Logit Models

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Basics

1.1 Logit

$$f(x) = log(\frac{p(y=1)}{1 - p(y=1)})$$

The basic idea of logistic regression:

$$p(y=1) = \frac{1}{1 + e^{-(\beta_0 + \beta_1 x_1 + \dots + \beta_n x_n)}} = \frac{e^{\beta_0 + \beta_1 x_1 + \dots + \beta_n x_n}}{1 + e^{\beta_0 + \beta_1 x_1 + \dots + \beta_n x_n}}$$

Thus, $e^{\beta_0+\beta_1x_1+...+\beta_nx_n}$ can be from $-\infty$ to $+\infty$, and p(y=1) will be always within the range of (0,1).

```
f<-function(x){exp(x)/(1+exp(x))}
data<-seq(-10,10,1)
plot(data,f(data),type = "b")</pre>
```



We can also write the function into another format as follows:

$$log\frac{p(y=1)}{1-p(y=1)}=\beta_0+\beta_1x_1+\ldots+\beta_nx_n$$

The following is an example testing whether that home teams are more likely to win in NFL games. The results show that the odd of winning is the same for both home and away teams.

```
mydata = read.csv(url('https://raw.githubusercontent.com/nfl-football-ops/Big-Data-Bow
mydata$result_new<-ifelse(mydata$HomeScore>mydata$VisitorScore,1,0)
summary(mydata$result_new)
##
                    Median
                              Mean 3rd Qu.
      Min. 1st Qu.
                                              Max.
    0.0000 0.0000 0.0000
                           0.4945 1.0000
                                           1.0000
mylogit1 = glm(result_new~1, family=binomial, data=mydata)
summary(mylogit1)
##
## Call:
## glm(formula = result_new ~ 1, family = binomial, data = mydata)
##
## Deviance Residuals:
```

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```
Min
              1Q Median
                              ЗQ
                                     Max
## -1.168 -1.168 -1.168
                          1.187
                                   1.187
##
## Coefficients:
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.02198
                          0.20967 -0.105
                                           0.917
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 126.14 on 90 degrees of freedom
## Residual deviance: 126.14 on 90 degrees of freedom
## AIC: 128.14
## Number of Fisher Scoring iterations: 3
```

1.2 Probit

$$\beta_0 + \beta_1 x_1 + \dots + \beta_n x_n = \Phi^{-1}(p)$$

Thus,

$$\Phi(\beta_0 + \beta_1 x_1 + \dots + \beta_n x_n) = p(y = 1)$$

MLE

The probability of y = 1 is as follows:

$$p = p(y = 1) = \frac{1}{1 + e^{-(\beta_0 + \beta_1 x_1 + \dots + \beta_n x_n)}} = \frac{e^{\beta_0 + \beta_1 x_1 + \dots + \beta_n x_n}}{1 + e^{\beta_0 + \beta_1 x_1 + \dots + \beta_n x_n}}$$

Thus, the likelihood function is as follows:

$$L = \prod p^{y_i} (1-p)^{1-y_i} = \prod \left(\frac{1}{1+e^{-(\beta_0+\beta_1 x_1 + \dots + \beta_n x_n)}}\right)^{y_i} \left(\frac{1}{1+e^{\beta_0+\beta_1 x_1 + \dots + \beta_n x_n}}\right)^{1-y_i}$$
$$= \prod \left(1+e^{-(\beta_0+\beta_1 x_1 + \dots + \beta_n x_n)}\right)^{-y_i} \left(1+e^{\beta_0+\beta_1 x_1 + \dots + \beta_n x_n}\right)^{-(1-y_i)}$$

Thus, the log-likelihood is as follows:

$$logL = \sum (-y_i \cdot log(1 + e^{-(\beta_0 + \beta_1 x_1 + \dots + \beta_n x_n)}) - (1 - y_i) \cdot log(1 + e^{\beta_0 + \beta_1 x_1 + \dots + \beta_n x_n}))$$

Typically, optimisers minimize a function, so we use negative log-likelihood as minimising that is equivalent to maximising the log-likelihood or the likelihood itself.

