STAT/BIOST 571: Homework 6

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Problem 1: Fitting and interpreting the results of a linear mixed effects model; robust standard error estimation (20 points)

Download the creatinine.csv dataset from the course website. This file contains repeated observational data for 619 subjects, some of whom have hypertension and some of whom have a hereditary kidney disease, as indicated by the group variable, according to the coding in the Table 1. The outcome variable

Group	Kidney disease	Hypertension	Sample size
1	Yes	Yes	294
2	Yes	No	103
3	No	Yes	73
4	No	No	149

Table 1: Measurements of serum creatinne reciprocals from 619 subjects in four groups

is scr, the reciprocal of serum creatinine. Serum creatinine is a measure of kidney function, with lower values indicating better kidney function. Higher values of the reciprocal reported in scr indicate better kidney function. The observations were taken at arbitrary times from each subject, with the number of observations ranging from 1 to 22. Ignoring hypertension status, we are interested in estimating the rate of change of scr for subjects with and without hereditary kidney disease. Thus, the only fixed effect covariates in your model should be age, kidney disease status, and possibly an interaction between these. In order to account for correlation within subjects, you will be fitting a linear mixed effects model with uncorrelated random slopes and intercepts and serial correlation of residuals that follows a spherical correlation model, including a nugget (correlation should be based on the timing of observations). Please use the lme() function in the nlme package in R to fit your models (i.e., do not code your own nonlinear optimization).

(a) Fit the model by ML and report the estimated values of all variance parameters.

The parameter estimates for fitting the model without and with an interaction term can be seen in Tables 3 and 4. The variance parameters can be found in Table 2. Their meaning is detailed in the subsequent paragraphs. The R model summaries can be found in the Appendix.

	Model				
Parameter	Without interaction (Equation 1)	With interaction (Equation 2)			
$\hat{\sigma}$	0.2633414	0.225834			
$\hat{\sigma}_{\gamma_0}$	0.04643211	0.1317468			
$\hat{\sigma}_{\gamma_1}$	0.00522239	0.004823202			
\hat{lpha}_r	7.8894707	4.6700641			
\hat{lpha}_n	0.1759323	0.2299764			

Table 2: Variance parameters for ML-fitted models.

Let t_{ij} be the age of the subject i at observation j. Let x_i indicate whether the subject has has kidney disease. Without an interaction term, the mean model is

$$Y_{ij} = (\beta_0 + \gamma_0) + \beta_2 x_i + (\beta_1 + \gamma_1) t_{ij} + \epsilon_{ij}.$$
 (1)

With the interaction term, the mean model is

$$Y_{ij} = (\beta_0 + \gamma_0) + \beta_2 x_i + (\beta_1 + \beta_3 x_i + \gamma_1) t_{ij} + \epsilon_{ij}.$$
 (2)

 γ_j are the random effects, where γ_0 is subject-specific adjustment to the intercept, and γ_1 is the subject-specific adjustment to the slope.

The covariance structure of subject i can be described by the matrix

$$\Sigma_i = \sigma^2 \left(Z_i G Z_i^{\mathsf{T}} + R_i \right) \tag{3}$$

 Z_i is a $m_i \times 2$ matrix, where the first column entries are all 1s, and the second column entries are ages for each subject t_{ij} .

G is a 2×2 diagonal matrix that describes the variance of the random effects γ_0 and γ_1 :

$$G = \frac{1}{\sigma^2} \begin{pmatrix} \sigma_{\gamma_0}^2 & 0\\ 0 & \sigma_{\gamma_1}^2 \end{pmatrix} \tag{4}$$

 R_i is an $m_i \times m_i$ matrix that describes the correlations between the ϵ_{ij} s for different js with a nugget parameter $0 \le \alpha_n < 1$ and range parameter $\alpha_r > 0$. $R_{ijj} = 1$ and $R_{ijj'} = (1 - \alpha_n) \exp\left(-\frac{|t_{ij} - t_{ij'}|}{\alpha_r}\right)$. Estimates for these parameters can be found in Table 2.

(b) Report point estimates and standard errors for all fixed effect coefficients in your model. Include three versions of standard error estimates: (i) robust/empirical sandwich SEs that correctly account for clustering of the data, (ii) bootstrap SEs that correctly account for clustering of the data, and (iii) model based SE estimates based on the assumed random effect model being correct.

Solution: See Table 3 for the point estimates and standard errors when no interaction term is included.

	Estimate	ML Standard Error	Sandwich Standard Error	Bootstrap Standard Error
(Intercept)	1.532222	0.039557	0.039348	0.041531
age	-0.012915	0.000964	0.000941	0.001030
kidney.disease	-0.281916	0.026260	0.025027	0.034516

Table 3: Standard error estimates for fixed effect parameters.

