STAT 571 HW7

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1 Relationship between fixed effects, random effects, GLS, and penalized regression; confounding; model misspecication

1.1 Part (a)

Figure 1 shows the marginal relationship between literacy and age regardless of individual. We connect the points of each individual.

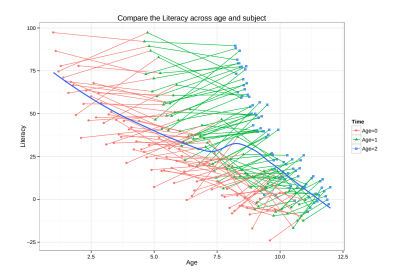


Figure 1 – The relationship between bias and the σ^2 . The blue lowess curve shows the marginal trend of literacy v.s. age.

Figure 2 shows the time correlation within each individual.

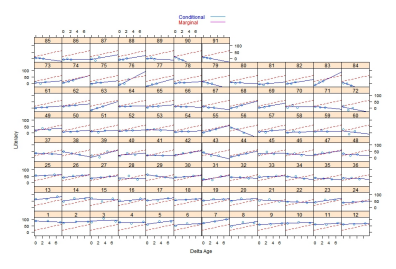


Figure 2 – Scatterplot for each individual. The within individual correlation over time is obvious.

1.2 Part (b and c)

We will compare the following two models: the fixed effect model (subject-specific model, i.e. there is an intercept β_i for each i),

$$Y_{ij} = (\beta_i) + \beta_L (x_{i1} - x_{ij}) + \epsilon_{ij}.$$

 $\epsilon_{ij} \stackrel{iid}{\sim} \text{some distribution with } (0, \sigma^2)$

and the linear mixed model

$$Y_{ij} = (\beta_0 + \gamma_i) + \beta_L (x_{i1} - x_{ij}) + \epsilon_{ij}$$

$$\gamma_i \stackrel{iid}{\sim} N(0, \sigma_{\gamma}^2)$$

$$\epsilon_{ij} \stackrel{iid}{\sim} N(0, \sigma^2)$$

The subject-specific model is better. Random effect model assumes that the random intercept is i.i.d. from some distribution. However, in this case since the data of X is fixed, the intercept for each subject is fixed, so random intercept model should not be used.

We simulated the data by the generating mechanism 5000 times. In each simulation, we fitted both models and recorded the estimates of β_L . The linear mixed model was done with REML.

1.3 Part (d)

Compare of the beta.L estimates in different models

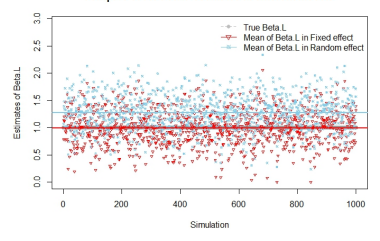


Figure 3 – The relationship between bias and the σ^2

In Figure 3 we see that the mean of $\hat{\beta}_L$ in the fixed effect model(subject-specific model) is closer to the true β_L .

Table 1 shows the Bias and standard error of of $\hat{\beta}_L$ in both models. The $\hat{\beta}_L$ in the fixed effect model seems to be unbiased, while the $\hat{\beta}_L$ in the random effect model does not.

Model	Bias	Standard deviation
subject-specific model	-1.27E-03	0.317
linear mixed model	-2.65E-01	0.322

Table 1 – Bias and standard error of \hat{eta}_L .

1.4 Part (e)

Table 2 shows the variables in the model. Both of the estimates of σ^2 are unbiased.

Model	Expected variance σ^2	Expected variance σ_{γ}^2
subject-specific model	99.94	-
linear mixed model	100.34	594.54

Table 2 - Estimated variance.

1.5 Part (f)

Calculate the "true" variance of $f(\cdot)$ function, given the "data" of the population.

$$Y_{ij} = (\beta_0 + \gamma_i) + \beta_L (x_{i1} - x_{ij}) + \epsilon_{ij}$$

$$EY_{ij} = (\beta_0 + \gamma_i) + \beta_L (x_{i1} - x_{ij})$$

$$\mu_{\gamma} = E[f(x_{i1})]$$

$$\sigma_{\gamma}^2 = var[f(x_{i1})] = \frac{1}{91} \left[\Sigma_i^{91} (f(x_{i1}) - \mu_{\gamma}) \right]^2$$

$$= 596.97$$

1.6 Part (g)

There are two ways to do. One is using the GLS regression and the other is penalized regression. From the below random effect model as asked, the only thing changed is the *exact* bias of $\hat{\beta}_L$,