```
#Source of R code: https://www.r-bloggers.com/logistic-regression/
mle.logreg = function(fmla, data)
{
    # Define the negative log likelihood function
    logl <- function(theta,x,y){
        y <- y</pre>
```

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```
x <- as.matrix(x)
    beta <- theta[1:ncol(x)]</pre>
    # Use the log-likelihood of the Bernouilli distribution, where p is
    # defined as the logistic transformation of a linear combination
    # of predictors, according to logit(p)=(x\%*\%beta)
   loglik <- sum(-y*log(1 + exp(-(x%*%beta))) - (1-y)*log(1 + exp(x%*%beta)))
   return(-loglik)
  # Prepare the data
  outcome = rownames(attr(terms(fmla), "factors"))[1]
 dfrTmp = model.frame(data)
 x = as.matrix(model.matrix(fmla, data=dfrTmp))
 y = as.numeric(as.matrix(data[,match(outcome,colnames(data))]))
  # Define initial values for the parameters
  theta.start = rep(0, (dim(x)[2]))
  names(theta.start) = colnames(x)
  # Calculate the maximum likelihood
 mle = optim(theta.start,logl,x=x,y=y, method = 'BFGS', hessian=T)
 out = list(beta=mle$par,vcov=solve(mle$hessian),ll=2*mle$value)
mydata = read.csv(url('https://stats.idre.ucla.edu/stat/data/binary.csv'))
mylogit1 = glm(admit~gre+gpa+as.factor(rank), family=binomial, data=mydata)
mydata$rank = factor(mydata$rank) #Treat rank as a categorical variable
fmla = as.formula("admit~gre+gpa+rank") #Create model formula
mylogit2 = mle.logreg(fmla, mydata) #Estimate coefficients
 print(cbind(coef(mylogit1), mylogit2$beta))
##
                                         [,2]
                            [,1]
## (Intercept)
                    -3.989979073 -3.772676422
                     0.002264426 0.001375522
## gre
                     0.804037549 0.898201239
## gpa
## as.factor(rank)2 -0.675442928 -0.675543009
## as.factor(rank)3 -1.340203916 -1.356554831
## as.factor(rank)4 -1.551463677 -1.563396035
```

Twitter Example

The following is part of my course project for Stat 536. It aims to replicate part of the findings from Barbera (2015) Birds of the Same Feather Tweet Together: Bayesian Ideal Point Estimation Using Twitter Data. Political Analysis 23 (1). Note that, the following model is much simpler than that in the original paper.

3.1 Model

Suppose that a Twitter user is presented with a choice between following or not following another target $j \in \{1, ..., m\}$. Let $y_j = 1$ if the user decides to follow j, and $y_j = 0$ otherwise.

$$y_j = \begin{cases} 1 & Following \\ 0 & NotFollowing \end{cases}$$

$$p(y_j = 1|\theta) = \frac{exp(-\theta_0|\theta_1 - x_j|^2)}{1 + exp(-\theta_0|\theta_1 - x_j|^2)}$$

We additionally know the priors of θ .

$$\theta_i \sim N(0, 10^2)(i = 0, 1)$$

The likelihood function is as follows.

$$L(Y|\theta) = \prod_{j=1}^{m} \left(\frac{exp(-\theta_0|\theta_1 - x_j|^2)}{1 + exp(-\theta_0|\theta_1 - x_j|^2)}\right)^{y_j} \left(1 - \frac{exp(-\theta_0|\theta_1 - x_j|^2)}{1 + exp(-\theta_0|\theta_1 - x_j|^2)}\right)^{(1-y_j)}$$

Thus, the posterior is as follows.