Let \hat{V}_i be the result of substituting the variance parameter estimates in Table 2 into Equation 3. Let $W_i = \hat{V}_i^{-1}$ be the weight matrix for each cluster. Let X_i be the matrix of cluster covariates, 1s in the first column, age in the second column, and an indicator for kidney disease in the third column. Let Y_i be the cluster response.

If we assume the random effect model is correct, then the covariance matrix for the paramter estimates $\hat{\beta}$ is

$$var\left(\hat{\beta}\right) = \left(\sum_{i=1}^{n} X_i^{\mathsf{T}} W_i X_i\right)^{-1},\tag{5}$$

and we take the square root of the diagonals to obtain the standard errors for the ML Standard Error column.

For the sandwich standard errors, we use the covariance matrix

$$\hat{\text{var}}\left(\hat{\beta}\right) = \left(\sum_{i=1}^{n} X_{i}^{\mathsf{T}} W_{i} X_{i}\right)^{-1} \left(\sum_{i=1}^{n} X_{i}^{\mathsf{T}} W_{i} \left(Y_{i} - X_{i} \hat{\beta}\right) \left(Y_{i} - X_{i} \hat{\beta}\right)^{\mathsf{T}} W_{i} X_{i}\right) \left(\sum_{i=1}^{n} X_{i}^{\mathsf{T}} W_{i} X_{i}\right)^{-1}, \quad (6)$$

which we use to get the Sandwich Standard Error column.

For the bootstrap standard errors, we resample clusters, and then fit a model to the resampled clusters to get samples $\hat{\beta}^{(1)}, \hat{\beta}^{(2)}, \dots, \hat{\beta}^{(L)}$. In this case, $L = 2^{10}$. Then, $\hat{\beta}^{(L)}$ is estimated by taking the unbiased covariance estimate of the samples. Taking the square root of the diagonals gives us the Bootstrap Standard Error column.

In this case, the standard errors assuming the random effects model is correct and the sandwich standard errors are very similar with the sandwich standard errors being slightly smaller. The bootstrap standard errors are largest. The difference is more than just Monte Carlo error, particularly for β_2 the coefficient for kidney disease, which hints at its interaction with other covariates.

(c) Now, give point estimates and three versions of standard error estimates for the marginal rates of change in scr in subjects with and without kidney disease. As in part (b), your three versions of SE estimates should be: (i) robust/empirical sandwich SEs that correctly account for clustering of the data, (ii) bootstrap SEs that correctly account for clustering of the data, and (iii) model based SE estimates based on the assumed random effect model being correct.

Solution: See Table 4 for the estimates and the standard errors for the model that includes the interaction term.

For subjects without kidney disease, we expect that the observed scr decreases by -0.003108 for each additional year of age. This effect is barely statistically significant: when using the

	Estimate	ML Standard Error	Sandwich Standard Error	Bootstrap Standard Error
(Intercept)	1.190676	0.054232	0.050269	0.050770
age	-0.003108	0.001436	0.001194	0.001217
kidney.disease	0.313575	0.071373	0.068325	0.070047
age:kidney.disease	-0.016649	0.001857	0.001705	0.001706

Table 4: Standard error estimates for fixed effect parameters with interaction term.

standard error that assumes the random effects model is correct, the upper bound of the 97.5% confidence interval is greater than 0.

For subjects with kidney disease, the expected observed baseline scr is 0.313575 higher. The expected change in observed scr for each additional year of age differs by -0.016649, so the expected observed scr decreases by -0.019757 for each additional year of age for subjects with kidney disease. These effects are all statistically significant. Indeed, we would expect that subjects with kidney disease experience a sharper decline in kidney function with age.

Standard errors that assume the random effect model is correct and sandwich standard errors are calculated with Equations 5 and 6 as in the previous part. To model the interation term, X_i has an additional column of $x_i t_{ij}$ s. The same bootstrap procedure of resampling clusters is used.

In this case, the bootstrap standard errors are almost identical to the sandwich standard errors. Both are smaller than the standard errors that assume the random effects model is correct. This indicates that there is some correlation between the random effects that can be leveraged to get estimates with less variance.

Appendix

Code for fitting models and generating tables is attached on the subsequent pages.

