# GLM models of the effects of MHC-I functional divergence on lifetime fitness in great reed warblers (Acrocephalus arundinaceus)

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This document presents the R code and output from the GLM models behind the results on the effects of MHC-I functional divergence on fitness in adult great reed warblers reported in the paper:

Roved J., Hansson B., Tarka M., Hasselquist D., & Westerdahl H. (2019). MHC-I functional divergence is positively associated with life span and fitness in male great reed warblers: support for the divergent allele advantage hypothesis. Molecular Ecology.

The data set is available at the datadryad.org repository, the link is specified in the paper mentioned above.

Note: The variable names employed in the models stem from our original analyses. The names of the corresponding variables in the data set that is publicly available at the respository are given below:

```
LifeSpan = Life span

ToSucLife_1014 = Lifetime no. fledglings (excluding unsuccessful first time breeders)

RecruLife_1014 = Lifetime no. recruiting offspring (excluding unsuccessful first time breeders)

Total.no.alleles = No. MHC-I alleles

PdistPBR = P-distance PBR

PdistPSS = P-distance PSS

Subset the data.

data.males <- subset(data, Sex=="male")

data.females <- subset(data, Sex=="female")

Load R packages.

library(MASS)
```

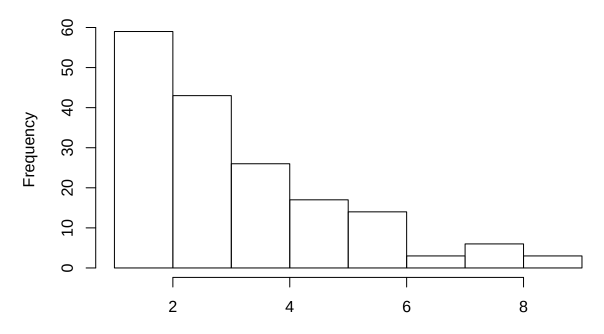
## Investigating the distribution of the fitness variables

We exclude samples for which we have no genetic data and unsuccessful first time breeders (by only selecting samples that have a value in the variables PdistPBR and ToSucLife\_1014).

### Life span

```
hist(data$LifeSpan[!is.na(data$PdistPBR)&!is.na(data$ToSucLife_1014)])
```

# :ogram of data\$LifeSpan[!is.na(data\$PdistPBR) & !is.na(data\$ToSucLife



data\$LifeSpan[!is.na(data\$PdistPBR) & !is.na(data\$ToSucLife\_1014)]

```
mean(data$LifeSpan[!is.na(data$PdistPBR)&!is.na(data$ToSucLife_1014)])

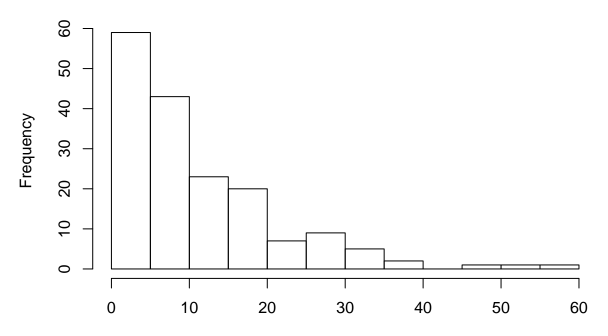
## [1] 3.473684

(sd(data$LifeSpan[!is.na(data$PdistPBR)&!is.na(data$ToSucLife_1014)]))^2

## [1] 3.603715

Lifetime number of fledglings
hist(data$ToSucLife_1014[!is.na(data$PdistPBR)&!is.na(data$ToSucLife_1014)])
```

# am of data\$ToSucLife\_1014[!is.na(data\$PdistPBR) & !is.na(data\$ToSuc



data\$ToSucLife\_1014[!is.na(data\$PdistPBR) & !is.na(data\$ToSucLife\_1014)]

```
mean(data$ToSucLife_1014[!is.na(data$PdistPBR)&!is.na(data$ToSucLife_1014)])
```

```
## [1] 12.05263
```

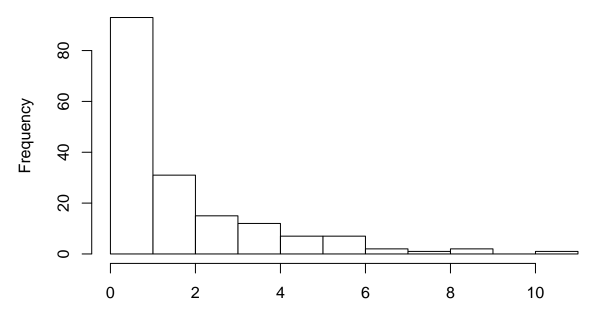
```
(sd(data$ToSucLife_1014[!is.na(data$PdistPBR)&!is.na(data$ToSucLife_1014)]))^2
```