$$L(Y|\theta) \cdot N(\theta_0|0,10) \cdot N(\theta_1|0,10)$$

$$\propto \prod_{j=1}^{m} \left(\frac{exp(-\theta_{0}|\theta_{1}-x_{j}|^{2})}{1+exp(-\theta_{0}|\theta_{1}-x_{j}|^{2})}\right)^{y_{j}} \left(1-\frac{exp(-\theta_{0}|\theta_{1}-x_{j}|^{2})}{1+exp(-\theta_{0}|\theta_{1}-x_{j}|^{2})}\right)^{(1-y_{j})} \cdot exp(-\frac{1}{2}(\frac{\theta_{0}}{10})^{2}) \cdot exp(-\frac{1}{2}(\frac{\theta_{1}}{10})^{2})$$

3.2 Simulating Data of Senators on Twitter

Assume that we have 100 senators, 50 Democrats and 50 Republicans, who we know their ideology. Assume that Democrats have negative ideology scores to indicate that they are more liberal, whereas Republicans have positive scores to indicate that they are more conservative. The following is data simulation for senators.

```
# Republicans are more conservative, and they have positive numbers.
Republicans<-c()
Republicans<-rnorm(50,1,0.5)
No_Republicans<-rep(1:50,1)
Part_1<-cbind(No_Republicans,Republicans)

# Democrats are more liberal, and they have negative numbers.
Democrats<-c()
Democrats<-rnorm(50,-1,0.5)
No_Democrats<-rep(51:100,1)
Part_2<-cbind(No_Democrats,Democrats)
Data_Elites<-rbind(Part_1,Part_2)
Data_Elites<-as.data.frame(Data_Elites)
colnames(Data_Elites) <- c("Elite_No","Elite_ideology")</pre>
```

```
Elite_No Elite_ideology
##
## 1
        1
               1.2887818
## 2
         2
               0.9773784
## 3
        3
               0.7437353
## 4
         4
               1.3321009
## 5
       5
               0.9609065
         6
## 6
               0.5025110
```

3.3 Simulating Data of Conservative Users on Twitter and Model Testing

Assume that we observe one Twitter user, who is more conservative. To simulate Twitter following data for this user, I assign this user to follow more Republican senators. Thus, if the Metropolis Hastings algorithm works as intended, we would expect to see a positive estimated value for their ideology. Importantly, as we can see in the histogram below, the estimated value indeed is positive, providing preliminary evidence for the statistical model and the algorithm. In addition, for the acceptance rate, we can see that the constant has a lower number than ideology, since we only accept a constant when it is positive.

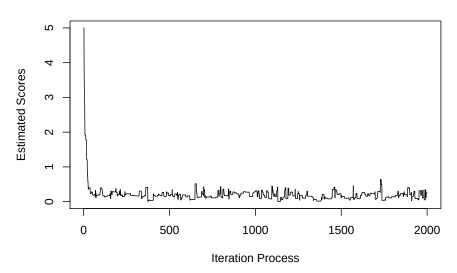
```
#This user approximately follows 45 Republican Senators and 10 Democrat Senators.
Data_user<-as.data.frame(matrix(c(ifelse(runif(50)<.1,0,1),ifelse(runif(50)<.8,0,1))), 100, 1)
colnames(Data_user)<-c("R_User")
Data_combined<-cbind(Data_Elites,Data_user)

X_data<-Data_combined$Elite_ideology
Y_data<-Data_combined$R_User

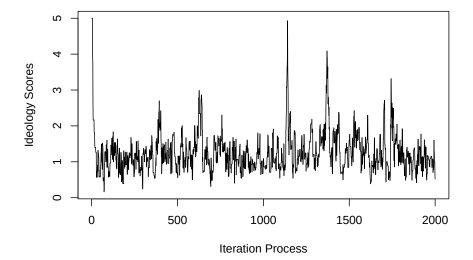
fit_C<-Bayes_logit(Y_data,X_data)
fit_C$acceptance_rate</pre>
```

```
## [1] 0.1660830 0.5637819
```