$$Y_{ij} = (\beta_0 + \gamma_i) + \beta_L (x_{i1} - x_{ij}) + \epsilon_{ij}.$$

• GLS

$$\begin{split} \hat{\beta}_{L} &= \left(X^{T}WX\right)^{-1}\left(X^{T}WY\right) \\ W_{i} &= V_{i}^{-1} \\ var(Y_{ij}) &= cov\left(\gamma_{i} + \epsilon_{ij}, \gamma_{i} + \epsilon_{ij}\right) = \sigma^{2} + \sigma_{\gamma}^{2} \\ cov(Y_{ij}, Y_{ij'}) &= cov\left(\gamma_{i} + \epsilon_{ij'}, \gamma_{i} + \epsilon_{ij'}\right) = \sigma_{\gamma}^{2} \\ \Sigma_{i} &= \begin{bmatrix} \sigma^{2} + \sigma_{\gamma}^{2} & \sigma_{\gamma}^{2} & \sigma_{\gamma}^{2} \\ \sigma_{\gamma}^{2} & \sigma^{2} + \sigma_{\gamma}^{2} & \sigma_{\gamma}^{2} \\ \sigma_{\gamma}^{2} & \sigma^{2}^{2} + \sigma_{\gamma}^{2} \end{bmatrix} \\ &= \left(\sigma^{2} + \sigma_{\gamma}^{2}\right) \begin{bmatrix} 1 & \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} & \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} \\ \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} & 1 & \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} \end{bmatrix} \\ V_{i} &= \begin{bmatrix} 1 & \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} & \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} & 1 \end{bmatrix} \\ V_{i} &= \begin{bmatrix} 1 & \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} & \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} \\ \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} & 1 & \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} \end{bmatrix} \end{split}$$

The exact bias is

$$E(\hat{\beta}_L) - \beta_L = \left(X^T W X\right)^{-1} \left(X^T W E(Y)\right) - \beta_L$$

$$= \left(\Sigma_i X_i^T W_i X_i\right)^{-1} \left(\Sigma_i X_i^T W_i E(Y_i)\right) - \beta_L,$$

$$= 1.273 - 1$$

$$= 0.273$$

where

$$E(Y) - \beta_L = (\beta_0 + \gamma_i) + \beta_L (x_{i1} - x_{ij}).$$

Penalized regression
 From the lecture notes, we have the solution from the penalized regression,

$$\begin{bmatrix} \beta \\ \gamma \end{bmatrix} = \arg \min \Sigma_{i}^{n} \left\{ (Y_{i} - X_{i}\beta - 1\gamma_{i})^{T} (Y_{i} - X_{i}\beta - 1\gamma_{i}) + \frac{\sigma^{2}}{\sigma_{\gamma}^{2}} \gamma_{i}^{2} \right\}$$

$$E \begin{bmatrix} \beta \\ \gamma \end{bmatrix} = \arg \min \Sigma_{i}^{n} \left\{ (EY_{i} - X_{i}\beta - 1\gamma_{i})^{T} (EY_{i} - X_{i}\beta - 1\gamma_{i}) + \frac{\sigma^{2}}{\sigma_{\gamma}^{2}} \gamma_{i}^{2} \right\}$$

- As in GLS case, we plug the mean of Y into the penalized package. The output is 0.27288.

1.7 Part (h)

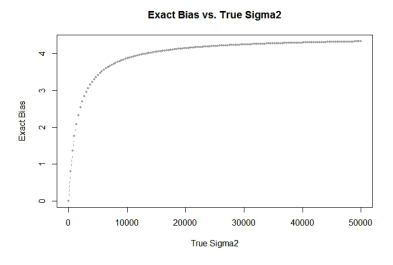


Figure 4 – The relationship between bias and the σ^2

Figure 4 shows that the exact bias is increasing as the σ^2 increases. But the slope is first decreasing.

1.8 Part (i)

$$Y_{ij} = (\beta_i) + \beta_L (x_{i1} - x_{ij}) + \epsilon_{ij}$$
 (1)

$$Y_{ij} = (\beta_0 + \gamma_i) + \beta_L (x_{i1} - x_{ij}) + \epsilon_{ij}$$
 (2)

$$Y_{ij} = (\beta_0) + \beta_L (x_{i1} - x_{ij}) + \epsilon_{ij}$$
(3)

$$E\begin{bmatrix} \beta \\ \gamma \end{bmatrix} = \arg \min \Sigma_i^n \left\{ (EY_i - X_i\beta - 1\gamma_i)^T (EY_i - X_i\beta - 1\gamma_i) + \frac{\sigma^2}{\sigma_\gamma^2} \gamma_i^2 \right\}$$
(4)

$$V_{i} = \begin{bmatrix} 1 & \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} & \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} \\ \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} & 1 & \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} \\ \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} & \frac{\sigma_{\gamma}^{2}}{\sigma^{2} + \sigma_{\gamma}^{2}} & 1 \end{bmatrix}$$

$$(5)$$

Table 3 shows the comparison of the GLS and the penalized regression in explaining for the patterns in Figure 4. When $\sigma^2 = 0$, the estimates of β is unbiased and thus the bias is zero. If σ^2 increases from zero, the bias increases and converges to some value.

