troutegg

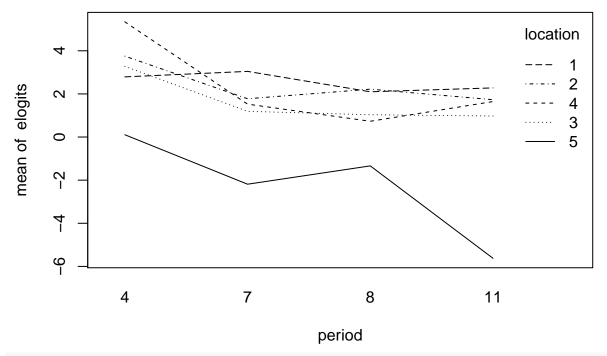
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Boxes of trout eggs were buried at five different stream locations and retrieved at 4 different times. The number of surviving eggs was recorded. Question of interest is whether survival of trout eggs depending on time and location.

```
time and location.
library(faraway)
data(troutegg)
?troutegg
str(troutegg)
  'data.frame':
                    20 obs. of 4 variables:
    $ survive : int 89 106 119 104 49 94 91 100 80 11 ...
              : int 94 108 123 104 93 98 106 130 97 113 ...
    $ location: Factor w/5 levels "1","2","3","4",...: 1 2 3 4 5 1 2 3 4 5 ...
   $ period : Factor w/ 4 levels "4","7","8","11": 1 1 1 1 1 2 2 2 2 2 ...
Look at data and check for interaction between location and period.
ftable(xtabs(cbind(survive,total) ~ location + period, troutegg))
##
                    survive total
## location period
```

```
## 1
              4
                              89
                                     94
##
              7
                              94
                                     98
##
              8
                              77
                                     86
##
              11
                             141
                                    155
                             106
                                    108
## 2
##
              7
                              91
                                    106
##
              8
                              87
                                     96
##
              11
                             104
                                    122
                             119
## 3
                                    123
              7
                             100
                                    130
##
##
              8
                              88
                                    119
##
              11
                              91
                                    125
## 4
                             104
                                    104
              7
                              80
                                     97
##
                              67
                                     99
##
              8
##
              11
                             111
                                    132
## 5
              4
                              49
                                     93
##
              7
                              11
                                    113
##
              8
                              18
                                     88
##
                                    138
```

```
elogits <- log((troutegg$survive+0.5)/(troutegg$total-troutegg$survive+0.5))
with(troutegg, interaction.plot(period, location, elogits))</pre>
```

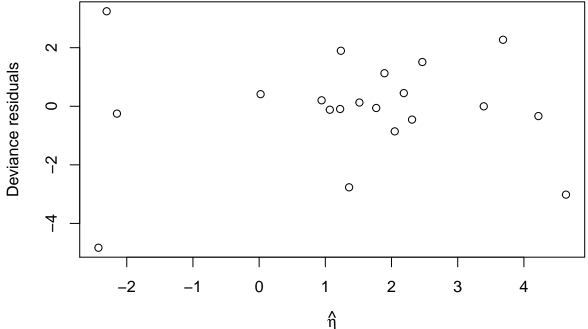


no obvious interaction

Fit a binomial regression with logit link. Residual deviance is very high - something is wrong with the model! bmod <- glm(cbind(survive,total-survive) ~ location+period, family=binomial, troutegg) summary(bmod)

```
##
## Call:
  glm(formula = cbind(survive, total - survive) ~ location + period,
##
       family = binomial, data = troutegg)
##
##
  Deviance Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                           Max
  -4.8305
           -0.3650 -0.0303
                                        3.2434
##
                               0.6191
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                 4.6358
                            0.2813
                                    16.479 < 2e-16 ***
                                    -1.694
                            0.2461
                                             0.0903
## location2
                -0.4168
## location3
                -1.2421
                            0.2194
                                    -5.660 1.51e-08 ***
## location4
                -0.9509
                            0.2288
                                    -4.157 3.23e-05 ***
## location5
                -4.6138
                            0.2502 - 18.439
                                            < 2e-16 ***
                                    -9.103
                                            < 2e-16 ***
## period7
                -2.1702
                            0.2384
                            0.2429
                                   -9.573
                                            < 2e-16 ***
## period8
                -2.3256
## period11
                -2.4500
                            0.2341 -10.466
                                            < 2e-16 ***
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 1021.469
                                on 19
                                       degrees of freedom
## Residual deviance:
                        64.495
                                on 12
                                       degrees of freedom
## AIC: 157.03
```

```
##
## Number of Fisher Scoring iterations: 5
pchisq(deviance(bmod),12, lower.tail=F)
## [1] 3.379416e-09
Check residuals - no trends.
plot(residuals(bmod) ~ predict(bmod,type="link"),
     xlab=expression(hat(eta)), ylab="Deviance residuals")
```



Let's estimate phi - should be close to 1 for a binomial regression, but isn't!

```
n <- 20
p <- 8
(phihat <- sum(residuals(bmod,type="pearson")^2)/(n-p))</pre>
```