LME and Creatinine

Joining Data

```
In [1]: library(data.table)
    library(nlme)
    library(parallel)
    library(xtable)

    creatinine.data <- data.table(read.csv('creat.csv'), key='group')
    head(creatinine.data)</pre>
```

```
        id
        group
        age
        scr

        1
        1
        35.765
        0.182

        1
        1
        37.990
        0.088

        3
        1
        51.083
        0.156

        3
        1
        52.386
        0.116

        3
        1
        52.805
        0.087

        3
        1
        52.997
        0.067
```

group	kidney.disease	hypertension
1	1	1
2	1	0
3	0	1
4	0	0

```
In [3]: creatinine.group.data <- creatinine.data[group.data]
    setkey(creatinine.group.data, id)
    head(creatinine.group.data)</pre>
```

id	group	age	scr	kidney.disease	hypertension
1	1	35.765	0.182	1	1
1	1	37.990	0.088	1	1
2	2	24.997	1.429	1	0
2	2	27.441	1.111	1	0
2	2	30.524	1.429	1	0
3	1	51.083	0.156	1	1

Fitting LME Model

```
In [4]: fit.scr.model <- function(data, interaction.term=FALSE) {
    formula = scr ~ age + kidney.disease
        if (interaction.term) { formula <- update(formula, . ~ . + age:kidney.disease)
    }
    lme(formula,
        random=reStruct(~age|id, pdClass='pdDiag'),
        correlation=corExp(form=~age|id, nugget=TRUE),
        method='ML', data=data,
        control=lmeControl(maxIter=100, msMaxIter=100, niterEM=50))
}</pre>
```

Covariance for Subject i

```
In [5]: make.covariance <- function(model, i) {
    # Error term, usually denoted epsilon
    error.correlation <- corMatrix(model$model$truct$corStruct)[[as.character(i)]]
    if (is.null(error.correlation)) { error.correlation <- 1 }
    # Random effects correlation, Z * G * tranpose(Z).
    Z <- cbind(1, model$data[J(i), age])
    random.correlation <- Z %*% as.matrix(model$model$truct$reStruct$id) %*% t(Z)
    # Convert correlation matrix into covariance matrix.
    (random.correlation + error.correlation)*(model$sigma*model$sigma)
}</pre>
```

β_i Covariance

Maximum Likelihood Estimate

This assumes that the random effects model is correct. I can also be used as the bread part of the sandwich estimator.

```
In [6]: make.covariates <- function(model, data) model.matrix(model$terms, data)

make.ml.parameter.covariance <- function(model) {
    groups <- unique(model$groups)$id
    chol2inv(chol(Reduce(`+`, lapply(groups, function(i) {
        X <- make.covariates(model, model$data[J(i)])
        t(X) %*% chol2inv(chol(make.covariance(model, i))) %*% X
    }))))
}</pre>
```

Sandwich Estimate

```
In [7]: make.response <- function(data) data$scr

make.sandwich.parameter.covariance <- function(model) {
    bread <- make.ml.parameter.covariance(model)
    meat <- Reduce(`+`, lapply(unique(model$groups)$id, function(i) {
        X <- make.covariates(model, model$data[J(i)])
        y <- make.response(model$data[J(i)])
        weights <- chol2inv(chol(make.covariance(model, i)))
        residuals <- as.numeric(make.response(model$data[J(i)]) - X %*% model$coef
    ficients$fixed)
        empirical.covariance <- outer(residuals, residuals)
        t(X) %*% weights %*% empirical.covariance %*% weights %*% X
    }))
    bread %*% meat %*% bread
}</pre>
```

Bootstrap Estimate

To account for clustering of the data, we resample clusters.

Models

Without Interaction Term

```
In [9]: | scr.model <- fit.scr.model(creatinine.group.data)</pre>
         summary(scr.model, adjustSigma=FALSE)
         Linear mixed-effects model fit by maximum likelihood
          Data: data
                 AIC
                          BIC
                                 logLik
           -53.94986 -11.00314 34.97493
         Random effects:
          Formula: ~age | id
          Structure: Diagonal
                                   age Residual
                 (Intercept)
         StdDev: 0.04643211 0.00522239 0.2633414
         Correlation Structure: Exponential spatial correlation
          Formula: ~age | id
          Parameter estimate(s):
             range
                     nugget
         7.8894707 0.1759323
         Fixed effects: list(formula)
                             Value Std.Error DF t-value p-value
         (Intercept)
                        1.5322224 0.03955663 965 38.73490
                        -0.0129154 0.00096383 965 -13.40006
         kidney.disease -0.2819162 0.02626024 617 -10.73548
          Correlation:
                        (Intr) age
                        -0.849
         age
         kidney.disease -0.352 -0.082
         Standardized Within-Group Residuals:
                 Min
                              Q1
                                        Med
                                                                  Max
                                                       Q3
         -2.49081741 -0.56731338 -0.07887231 0.43113427 4.95897388
         Number of Observations: 1585
         Number of Groups: 619
In [10]: | ml.parameter.covariance <- make.ml.parameter.covariance(scr.model)</pre>
         sandwich.parameter.covariance <- make.sandwich.parameter.covariance(scr.model)</pre>
         bootstrap.parameter.samples <- do.call(rbind, mclapply(</pre>
             replicate(1024, creatinine.group.data, simplify=FALSE), function(data) {
                 resampled.data <- resample.clusters(data)</pre>
                 fit.scr.model(resampled.data)$coefficients$fixed
             }, mc.cores=4))
         bootstrap.parameter.covariance <- cov(bootstrap.parameter.samples)</pre>
```