Case	Explanation			
$\sigma^2 \approx 0$	$\frac{\sigma^2}{\sigma_{\gamma}^2}$ is too small and Eq. (4) is a unconstraint problem for a subject-specific fixed effect model of Eq. (1).			
$\sigma^2 \approx 0$	In GLS, Eq. (5) would make the all the within-cluster observations highly dependent.			
$\sigma^2 \approx +\infty$	$\frac{\sigma^2}{\sigma_{\gamma}^2}$ is too large and Eq. (4) would make the $\gamma_i=0$. This is the OLS model of Eq. (3).			
$\sigma^2 \approx +\infty$	In GLS, Eq. (5) would make the all the within-cluster observations independent. This is the OLS model of Eq. (3).			

Table 3 - Summary of the GLS and the penalized regression

1.9 Part (j)

The GLS gives unbiased estimate of β with any working covariance matrix. In this case, the working covariance is the true covariance by our derivation. Thus the model of GLS is better than the OLS due to a correct covariance matrix.

However, the GLS does not correctly specify the baseline age. So, the GLS does not work well, as the linear mixed model does, even the GLS returns an unbiased estimate. The OLS, as we have discussed, is wrong at both mean model and covariance matrix, so it work worse than the GLS.

2 Fitting and interpreting the results of a linear mixed effects model; deriving the REML and ML likelihood functions; robust standard error estimation

2.1 Part (a)

Table 4 shows the variables in the model.

Variable name	Explanation
scr	The response
kidney	Kidney = 1 means the subject with hereditary disease.
age	Age of the subject
id	Identification number for subject

Table 4 - Summary of the variables in the model

We denote jth ($j = 1, ..., m_i$) observation of the ith (i = 1, ..., 619) subject (id) as Y_{ij} . We fit the following model of scr on age, kidney disease status, and their interaction. Since we are interested to estimate the rate of change in serum creatinine scr for subjects with and without hereditary kidney disease, then the random effects are on age. We include an random slope and an random intercept to account for correlation within subject. The variable kidney is a indicator, and the value equal to 1 means that the subject has kidney disease. In our data, the kidney disease status did not change within subject.

$$\begin{array}{rcl} Y_{ij} & = & \beta_0 + \beta_1 \cdot age_{ij} + \beta_2 \cdot kidney_i + \beta_{12} \cdot age_{ij} \times kidney_i \\ & & + \gamma_{i0} + \gamma_{i1} \cdot age_{ij} + W_i(t_{ij}) + \epsilon_{ij} \end{array}$$

$$\left(\begin{array}{c} \gamma_{i0} \\ \gamma_{i1} \end{array} \right) \stackrel{iid}{\sim} N \left[\left(\begin{array}{c} 0 \\ 0 \end{array} \right), G = \left(\begin{array}{c} G_{11} = \sigma_{\gamma_0}^2 & 0 \\ 0 & G_{22} = \sigma_{\gamma_1}^2 \end{array} \right) \right]$$

We include serial dependence W_i to construct the covariance of ε_i , which is a multivariate Gaussian with zero mean and covariance as below,

$$\begin{array}{rcl} E(\epsilon_{ij}) & = & 0 \\ var(\epsilon_{ij}) & = & \sigma^2 \\ \\ cov(\epsilon_{ij}, \epsilon_{ij'}) & = & \begin{cases} (1-nugget) \cdot \sigma^2 \left[1 - 1.5 \cdot \frac{|age_{ij} - age_{ij'}|}{range} + 0.5 \left(\frac{|age_{ij} - age_{ij'}|}{range} \right)^3 \right] & |age_{ij} - age_{ij'}| < range \\ 0 & |age_{ij} - age_{ij'}| \geqslant range \end{cases} \end{array}$$

where σ^2 is the residual variance, and the *nugget* parameter that measures the portion of σ^2 that belongs to the nugget effect.

Moreover, the matrix notation $\epsilon_i \sim N(0, R_i)$ where R_i is the covariance matrix of ϵ_i given as below,

$$R_{i,jj'} = cov(\epsilon_{ij}, \epsilon_{ij'})$$

 $R_{i,jj} = var(\epsilon_{ij}).$

2.2 Part (b)

Within a subject i, let X_i be the design matrix of fixed effect covariates, and Z_i be the design matrix of random effect covariates, as below

$$X_{i} = \begin{cases} 1 & age_{i1} & kidney_{i} & age_{i1} \times kidney_{i} \\ 1 & age_{i2} & kidney_{i} & age_{i2} \times kidney_{i} \\ \vdots & \vdots & \vdots & \vdots \\ 1 & age_{im} & kidney_{i} & age_{im} \times kidney_{im} \end{cases}$$

$$Z_{i} = \begin{cases} 1 & age_{i1} \\ 1 & age_{i2} \\ \vdots & \vdots \\ 1 & age_{im} \end{cases} .$$