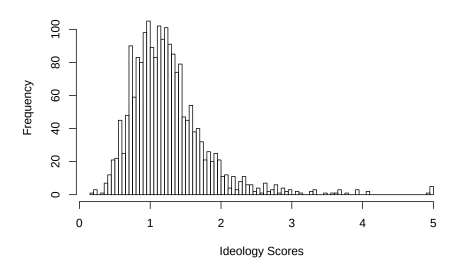
Constant (Conservative Users)



Estimated Ideology Scores (Conservative Users)



Estimated Ideology Scores (Conservative Users)



3.4 Simulating Data of Liberal Users on Twitter and Model Testing

To further verify the Metropolis Hastings algorithm, I plan to test the opposite estimate. Specifically, assume that we observe another user, who is more liberal. To simulate Twitter following data for this user, I assign this user to follow more Democrat senators. In this case, we would expect to see a negative value for their estimated ideology. As we can see in the histogram shown below, as expected, the estimated value is negative, providing convergent evidence for the model and the algorithm.

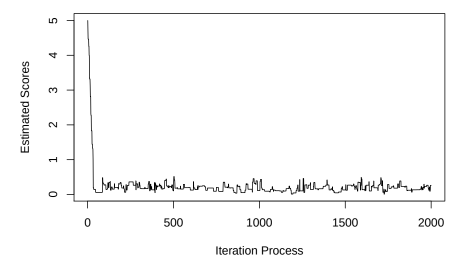
```
#This user approximately follows 10 Republican Senators and 45 Democrat Senators.
Data_user<-as.data.frame(matrix(c(ifelse(runif(50)<.8,0,1),ifelse(runif(50)<.1,0,1))), 100, 1)
colnames(Data_user)<-c("L_User")
Data_combined<-cbind(Data_Elites,Data_user)

X_data<-Data_combined$Elite_ideology
Y_data<-Data_combined$L_User</pre>
```

```
fit_L<-Bayes_logit(Y_data,X_data)
fit_L$acceptance_rate</pre>
```

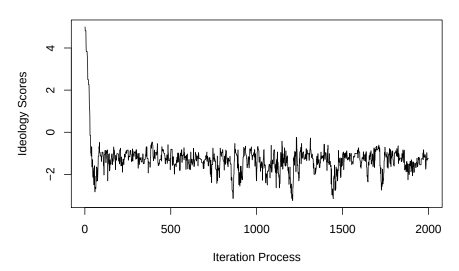
[1] 0.1605803 0.5297649

Constant (Liberal Users)

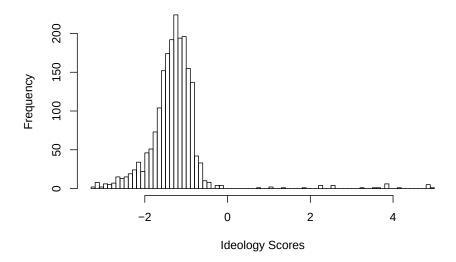


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Estimated Ideology Scores (Liberal Users)



Estimated Ideology Scores (Liberal Users)



Linear Mixed Models

The following is a shortened version of Jonathan Rosenblatt's LMM tutorial. http://www.john-ros.com/Roourse/lme.html.

In addition, another reference is from Douglas Bates's R package document. https://cran.r-project.org/web/packages/lme4/vignettes/lmer.pdf?fbclid= IwAR1nmmRP9A0BrhKdgBibNjM5acR spTpXV8QlQGdmTWyQz3ZtV3LYn6kCbQ

Assume that y is a function of x and u, where x is the fixed effect and u is the random effect. Thus, we can get,

$$y|x, u = x'\beta + z'u + \epsilon$$

For random effect, one example can be that you want to test the treatment effect, and sample 8 observations from 4 groups. You measure before and after the treatment. In this case, x represents the treatment effect, whereas z represents the group effect (i.e., random effect). Note that, in this case, it reminds the paired t-test. Remember in SPSS, why do we do paired t-test? Typically, it is the case when we measure a subject (or, participant) twice. In this case, we can consider each participant as an unit of random effect (rather than as group in the last example.)

