parameter for poisson family taken to be 1)
    Null deviance: 2521.75 on 235 degrees of freedom
Residual deviance: 958.46 on 232 degrees of freedom
AIC: 1732
Number of Fisher Scoring iterations: 5
```

Figure 13.8 R output of the summary method for the epilepsy_glm model.

```
R> summary(epilepsy_gee1)
Model:
Link:
                            Logarithm
 Variance to Mean Relation: Poisson
 Correlation Structure:
                            Independent
Coefficients:
             Estimate Naive S.E. Naive z Robust S.E. Robust z
              -0.1306
                        0.135619 -0.963
                                             0.36515
                                                       -0.358
(Intercept)
                        0.000509 44.476
                                             0.00124
                                                       18.332
base
               0.0227
               0.0227
                        0.004024
                                  5.651
                                             0.01158
                                                        1.964
trtProgabide -0.1527
                        0.047805 -3.194
                                             0.17111
                                                       -0.892
Estimated Scale Parameter: 1
```

Figure 13.9 R output of the summary method for the epilepsy_gee1 model (slightly abbreviated).

R> summary(epilepsy_gee2) Model: Link: Logarithm Variance to Mean Relation: Poisson Correlation Structure: Exchangeable Coefficients: Estimate Naive S.E. Naive z Robust S.E. Robust z 0.200442 -0.652 -0.1306 0.36515 -0.358(Intercept) 18.332 0.0227 0.000753 30.093 0.00124 base 0.005947 3.824 0.01158 0.0227 1.964 age trtProgabide -0.1527 0.070655 -2.161 0.17111 -0.892 Estimated Scale Parameter: 1

Figure 13.10 R output of the summary method for the epilepsy_gee2 model (slightly abbreviated).

```
R> summary(epilepsy_gee3)
. . .
Model:
Link:
                           Logarithm
Variance to Mean Relation: Poisson
 Correlation Structure: Exchangeable
. . .
Coefficients:
            Estimate Naive S.E. Naive z Robust S.E. Robust z
(Intercept)
             -0.1306 0.4522 -0.289 0.36515 -0.358
                         0.0017 13.339
base
              0.0227
                                           0.00124
                                                     18.332
                                1.695
age
              0.0227
                         0.0134
                                           0.01158
                                                     1.964
                        0.1594 -0.958
trtProgabide -0.1527
                                           0.17111
                                                     -0.892
Estimated Scale Parameter: 5.09
```

Figure 13.11 R output of the summary method for the epilepsy_gee3 model (slightly abbreviated).

R> summary(re	R> summary(resp_lmer)			
Fixed effect	Fixed effects:			
	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	-1.6546	0.7762	-2.13	0.033
baselinegood	3.0890	0.5986	5.16	2.5e-07
month.L	-0.2035	0.2796	-0.73	0.467
month.Q	-0.0282	0.2791	-0.10	0.919
month.C	-0.3557	0.2808	-1.27	0.205
trttrt	2.1662	0.5516	3.93	8.6e-05
gendermale	0.2384	0.6661	0.36	0.720
age	-0.0256	0.0199	-1.28	0.200
centre2	1.0385	0.5418	1.92	0.055

Figure 13.12 R output of the summary method for the resp_lmer model (abbreviated).

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Bibliography

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