Let $\beta = (\beta_0, \beta_1, \beta_2, \beta_3)^T$ and $\gamma_i = (\gamma_{i0}, \gamma_{i1})^T$. Our data generating model is below,

$$Y_{i} = X_{i}\beta + Z_{i}\gamma_{i} + \epsilon_{i}$$

$$\gamma_{i} \sim N(0, G)$$

$$\epsilon_{i} \sim N(0, R_{i})$$

Or equivalently if we set $\alpha = \left(\sigma^2, nugget, range, \sigma_{\gamma_0}^2, \sigma_{\gamma_1}^2\right)^T$, we have

$$\begin{array}{rcl} Y_i & = & X_i\beta + Z_i\gamma_i + \epsilon_i \\ cov(\gamma_i) & = & G(\alpha) \\ cov(\epsilon_i) & = & R_i(\alpha) \\ cov(Y_i) & = & Z_iG(\alpha)Z_i^T + R_i(\alpha) = \Sigma_i(\alpha) \end{array}$$

The ML log likelihood function from profiling likelihood (3.48) is below,

$$l(\alpha) = -\Sigma_i^n log |\Sigma_i| - \Sigma_i^n (Y_i - X_i \hat{\beta})^T \Sigma_i^{-1} (Y_i - X_i \hat{\beta})$$

$$\hat{\beta} = \beta(\alpha) = \left(\Sigma_i^n X_i^T \Sigma_i^{-1} X_i\right)^{-1} \left(\Sigma_i^n X_i^T \Sigma_i^{-1} Y_i\right).$$
(6)

The REML log likelihood is below,

$$l(\alpha) = -\Sigma_{i}^{n} log \left| \Sigma_{i} \right| - \Sigma_{i}^{n} log \left| X_{i}^{T} \Sigma_{i}^{-1} X_{i} \right| - \Sigma_{i}^{n} \left(Y_{i} - X_{i} \hat{\beta} \right)^{T} \Sigma_{i}^{-1} \left(Y_{i} - X_{i} \hat{\beta} \right)$$

$$\hat{\beta} = \beta(\alpha) = \left(\Sigma_{i}^{n} X_{i}^{T} \Sigma_{i}^{-1} X_{i} \right)^{-1} \left(\Sigma_{i}^{n} X_{i}^{T} \Sigma_{i}^{-1} Y_{i} \right).$$
(7)

To obtain ML and REML $\hat{\alpha}$'s, we maximize Eq. (6) and Eq. (7) respectively.

2.3 Part (c)

The results are shown in the below Table 5. The estimates of $\alpha = \left(\sigma^2, nugget, range, \sigma_{\gamma_0}^2, \sigma_{\gamma_1}^2\right)^T$ via REML are slightly higher than the ones using MLE.

Table 5 - Estimated variance parameters for ML and REML

Linear mixed-effects model fit by maximum likelihood Linear mixed-effects model fit by REML Data: data Log-likelihood: 71.38007 Data: data Log-restricted-likelihood: 53.66149 Fixed: scr ~ age * kidney (Intercept) age kidney age:kidney Fixed: scr ~ age * kidney (Intercept) age kidney age:kidney 1.196490051 -0.003275613 0.314170354 -0.016642243 1.19663113 -0.00327953 0.31431681 -0.01664654 Random effects: Formula: ~1 + age | id Random effects: Formula: ~1 + age | id Structure: Diagonal Structure: Diagonal (Intercept) age Residual (Intercept) age Residual StdDev: 0.1559501 0.004816316 0.2117685 (Sqaure) StdDev: 0.1553242 0.004798277 0.2115859 (Sqaure) Correlation Structure: Spherical spatial correlation Correlation Structure: Spherical spatial correlation Formula: ~age | id Formula: ~age | id Parameter estimate(s): Parameter estimate(s): range nugget range nugget **7.36**76230 **0.27**63843 **7.37**76035 **0.27**58132 Number of Observations: 1585 Number of Groups: 619 Number of Observations: 1585 Number of Groups: 619 > fit.reml\$sigma^2 > fit.ml\$sigma^2 [1] 0.0448 [1] 0.0448

 $\hat{\sigma}^{\overline{2}}$ $\sigma_{\gamma_0}^2$ $\sigma_{\gamma_1}^2$ nuĝget range 0.0448 0.0243 2.319e-05 7.38 REML 0.276 ML 0.0448 0.0241 2.302e-05 0.276 7.37

2.4 Part (d)

Table 6 shows the values of three versions of standard estimates for ML and REML. Values in Table 6a are generally smaller than in Table 6b.

ML					
	Estimates	SE:model-based	SE:empirical sand	SE:model-based sand	
(intercept)	1.1965	5.3871E-02	5.0702E-02	5.3871E-02	
age	-3.2756E-03	1.4232E-03	1.2062E-03	1.4232E-03	
kidney	3.1417E-01	7.0909E-02	6.8759E-02	7.0909E-02	
age:kidney	-1.6642E-02	1.8420E-03	1.7155E-03	1.8420E-03	
(a) ML					
	REML				
Estimates SE:model-based SE:empirical sand SE:model-based sand					
(intercept)	1.1966	5.3998E-02	5.0725E-02	5.3998E-02	
age	-3.2795E-03	1.4267E-03	1.2068E-03	1.4267E-03	
kidney	3.1432E-01	7.1077E-02	6.8775E-02	7.1077E-02	
age:kidney	-1.6647E-02	1.8466E-03	1.7159E-03	1.8466E-03	
(b) REML					

Table 6 - Summary of three versions of standard estimates for ML and REML

4

The REML standard error estimates are larger than the ML, because the estimated variance parameters are also slightly larger as we have seen in Part (C).