## [1] 99.45015

### Lifetime number of recruits

hist(data\$RecruLife\_1014[!is.na(data\$PdistPBR)&!is.na(data\$ToSucLife\_1014)])

# am of data\$RecruLife\_1014[!is.na(data\$PdistPBR) & !is.na(data\$ToSuc



data\$RecruLife\_1014[!is.na(data\$PdistPBR) & !is.na(data\$ToSucLife\_1014)]

```
mean(data$RecruLife_1014[!is.na(data$PdistPBR)&!is.na(data$ToSucLife_1014)])
## [1] 1.900585
(sd(data$RecruLife_1014[!is.na(data$PdistPBR)&!is.na(data$ToSucLife_1014)]))^2
## [1] 4.395941
```

Life span, lifetime number of fledglings, and lifetime number of recruits all appear to follow negative binomial distributions.

For the GLM models, we have estimated the overdispersion parameter of the negative binomial distributions using a maximum likelihood based method (kfit function) presented in [Michael J Crawley, The R Book, Wiley, 2007].

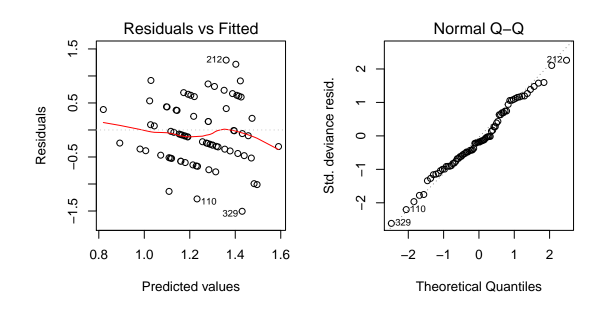
Function to test for overdispersion in GLM models using Pearson residuals (courtesy of Ben Bolker).

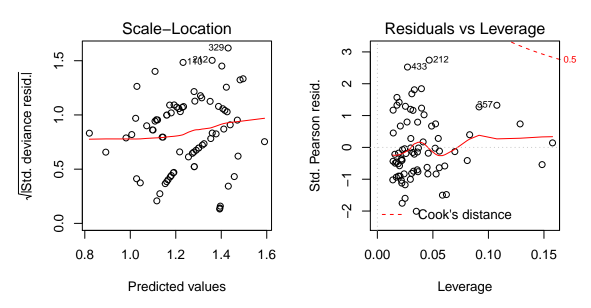
```
overdisp_fun <- function(model) {
  rdf <- df.residual(model)
  rp <- residuals(model,type="pearson")
  Pearson.chisq <- sum(rp^2)
  prat <- Pearson.chisq/rdf
  pval <- pchisq(Pearson.chisq, df=rdf, lower.tail=FALSE)
  c(chisq=Pearson.chisq,ratio=prat,rdf=rdf,p=pval)
}</pre>
```

# Modeling the effects of MHC-I functional divergence on life span

### GLM of life span on total number of alleles and P-distance PBR in males only

```
fm1_PBR <- glm(LifeSpan~Total.no.alleles+PdistPBR,negative.binomial(theta=5.14632,link="log"),data=data
s1_PBR <- summary(fm1_PBR)</pre>
s1_PBR
##
## Call:
## glm(formula = LifeSpan ~ Total.no.alleles + PdistPBR, family = negative.binomial(theta = 5.14632,
      link = "log"), data = data.males[!is.na(PdistPBR) & !is.na(ToSucLife_1014),
##
      ])
##
## Deviance Residuals:
      Min
##
                                           Max
                 1Q
                     Median
                                   3Q
## -1.5050 -0.4432 -0.1066
                               0.3828
                                        1.2933
##
## Coefficients:
##
                    Estimate Std. Error t value Pr(>|t|)
                                1.30462 -2.118 0.03755 *
## (Intercept)
                    -2.76368
## Total.no.alleles 0.02781
                                0.01811
                                          1.535 0.12902
                                          3.176 0.00219 **
## PdistPBR
                    11.66330
                                3.67272
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for Negative Binomial(5.1463) family taken to be 0.3432786)
##
      Null deviance: 28.754 on 75 degrees of freedom
##
## Residual deviance: 25.249 on 73 degrees of freedom
## AIC: 303.35
## Number of Fisher Scoring iterations: 4
overdisp_fun(fm1_PBR)
##
                                rdf
        chisq
                   ratio
## 25.0593297 0.3432785 73.0000000 1.0000000
par(mfrow=c(2,2))
plot(fm1_PBR)
```