[1] 5.330322

Fitting quasibinomial model.

```
# fitting quasibinomial model
qbmod <- glm(cbind(survive,total-survive) ~ location+period, family=quasibinomial, troutegg)</pre>
summary(qbmod)
##
## Call:
   glm(formula = cbind(survive, total - survive) ~ location + period,
       family = quasibinomial, data = troutegg)
##
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                            Max
                                0.6191
  -4.8305 -0.3650 -0.0303
                                         3.2434
##
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
```

```
## (Intercept)
               4.6358
                            0.6495
                                    7.138 1.18e-05 ***
                            0.5682 -0.734 0.477315
## location2
               -0.4168
## location3
               -1.2421
                            0.5066 -2.452 0.030501 *
## location4
               -0.9509
                            0.5281
                                   -1.800 0.096970 .
## location5
               -4.6138
                            0.5777
                                   -7.987 3.82e-06 ***
## period7
               -2.1702
                            0.5504
                                   -3.943 0.001953 **
               -2.3256
## period8
                            0.5609 -4.146 0.001356 **
## period11
               -2.4500
                            0.5405 -4.533 0.000686 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for quasibinomial family taken to be 5.330358)
##
                               on 19 degrees of freedom
##
       Null deviance: 1021.469
## Residual deviance:
                        64.495 on 12 degrees of freedom
## AIC: NA
##
## Number of Fisher Scoring iterations: 5
0.2461*sqrt(phihat)
## [1] 0.5681831
Fitting quasibinomial model by rescaling variances by phihat
summary(bmod, dispersion=phihat)
##
## Call:
## glm(formula = cbind(survive, total - survive) ~ location + period,
       family = binomial, data = troutegg)
##
## Deviance Residuals:
##
      Min
                1Q
                    Median
                                   3Q
                                          Max
## -4.8305 -0.3650 -0.0303
                              0.6191
                                        3.2434
##
## Coefficients:
##
              Estimate Std. Error z value Pr(>|z|)
## (Intercept)
               4.6358
                           0.6495
                                    7.138 9.49e-13 ***
                            0.5682 -0.734
## location2
               -0.4168
                                            0.4632
               -1.2421
## location3
                            0.5066 - 2.452
                                            0.0142 *
## location4
                            0.5281 -1.800
               -0.9509
                                            0.0718 .
## location5
               -4.6138
                            0.5777
                                   -7.987 1.39e-15 ***
## period7
               -2.1702
                            0.5504 -3.943 8.05e-05 ***
## period8
               -2.3256
                            0.5609 -4.146 3.38e-05 ***
                            0.5405 -4.533 5.82e-06 ***
## period11
               -2.4500
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 5.330322)
##
      Null deviance: 1021.469 on 19 degrees of freedom
## Residual deviance:
                       64.495
                               on 12 degrees of freedom
## AIC: 157.03
##
## Number of Fisher Scoring iterations: 5
```

```
Test for significance of location.
```

```
summary(qbmod2)
##
## Call:
## glm(formula = cbind(survive, total - survive) ~ period, family = quasibinomial,
       data = troutegg)
##
## Deviance Residuals:
##
       Min
                   10
                        Median
                                       30
                                                Max
## -17.3778
               0.8058
                        3.0192
                                   4.5949
                                             7.1826
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                 2.139
                             1.070
                                    1.999
                                            0.0629 .
## period7
                -1.333
                             1.277 -1.044
                                             0.3118
                 -1.336
                             1.298 -1.029
## period8
                                             0.3186
## period11
                -1.453
                             1.233 -1.178
                                            0.2561
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for quasibinomial family taken to be 56.33299)
##
##
       Null deviance: 1021.47 on 19 degrees of freedom
## Residual deviance: 913.56 on 16 degrees of freedom
## AIC: NA
##
## Number of Fisher Scoring iterations: 5
anova(qbmod, qbmod2, test="F")
## Analysis of Deviance Table
##
## Model 1: cbind(survive, total - survive) ~ location + period
## Model 2: cbind(survive, total - survive) ~ period
    Resid. Df Resid. Dev Df Deviance
                                           F
## 1
            12
                    64.50
## 2
                   913.56 -4 -849.06 39.822 7.779e-07 ***
            16
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# check calculation of F statistic for testing location
(913.56 - 64.5)/4/phihat
## [1] 39.82218
Or using
bmod2 <- glm(cbind(survive,total-survive) ~ period, family=binomial, troutegg)</pre>
anova(bmod, bmod2, dispersion=phihat, test="F")
## Analysis of Deviance Table
##
## Model 1: cbind(survive, total - survive) ~ location + period
## Model 2: cbind(survive, total - survive) ~ period
```

qbmod2 <- glm(cbind(survive,total-survive) ~ period, family=quasibinomial, troutegg)

```
## Resid. Df Resid. Dev Df Deviance F Pr(>F)
## 1     12     64.50
## 2     16     913.56 -4 -849.06     39.822 7.779e-07 ***
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```