Model-based sandwich standard errors and model-based standard errors in each of the tables are the same.

If we use "summary(fit, adjustSigma=F)", then columns 2 and 4 should not be EXACTLY the same, but they are close because of a degree-of-freedom correction.

2.5 Part (e)

Table 7 gives point estimates and robust sandwich standard error estimates that account for clustering for the marginal rates of change in *scr* in subjects with and without kidney disease for the model fit by REML. From the derivation below, we have construted the combination to get the rate of change in *scr* in subjects with and without kidney disease, as well as the robust sandwich standard errors of the rate of change.

$$\hat{\beta}_{age}(Desease = T) = \hat{\beta}_1 + \hat{\beta}_{12}$$

$$= -0.01991$$

$$Var(\hat{\beta}_{age}(Desease = T)) = Var_{robust-sand}(\hat{\beta}_1 + \hat{\beta}_{12})$$

$$= (0,1,0,1) \times Var(\hat{\beta}) \times (0,1,0,1)^T$$

$$= (1.2197E - 03)^2$$

	KEML	
	Estimates	SE: robust/empirical sand
Without kidney disease	-3.2795E-03	1.2068E-03
With kidney disease	-1.9918E-02	1.2197E-03

DEL

Table 7 - Point estimates and robust sandwich standard error estimates from REML

2.6 Part (f)

The subject 20 in the data has kidney disease at three different ages that are not 60 years old. The model should include age and kidney disease status and subject id. To predict scr for a similar subject with kidney disease at age 60, we need not only the fixed effect coeffcient estimates but the estimated values of the random intercept and slope of subject 20.

Table 8 shows both estimates of fixed and random effects for the subject at age 60.

	Fixed	Random	Predicted value
With kidney disease	0.3156	0.6521	0.9676

Table 8 – A similar subject to subject 20 at age 60.