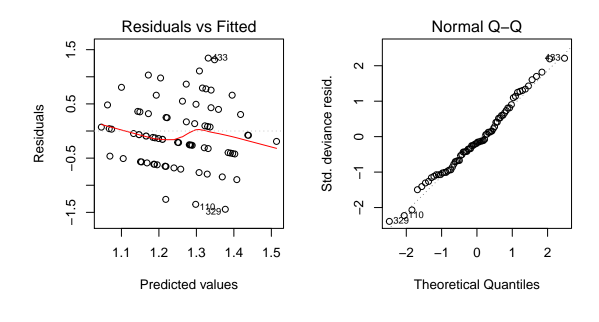
GLM of life span on total number of alleles and P-distance PSS in males only

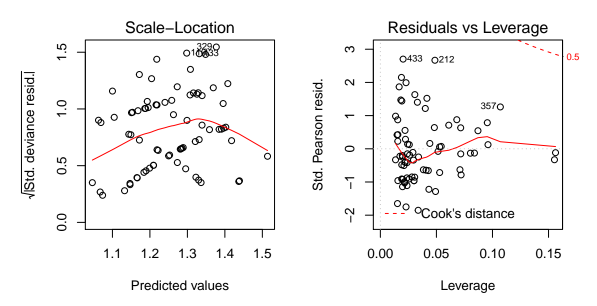
##

```
s1_PSS <- summary(fm1_PSS)</pre>
s1_PSS
##
## Call:
   glm(formula = LifeSpan ~ Total.no.alleles + PdistPSS, family = negative.binomial(theta = 5.14632,
       link = "log"), data = data.males[!is.na(PdistPSS) & !is.na(ToSucLife_1014),
##
##
       ])
##
## Deviance Residuals:
##
       Min
                  1Q
                       Median
                                     3Q
                                             Max
##
  -1.4432
            -0.4743
                     -0.1057
                                0.3282
                                          1.3444
```

fm1\_PSS <- glm(LifeSpan~Total.no.alleles+PdistPSS,negative.binomial(theta=5.14632,link="log"),data=data

```
## Coefficients:
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) -2.31280 1.75402 -1.319 0.1914
## Total.no.alleles 0.02288 0.02022 1.132 0.2615
                    9.01796 4.32516 2.085 0.0406 *
## PdistPSS
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for Negative Binomial(5.1463) family taken to be 0.3774148)
##
       Null deviance: 28.754 on 75 degrees of freedom
## Residual deviance: 27.094 on 73 degrees of freedom
## AIC: 305.19
##
\mbox{\tt \#\#} Number of Fisher Scoring iterations: 4
overdisp_fun(fm1_PSS)
        chisq
                   ratio
                                rdf
## 27.5512777 0.3774148 73.0000000 0.9999997
par(mfrow=c(2,2))
plot(fm1_PSS)
```





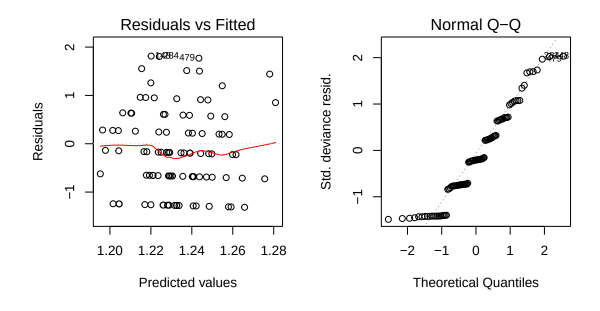
GLM of life span on total number of alleles and P-distance PBR in females only

