Encephalitis - Loglinear Poisson Model and Normal Distribution Model

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First of all, the encephalitis data are loaded:

```
> library(catdata)
> data(encephalitis)
> attach(encephalitis)
   Some variables are renamed and recoded before fitting the model.
> BAV <- country
> BAV[BAV==2] <-0
> TIME <- year
The number of infections (count) is modeled in dependence on country and
TIME. A Loglinear Poisson Model is fitted.
> enc1 <- glm(count ~ TIME+I(TIME^2)+BAV+TIME*BAV, family = poisson)
> summary(enc1)
Call:
glm(formula = count ~ TIME + I(TIME^2) + BAV + TIME * BAV, family = poisson)
Deviance Residuals:
    Min
              1Q
                  Median
                                 ЗQ
                                         Max
-1.7747 -0.4820
                   0.0403
                             0.5141
                                      1.2125
Coefficients:
             Estimate Std. Error z value Pr(>|z|)
```

(Dispersion parameter for poisson family taken to be 1)

Null deviance: 77.434 on 25 degrees of freedom Residual deviance: 12.855 on 21 degrees of freedom

```
AIC: 105.74
Number of Fisher Scoring iterations: 4
For comparison the linear Normal Model with the identity link is fitted.
> enc2 <- glm(count ~ TIME+I(TIME^2)+BAV+TIME*BAV, family = gaussian("identity"))</pre>
> summary(enc2)
Call:
glm(formula = count ~ TIME + I(TIME^2) + BAV + TIME * BAV, family = gaussian("identity"))
Deviance Residuals:
   Min 1Q Median
                                3Q
                                        Max
-4.1325 -1.4000 -0.0303
                           1.4372
                                     4.2604
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.39710 1.69760
                                0.234 0.817312
TIME
            1.15424
                     0.47280
                                2.441 0.023577 *
                       0.03027 -2.166 0.042002 *
I(TIME^2)
            -0.06554
BAV
            -4.41444
                        1.79700 -2.457 0.022816 *
TIME:BAV
            0.85309
                        0.20713
                                4.119 0.000489 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for gaussian family taken to be 4.847447)
   Null deviance: 399.54 on 25 degrees of freedom
Residual deviance: 101.80 on 21 degrees of freedom
AIC: 121.27
Number of Fisher Scoring iterations: 2
Fit of loglinear Normal Model. That means a normal model with log-link.
> enc3 <- glm(count ~ TIME+I(TIME^2)+BAV+TIME*BAV, family = gaussian("log"),
              start=enc1$coef)
> summary(enc3)
Call:
glm(formula = count ~ TIME + I(TIME^2) + BAV + TIME * BAV, family = gaussian("log"),
   start = enc1$coef)
Deviance Residuals:
   Min
         10
                 Median
                                3Q
                                        Max
-5.3340 -0.8481
                  0.0009
                            1.1639
                                     3.9298
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
```

(Intercept) -0.223708 0.592457 -0.378 0.70952

```
TIME 0.499564 0.134163 3.724 0.00126 **
I(TIME^2) -0.029337 0.007919 -3.704 0.00131 **
BAV -1.478283 0.621729 -2.378 0.02700 *
TIME:BAV 0.198575 0.062320 3.186 0.00444 **
```

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for gaussian family taken to be 3.701177)

Null deviance: 399.538 on 25 degrees of freedom Residual deviance: 77.724 on 21 degrees of freedom

AIC: 114.26

Number of Fisher Scoring iterations: 4