Appendix

```
## Problem 1
library(nlme)
library(Matrix)
library(penalized)
fx = function (x){
  out = (10-x)^2
 return(out)
# n.subject = 91
# m.time = 3
genXY = function(n.subject= 91,m.time= 3,sigma2=100,beta.L=1){
 x = matrix(0,n.subject,m.time)
  for (i in 1:n.subject){
    for (j in 1:m.time){
      if (j == 1){
        # baseage
        x[i,j] = 1+0.1*(i-1)
        delta.xi = (1+ (10-x[i,1])/10)^2
      } else{
        # j=2, j=3
        x[i,j] = x[i,1] + (j-1) * delta.xi
    }
  }
  xx = cbind(x, x - x[,1])
  colnames(xx) = c("base.age", "age2", "age3", "base.age-base.age", "delta.age", "2delta.age")
  beta.L = 1
   sigma2 = 100
  sigma = sqrt(sigma2)
  y = y.m = subject.long = time.long =rep(0,n.subject * m.time)
  base.age.long = delta.age.long =rep(0,n.subject * m.time)
  for (i in 1:n.subject){
    for (j in 1:m.time){
     y.m[j + (i-1)*m.time] = fx(x[i,1]) + beta.L * (x[i,j]-x[i,1])
     y[j + (i-1)*m.time] = rnorm(1,0,sigma) + y.m[j + (i-1)*m.time]
     subject.long[j + (i-1)*m.time] = i
     time.long[j + (i-1)*m.time] = j
     base.age.long[j + (i-1)*m.time] = xx[i,1]
     delta.age.long[j + (i-1)*m.time] = xx[i,3+j]
    }
  }
  df.long = data.frame(subject = subject.long,
                       time=time.long,base.age = base.age.long,
                       delta.age =delta.age.long, literacy = y,
                       y.m = y.m)
  return(list(df.long = df.long,x=x,xx=xx,y=y,y.m=y.m))
}
xx=genXY()$xx
y.m = genXY()$y.m
```

```
# sim.out = genXY()
n.sim = 1500
sim.out = list()
bl.lm.0 = bl.lm.fix = bl.lme.ran = rep(0,n.sim)
sig2.lm.fix = sig2.lme.ran = rep(0,n.sim)
sig2.subj.ran = rep(0,n.sim)
for (i in 1:n.sim){
  sim.out[[i]] = genXY()
  tmr = sim.out[[i]]
  df.long = tmr$df.long
  lm.fit = lm(literacy~ delta.age,data=df.long)
  bl.lm.0[i] = lm.fit$coef[2]
  # fit fixed effects linear reg
  lm.fit.fix = lm(literacy~ delta.age+as.factor(subject),data=df.long)
  bl.lm.fix[i] = lm.fit.fix$coef[2]
  sig2.lm.fix[i] = summary(lm.fit.fix)$sigma^2
  # fit random effects linear reg
  lme.fit.ran = lme(literacy~delta.age,
                     random=~1|as.factor(subject),
                     data=df.long)
  bl.lme.ran[i]=fixef(lme.fit.ran)["delta.age"]
  sig2.subj.ran[i]=getVarCov(lme.fit.ran)[[1]]
  sig2.lme.ran[i] = summary(lme.fit.ran)$sigma^2
}
library(lattice)
library(latticeExtra)
a1=xyplot(literacy ~ delta.age | factor(subject),
          data=df.long,
          main=NULL,
          ylab="Literacy",xlab="Delta Age",
          panel = function(x,y) {
            j = panel.number()
            panel.xyplot(x,y)
            panel.lmline(x,y,col="blue") # fit scatter per grp
            panel.abline(lm(literacy ~ delta.age,data=df.long),lty=2,col="Red") # ignore grp
          },
          auto.key=list(text=c("Conditional", "Marginal"),
                         lines=T,points=F, space = "top",col=c("blue","red")
          ))
print(a1)
df = data.frame(x=rep(xx[,1],3),y=c(y1,y2,y3),group=rep(1:3,each=length(xx[,1])))
m=ggplot(df)
m=m+
  geom_point(aes(x=x,y=y,group=factor(group),colour=factor(group),shape=factor(group)),size=4,alpha = 0
  stat\_binhex(aes(x=x,y=y,group=factor(group),colour=factor(group),shape=factor(group)))
  theme_bw()+xlab("Baseline Age")+ylab("F(xi1)")+
```

```
ggtitle("Compare the F(xi1) across true, fixed effect and random effect")+
 # guides(colour=guide_legend(title="Time"))+
     quides(shape=quide legend(title="Time"))+
  # geom_smooth(aes(x=age,y=literacy,group=factor(time),shape=factor(time)),se=FALSE,size=1)
\# geom_line(aes(x=age,y=literacy,group=factor(subject),colour=factor(time),shape=factor(time)))+
  scale_colour_discrete(name="Model",
                      breaks=1:3,
                      labels=c("True","fixed effect","random effect"))+
 scale_shape_discrete(name="Model",
                     breaks=1:3,
                     labels=c("True","fixed effect","random effect"))
m2 = m + geom_smooth(aes(x=x,y=y),data=subset(df,group==1),se=TRUE,size=1,method="auto")
m2
# lme.fit.ran$coef
\# tmr = fixef(lme.fit.ran)[1] + ranef(lme.fit.ran)[,1]
# points(x, tmr, pch=22)
sim.table = cbind(1:length(bl.lm.0),bl.lm.0,bl.lm.fix,sig2.lm.fix,bl.lme.ran,sig2.lme.ran,sig2.subj.ran
# table for ggplot2
apply(sim.table,2,mean)
apply(sim.table,2,sd)
sim.table2 = as.data.frame(cbind(rep(1:length(bl.lm.0),3),
                               c(rep(1,length(bl.lm.fix)),bl.lm.fix, bl.lme.ran),
                               rep(0:2,each=length(bl.lm.0))))
colnames(sim.table2) = c("simulation","beta.L","model")
?as.data.frame