4.1 Calculate mean

The following code generates 4 numbers (N(0,10)) for 4 groups. Then, replicate it within each group. That is, in the end, there are 8 observations.

Note that, in the following code, there are no "independent variables". Both the linear model and mixed model are actually just trying to calculate the mean. Note that $\frac{1}{y-1+1}$ and $\frac{1}{y-1+1}$ will generate the same results.

```
set.seed(123)
n.groups <- 4 # number of groups
n.repeats <- 2 # samples per group</pre>
#Generating index for observations belong to the same group
groups <- as.factor(rep(1:n.groups, each=n.repeats))</pre>
n <- length(groups)</pre>
#Generating 4 random numbers, assuming normal distribution
z0 <- rnorm(n.groups, 0, 10)</pre>
z <- z0[as.numeric(groups)] # generate and inspect random group effects
## [1] -5.6047565 -5.6047565 -2.3017749 -2.3017749 15.5870831 15.5870831 0.7050839
## [8] 0.7050839
epsilon <- rnorm(n,0,1) # generate measurement error</pre>
beta0 <- 2 # this is the actual parameter of interest! The global mean.
y <- beta0 + z + epsilon # sample from an LMM
# fit a linear model assuming independence
# i.e., assume that there is no "group things".
lm.5 < - lm(y~1)
# fit a mixed-model that deals with the group dependence
#install.packages("lme4")
library(lme4)
lme.5.a <- lmer(y~1+1|groups)</pre>
lme.5.b <- lmer(y~1|groups)</pre>
lm.5
##
## Call:
## lm(formula = y ~ 1)
##
## Coefficients:
## (Intercept)
##
         4.283
lme.5.a
## Linear mixed model fit by REML ['lmerMod']
## Formula: y ~ 1 + 1 | groups
## REML criterion at convergence: 36.1666
## Random effects:
## Groups Name
                          Std.Dev.
```

```
## groups
             (Intercept) 8.8521
## Residual
                        0.8873
## Number of obs: 8, groups: groups, 4
## Fixed Effects:
## (Intercept)
        4.283
lme.5.b
## Linear mixed model fit by REML ['lmerMod']
## Formula: y ~ 1 | groups
## REML criterion at convergence: 36.1666
## Random effects:
                        Std.Dev.
## Groups Name
## groups (Intercept) 8.8521
## Residual
                       0.8873
## Number of obs: 8, groups: groups, 4
## Fixed Effects:
## (Intercept)
        4.283
##
```

4.2 Test the treatment effect

As we can see that, LLM and paired t-test generate the same t-value.

2

3

3

4

4

2

1

2

1

2

[4,] -1.5668361

[5,] 16.9002303

[6,] 17.1414212 ## [7,] 3.9291657

[8,] 3.0648977

```
times<-rep(c(1,2),4) # first time and second time
times

## [1] 1 2 1 2 1 2 1 2

data_combined<-cbind(y,groups,times)
data_combined

## y groups times
## [1,] -3.4754687 1 1
## [2,] -1.8896915 1 2
## [3,] 0.1591413 2 1</pre>
```

```
lme_diff_times<- lmer(y~times+(1|groups))</pre>
t_results<-t.test(y~times, paired=TRUE)</pre>
lme_diff_times
## Linear mixed model fit by REML ['lmerMod']
## Formula: y ~ times + (1 | groups)
## REML criterion at convergence: 35.0539
## Random effects:
## Groups Name
                      Std.Dev.
## groups (Intercept) 8.845
## Residual
                         1.013
## Number of obs: 8, groups: groups, 4
## Fixed Effects:
## (Intercept)
                     times
       4.5691
                    -0.1908
print("The following results are from paired t-test")
## [1] "The following results are from paired t-test"
t_results$statistic
##
## 0.2664793
```