	Estimate	ML Standard Error	Sandwich Standard Error	Bootstrap Standard Error
(Intercept)	1.53222240	0.0395566329	0.0393479582	0.041530790
age	-0.01291536	0.0009638288	0.0009409584	0.001030028
kidney.disease	-0.28191623	0.0262602392	0.0250273809	0.034516098

With Interaction Term

```
In [12]:
         scr.model.interaction <- fit.scr.model(creatinine.group.data, interaction.term=TRU</pre>
         summary(scr.model.interaction, adjustSigma=FALSE)
         Linear mixed-effects model fit by maximum likelihood
          Data: data
                 AIC
                           BIC
                                 logLik
           -127.9923 -79.67728 72.99617
         Random effects:
          Formula: ~age | id
          Structure: Diagonal
                 (Intercept)
                                     age Residual
                   0.1317468 0.004823202 0.225834
         StdDev:
         Correlation Structure: Exponential spatial correlation
          Formula: ~age | id
          Parameter estimate(s):
             range
                     nugget
         4.6700641 0.2299764
         Fixed effects: list(formula)
                                 Value Std.Error DF t-value p-value
                             1.1906763 0.05423200 964 21.955237 0.0000
         (Intercept)
                            -0.0031083 0.00143568 964 -2.165016 0.0306
         age
                             0.3135748 0.07137345 617 4.393438 0.0000
         kidney.disease
         age:kidney.disease -0.0166492 0.00185709 964 -8.965190 0.0000
          Correlation:
                            (Intr) age
                                          kdny.d
         age
                            -0.928
         kidney.disease
                            -0.760 0.705
         age:kidney.disease 0.718 -0.773 -0.935
         Standardized Within-Group Residuals:
                              Q1
                                         Med
                                                      Q3
         -2.30553348 -0.53490304 -0.07976542 0.36022352 5.46804808
         Number of Observations: 1585
         Number of Groups: 619
In [13]: ml.parameter.covariance.interaction <- make.ml.parameter.covariance(scr.model.inte
         raction)
         sandwich.parameter.covariance.interaction <- make.sandwich.parameter.covariance(sc</pre>
         r.model.interaction)
         bootstrap.parameter.samples.interaction <- do.call(rbind, mclapply(</pre>
             replicate(1024, creatinine.group.data, simplify=FALSE), function(data) {
                 resampled.data <- resample.clusters(data)</pre>
                 fit.scr.model(resampled.data, interaction.term=TRUE)$coefficients$fixed
             }, mc.cores=4))
         bootstrap.parameter.covariance.interaction <- cov(bootstrap.parameter.samples.inte
         raction)
```

```
In [14]: standard.errors.interaction <- data.frame(</pre>
              `Estimate`=scr.model.interaction$coefficients$fixed,
              `ML Standard Error`=sqrt(diag(ml.parameter.covariance.interaction)),
              `Sandwich Standard Error`=sqrt(diag(sandwich.parameter.covariance.interaction
              \verb|`Bootstrap Standard Error` = & \texttt{sqrt}( \\ \texttt{diag}( \texttt{bootstrap.parameter.covariance.interactio})|
          n)),
              check.names=FALSE)
          print(xtable(standard.errors.interaction,
                        caption='Standard error estimates for fixed effect parameters with in
          teraction term.',
                        label='tab:standard_errors_interaction',
                        digits=c(0, 6, 6, 6, 6)),
                booktabs=TRUE, file='standard_errors_interaction.tex',
                sanitize.colnames.function=identity,
                sanitize.rownames.function=identity,
                size='small')
          standard.errors.interaction
```

	Estimate	ML Standard Error	Sandwich Standard Error	Bootstrap Standard Error
(Intercept)	1.190676306	0.054231995	0.050269334	0.050769991
age	-0.003108265	0.001435677	0.001193725	0.001216667
kidney.disease	0.313574816	0.071373449	0.068324860	0.070047199
age:kidney.disease	-0.016649153	0.001857089	0.001705009	0.001706189