sim.table=as.data.frame(sim.table)
head(sim.table)
apply(sim.table,2,mean)
           bl.lm.fix sig2.lm.fix bl.lme.ran sig2.lme.ran sig2.subj.ran
# bl.lm.0
# 750.5000000
                594.5482125
apply(sim.table,2,sd)
# bl.lm.0 bl.lm.fix siq2.lm.fix bl.lme.ran siq2.lme.ran siq2.subj.ran
               # 433.1570154
                                                                                  33.8165254
tail(sim.table2)
plot(1:n.sim,rep(1,n.sim),
    xlab = "Simulation",
    ylab = "Estimates of Beta.L",
    ylim = c(min(bl.lm.fix,bl.lme.ran), max(bl.lm.fix,bl.lme.ran)*1.25),
    pch = 19,"b", lty = 2, lwd = 2, col = grey(0.6),
    main="Compare of the beta.L estimates in different models")
points(1:n.sim,bl.lm.fix,pch = 6,lty = 1, col = "red", cex = 0.5)
points(1:n.sim,bl.lme.ran,pch = 13,lty = 1, col = "skyblue",cex = 0.5)
abline(h=mean(bl.lm.fix),lty = 1, col = "red",lwd=2)
abline(h=mean(bl.lme.ran),lty = 1, col = "skyblue",lwd=2)
legend("topright",
      legend =c("True Beta.L", "Mean of Beta.L in Fixed effect", "Mean of Beta.L in Random effect"),
```

```
lty = c(2,1,1),
       col = c("grey", "red", "skyblue"),
       pch= c(19,6,13),
       bty="n")
# m=qqplot(sim.table2)
\# m=m+qeom_point(aes(x=simulation,y=beta.L,qroup=factor(model),colour=factor(model),shape=factor(model)
   theme_bw()+xlab("Simulation")+ylab("Estimates of Beta.L")+
   qqtitle("Compare of the beta.L estimates in different models")+
   guides(colour=guide_legend(title="Model"))+
   guides(shape=guide_legend(title="Model"))+
   qeom smooth(aes(x=simulation,y=beta.L,qroup=factor(model),colour=factor(model)),se=FALSE,size=1)
# m
df.long$age = df.long$base.age+df.long$delta.age
col = c("red","blue","yellow")
tmr = df.long[which(df.long$subject==1),]
m=ggplot(df.long)
m=m+geom_point(aes(x=age,y=literacy,group=factor(time),colour=factor(time),shape=factor(time)))+
  theme_bw()+xlab("Age")+ylab("Literacy")+
  ggtitle("Compare the Literacy across age and subject")+
   quides(colour=quide_legend(title="Time"))+
  quides(shape=quide_legend(title="Time"))+
# qeom_smooth(aes(x=aqe,y=literacy,qroup=factor(time),shape=factor(time)),se=FALSE,size=1)
  geom_line(aes(x=age,y=literacy,group=factor(subject),colour=factor(time),shape=factor(time)))+
 scale_colour_discrete(name="Time",
                        breaks=1:3,
                        labels=c("Age=0","Age=1","Age=2"))+
  scale_shape_discrete(name ="Time",
                       breaks=c(1:3),
                       labels=c("Age=0","Age=1","Age=2"))
m2 = m + geom_smooth(aes(x=age,y=literacy),se=FALSE,size=1)
ggsave(m,file="literacy.pdf")
ggsave(m2,file="literacy2.pdf")
# warnings()
exact.bias = function (sigma2,df.long,beta.L){
  ## exact
 true.sigma2 = sigma2
 true.sigma2.gamma = var(fx(xx[,1]))*90/91
  # 596.9742
# find the exact bias using GLS
  X = cbind(1, df.long[,4])
  Y = df.long\$y.m
  Sigma = matrix(true.sigma2.gamma,nrow=3,ncol=3)
  diag(Sigma) = true.sigma2.gamma+true.sigma2
```

```
W = solve(Sigma)
  Sigma.all = matrix(0,91*3,91*3)
  for (i in 1:n.subject){
    Sigma.all[3*(i-1)+(1:3), 3*(i-1)+(1:3)] = W
  true.beta.l=(solve(t(X))*%Sigma.all\*%X))*%t(X))*%Sigma.all\*%Y)[2]
  print(true.beta.1)
  #1.273
  # find the exact bias by penalized OLS
  pen.fit = penalized(df.long$y.m,
                       penalized=~as.factor(df.long$subject),
                       unpenalized=~df.long$delta.age,
                       lambda2=true.sigma2 /true.sigma2.gamma)
  true.beta.l.pen = coef(pen.fit)[2]
  print(true.beta.l.pen)
 bias=true.beta.l-beta.L
  return(bias)
}
beta.L = 1
n.sigma2 = 150
sigma2.lst = bias.lst = seq(1,50000,l=n.sigma2)
for (i in 1:n.sigma2 ){
  sigma2 = sigma2.lst [i]
  df.long=genXY(sigma2=sigma2,beta.L=beta.L)$df.long
  bias.lst[i] = exact.bias(sigma2=sigma2,df.long=df.long,beta.L=beta.L)
plot(sigma2.lst,bias.lst,
     xlab = "True Sigma2",
     ylab = "Exact Bias",
      ylim = c(min(bl.lm.fix,bl.lme.ran), max(bl.lm.fix,bl.lme.ran)*1.25),
     pch = 19,"b", lty = 2, lwd = 1, col = grey(0.6), cex = 0.6,
     main="Exact Bias vs. True Sigma2")
## Problem 2
library(nlme)
data = read.csv("creat.csv")
head(data)
data$group = as.numeric(data$group)
data$kidney[data$group<3]=1
data$kidney[data$group>2]=0
data$interaction = data$kidney*data$age
fit.ml=lme(fixed = scr~age*kidney,
    random = reStruct(~1+age| id, pdClass="pdDiag"),
    corr=corSpher(form = ~age | id, nugget=TRUE),
   method = "ML",
    data=data
    )
```

```
# fit.ml$siqma^2
fit.reml=lme(fixed = scr~age*kidney,
    random = reStruct(~1+age| id, pdClass="pdDiag"),