4.3 Another example

```
data(Dyestuff, package='lme4')
attach(Dyestuff)

## The following objects are masked from Dyestuff (pos = 5):
##
## Batch, Yield

Dyestuff
```

```
##
      Batch Yield
## 1
         A 1545
## 2
         A 1440
## 3
         A 1440
## 4
         A 1520
## 5
         A 1580
## 6
         B 1540
## 7
         B 1555
## 8
         B 1490
## 9
         B 1560
## 10
         B 1495
## 11
         C 1595
## 12
         C 1550
## 13
         C 1605
## 14
         C 1510
         C 1560
## 15
## 16
         D 1445
## 17
         D 1440
## 18
         D 1595
         D 1465
## 19
## 20
         D 1545
## 21
         E 1595
## 22
        E 1630
## 23
         E 1515
## 24
         E 1635
## 25
         E 1625
## 26
         F 1520
## 27
         F 1455
## 28
         F 1450
## 29
         F 1480
## 30
         F 1445
lme_batch<- lmer( Yield ~ 1 + (1|Batch) , Dyestuff )</pre>
summary(lme_batch)
## Linear mixed model fit by REML ['lmerMod']
## Formula: Yield ~ 1 + (1 | Batch)
##
     Data: Dyestuff
##
## REML criterion at convergence: 319.7
## Scaled residuals:
              1Q Median
      Min
                               3Q
                                      Max
## -1.4117 -0.7634 0.1418 0.7792 1.8296
##
```

```
## Random effects:
## Groups Name Variance Std.Dev.
## Batch (Intercept) 1764 42.00
## Residual 2451 49.51
## Number of obs: 30, groups: Batch, 6
##
## Fixed effects:
## Estimate Std. Error t value
## (Intercept) 1527.50 19.38 78.8
```

4.4 Full LMM model

```
data(sleepstudy, package='lme4')
attach(sleepstudy)
## The following objects are masked from sleepstudy (pos = 5):
##
##
      Days, Reaction, Subject
head(sleepstudy)
    Reaction Days Subject
## 1 249.5600 0
                     308
## 2 258.7047 1
                     308
## 3 250.8006 2
                    308
## 4 321.4398 3 308
## 5 356.8519 4 308
## 6 414.6901 5 308
nrow(sleepstudy)
## [1] 180
#lmer ( Reaction ~ Days + ( Days | Subject ) , data= sleepstudy )
lmer ( Reaction ~ Days + ( Days | Subject ) , data= sleepstudy )
## Linear mixed model fit by REML ['lmerMod']
## Formula: Reaction ~ Days + (Days | Subject)
     Data: sleepstudy
## REML criterion at convergence: 1743.628
```

```
## Random effects:
   Groups
                         Std.Dev. Corr
             Name
   Subject
             (Intercept) 24.737
##
             Days
                          5.923
                                   0.07
## Residual
                         25.592
## Number of obs: 180, groups: Subject, 18
## Fixed Effects:
## (Intercept)
                       Days
        251.41
##
                      10.47
```

4.5 Serial correlations in time and space

The hierarchical model of $y|x, u = x'\beta + z'u + \epsilon$ can work well for correlations within blocks, but not for correlations in time as the correlations decay in time. The following uses nlme package to calculate time serial data.

```
library(nlme)
head(nlme::Ovary,n=50)
```

```
## Grouped Data: follicles ~ Time | Mare
##
      Mare
                  Time follicles
## 1
         1 -0.13636360
                              20
## 2
         1 -0.09090910
                              15
## 3
         1 -0.04545455
                              19
## 4
         1 0.00000000
                              16
## 5
         1
           0.04545455
                              13
## 6
         1 0.09090910
                              10
## 7
         1 0.13636360
                              12
## 8
         1 0.18181820
                              14
## 9
         1
           0.22727270
                              13
## 10
         1 0.27272730
                              20
## 11
         1 0.31818180
                              22
## 12
         1 0.36363640
                              15
## 13
         1 0.40909090
                              18
## 14
         1 0.45454550
                              17
## 15
         1 0.50000000
                              14
## 16
         1 0.54545450
                              18
## 17
         1 0.59090910
                              14
## 18
         1 0.63636360
                              16
## 19
         1 0.68181820
                              17
## 20
         1 0.72727270
                              18
## 21
         1 0.77272730
                              18
## 22
        1 0.81818180
                              17
## 23
         1 0.86363640
                              14
```