    corr=corSpher(form = ~age | id, nugget=TRUE),
    method = "REML",
    data=data
)
# fit.reml$sigma^2
## 2d)
# > head(data)
# id group age scr kidney interaction
# 1 1 1 35.765 0.182 1
                                        35.765
# 2 1 1 37.990 0.088 1 37.990

# 3 2 2 24.997 1.429 1 24.997

# 4 2 2 27.441 1.111 1 27.441

# 5 2 2 30.524 1.429 1 30.524

# 6 3 1 51.083 0.156 1 51.083
ses = function (fit,data){
  A = matrix(0,4,4)
  B = matrix(0,4,4)
  B.mb = matrix(0,4,4)
  #get variance components
  range = as.numeric(coef(fit$modelStruct, unconstrained=F)[3])
  #nugget effect
  nugget = as.numeric(coef(fit$modelStruct, unconstrained=F)[4])
  # beta est.
  beta = fixef(fit)
  # how many subj.
  id.lst = unique(data$id)
  n = length(id.lst)
  # get var(gamma)
  G = matrix(c(getVarCov(fit)),2,2)
  #residual variance
  sigma2 = fit$sigma^2
  # loop in the subjects
  for(i in 1:n){
    yi = data$scr[which(data$id==id.lst[i])]
    mi = length(yi)
    tmr = data[which(data$id==id.lst[i]),c(3,5,6)]
    xi = cbind(rep(1,mi),tmr)
    # design mat
    zi = model.matrix(~1+xi$age)
    Ri = matrix(0,mi,mi)
    ## calculate rho matrix : loop within subject
    for(j in 1:mi){
      for(k in 1:mi){
         # eps over age
```

```
if(abs(xi$age[j]-xi$age[k]) < range){</pre>
         rho = (1-nugget)*( 1 - 1.5*abs(xi$age[j]-xi$age[k])/range +
                                0.5*abs(xi$age[j]-xi$age[k])^3/range^3
       }else{
         rho = 0
       if(j == k){
        rho = 1
       Ri[j, k] = sigma2*rho
     }
    if(mi==1){
     si = zi\%\%\%\%\%\%t(zi) + Ri # Simga_i matrix in the hw
     si = zi\%*\%G\%*\%t(zi) + Ri
   xi = as.matrix(xi)
   b = as.matrix(beta,1,4)
   resi = yi - xi%*%b
   B = B + t(xi)\%*\%solve(si)\%*\%resi\%*\%t(resi)\%*\%solve(si)\%*\%xi
   B.mb = B.mb + t(xi)%*%solve(si)%*%xi # canceled out.
   A = A + t(xi)\%*\%solve(si)\%*\%xi
  se.mb = sqrt(diag(summary(fit)$varFix))
  sand = solve(A)%*%B%*%solve(A)
  sand.mb = solve(A)%*%B.mb%*%solve(A)
  output = matrix(0,4,4)
  summary(fit)
  output[,1] = beta
  output[,2] = se.mb
  output[,3] = sqrt(diag(sand))
  output[,4] = sqrt(diag(sand.mb))
  colnames( output ) = c("Estimate", "SE:model-based", "SE:empirical sand", "SE:model-based sand")
  rownames( output ) = c("(intercept)", "age", "kidney", "age:kidney")
  output
 return(list(output=output, sand=sand))
}
T1=ses(fit.ml,data)$output # table of ML
# Estimate SE:model-based SE:empirical sand SE:model-based sand
# (intercept) 1.196490051 0.053870837 0.050701519
                                                                 0.053870837
             # age
                                              0.001206204
                                                                 0.001423158
# kidney
            0.314170354 0.070908899
                                              0.068759444
                                                                 0.070908899
# age:kidney -0.016642243 0.001841956
                                              0.001715466
                                                                 0.001841956
T2=ses(fit.reml,data)$output# table of REML
# Estimate SE:model-based SE:empirical sand SE:model-based sand
# (intercept) 1.19663113 0.053997862 0.050725204
                                                              0.053997862
            -0.00327953 0.001426745
# age
                                           0.001206847
                                                                0.001426745
             0.31431681 0.071077369
# kidney
                                            0.068774636
                                                                0.071077369
# age:kidney -0.01664654 0.001846595
                                            0.001715854
                                                                0.001846595
```

```
reml.sand = ses(fit.reml,data)$sand
reml.sand
## 2e)
# without kidney disease
T2[,c(1,3)]
# Estimate SE:empirical sand
# (intercept) 1.19663113 0.050725204
# with kidney disease
sqrt(c(0,1,0,1)%*%reml.sand%*%c(0,1,0,1)) #0.001219702
fixef(fit.ml)[2]+fixef(fit.ml)[4] #-0.01991786
## 2f)
data[which(data$id==20),]
# id group age scr kidney interaction
# 42 20 1 30.034 1.429 1 30.034
fixef(fit.ml) #fixed
# subject 20 at 60 years old
age.pred = 60
kidney.pred = 1
inter.pred = age.pred*kidney
fixed.20 = c(1,age.pred,kidney.pred,inter.pred) *%as.numeric(fixef(fit.ml)) #0.3156
rownames(ranef(fit.ml))[20]
random.20 = unlist(ranef(fit.ml)[20,])%*%c(1,age.pred)
random.20
random.20 + fixed.20# 0.9676927
```