##

```
## 24
        1 0.90909090
                             12
## 25
        1 0.95454550
                             12
## 26
                             14
        1 1.00000000
## 27
        1 1.04545500
                             10
        1 1.09090900
## 28
                             11
## 29
        1 1.13636400
                             16
## 30
      2 -0.15000000
                              6
## 31
      2 -0.10000000
                              6
## 32
        2 -0.05000000
                              8
## 33
                              7
        2 0.00000000
        2 0.05000000
## 34
                             16
## 35
        2 0.10000000
                             10
## 36
        2 0.15000000
                             13
## 37
        2 0.20000000
                              9
                              7
## 38
        2 0.25000000
## 39
        2 0.30000000
                              6
## 40
        2 0.35000000
                              8
## 41
                              8
        2 0.40000000
## 42
        2 0.45000000
                              6
## 43
        2 0.50000000
                              8
## 44
        2 0.55000000
                              7
                              9
## 45
      2 0.60000000
## 46
        2 0.65000000
                              6
        2 0.70000000
## 47
                              4
## 48
        2 0.75000000
                              5
## 49
        2 0.80000000
                              8
## 50
        2 0.85000000
                             11
fm10var.lme <- nlme::lme(fixed=follicles ~ sin(2*pi*Time) + cos(2*pi*Time),</pre>
                  data = Ovary,
                  random = pdDiag(~sin(2*pi*Time)),
                  correlation=corAR1() )
summary(fm10var.lme)
## Linear mixed-effects model fit by REML
## Data: Ovary
##
         AIC
                 BIC
                      logLik
##
    1563.448 1589.49 -774.724
##
## Random effects:
## Formula: ~sin(2 * pi * Time) | Mare
   Structure: Diagonal
##
          (Intercept) sin(2 * pi * Time) Residual
## StdDev: 2.858385
                               1.257977 3.507053
```

```
## Correlation Structure: AR(1)
## Formula: ~1 | Mare
## Parameter estimate(s):
        Phi
## 0.5721866
## Fixed effects: follicles ~ sin(2 * pi * Time) + cos(2 * pi * Time)
                       Value Std.Error DF t-value p-value
## (Intercept)
                12.188089 0.9436602 295 12.915760 0.0000
## sin(2 * pi * Time) -2.985297 0.6055968 295 -4.929513 0.0000
## cos(2 * pi * Time) -0.877762 0.4777821 295 -1.837159 0.0672
## Correlation:
##
                     (Intr) s(*p*T
## sin(2 * pi * Time) 0.000
## cos(2 * pi * Time) -0.123 0.000
##
## Standardized Within-Group Residuals:
          Min
                       Q1
                                 Med
                                              Q3
## -2.34910093 -0.58969626 -0.04577893 0.52931186 3.37167486
## Number of Observations: 308
## Number of Groups: 11
```

Generalized Linear Mixed Models

5.1 Basics

The following is the note from Charle E. McCulloch's "Maximum likelihood algorithems for Generalized Linear Mixed Models"

5.2 Some References

http://www.biostat.umn.edu/~baolin/teaching/linmods/glmm.html

 $http://www.biostat.umn.edu/{\sim}baolin/teaching/probmods/GLMM_mcmc. \\ html$

https://bbolker.github.io/mixed models-misc/glmmFAQ.html