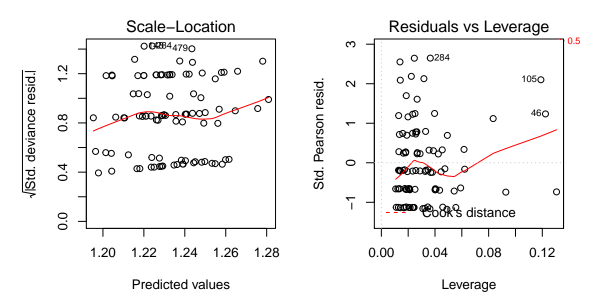
##

```
s2_PBR <- summary(fm2_PBR)</pre>
s2_PBR
##
## Call:
   glm(formula = LifeSpan ~ Total.no.alleles + PdistPBR, family = negative.binomial(theta = 5.14632,
       link = "log"), data = data.females[!is.na(PdistPBR) & !is.na(ToSucLife_1014),
##
##
       ])
##
## Deviance Residuals:
##
       Min
                  1Q
                       Median
                                     3Q
                                             Max
##
  -1.3148
            -0.6869
                      -0.1884
                                0.5797
                                          1.8134
```

fm2\_PBR <- glm(LifeSpan~Total.no.alleles+PdistPBR,negative.binomial(theta=5.14632,link="log"),data=data

```
## Coefficients:
##
                    Estimate Std. Error t value Pr(>|t|)
                  1.264632 1.786757 0.708
## (Intercept)
                                                  0.481
## Total.no.alleles 0.005697 0.022879 0.249
                                                  0.804
## PdistPBR
                  -0.346263 5.370565 -0.064
                                                  0.949
##
## (Dispersion parameter for Negative Binomial(5.1463) family taken to be 0.8213017)
##
      Null deviance: 72.819 on 94 degrees of freedom
## Residual deviance: 72.750 on 92 degrees of freedom
## AIC: 404.93
## Number of Fisher Scoring iterations: 4
overdisp_fun(fm2_PBR)
##
       chisq
                  ratio
                              rdf
## 75.5597338  0.8213015 92.0000000  0.8930381
par(mfrow=c(2,2))
plot(fm2_PBR)
```





GLM of life span on total number of alleles and P-distance PSS in females only

##

##

-1.4313

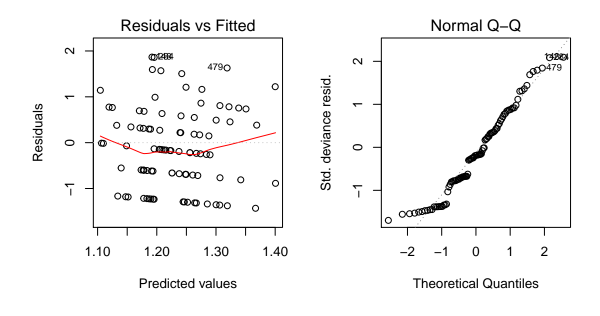
-0.6993

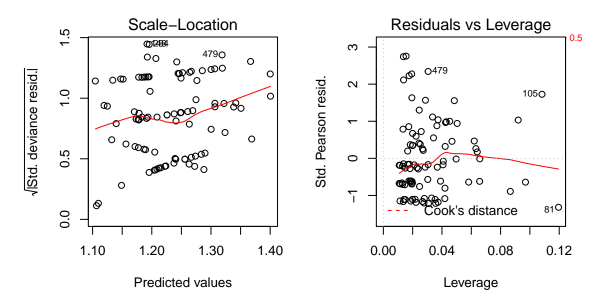
-0.1710

0.4724

```
fm2_PSS <- glm(LifeSpan~Total.no.alleles+PdistPSS,negative.binomial(theta=5.14632,link="log"),data=data
s2_PSS <- summary(fm2_PSS)</pre>
s2_PSS
##
## Call:
   glm(formula = LifeSpan ~ Total.no.alleles + PdistPSS, family = negative.binomial(theta = 5.14632,
       link = "log"), data = data.females[!is.na(PdistPSS) & !is.na(ToSucLife_1014),
##
##
       ])
##
  Deviance Residuals:
##
##
       Min
                  1Q
                       Median
                                    3Q
                                            Max
```

```
## Coefficients:
##
                    Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                  3.179939 2.035643 1.562
## Total.no.alleles -0.001591 0.022660 -0.070
                                                  0.944
## PdistPSS
                  -5.357085 5.362257 -0.999
                                                  0.320
##
## (Dispersion parameter for Negative Binomial(5.1463) family taken to be 0.8063923)
##
      Null deviance: 72.819 on 94 degrees of freedom
## Residual deviance: 71.945 on 92 degrees of freedom
## AIC: 404.13
## Number of Fisher Scoring iterations: 4
overdisp_fun(fm2_PSS)
##
       chisq
                  ratio
                               rdf
## 74.1880032  0.8063913  92.0000000  0.9129155
par(mfrow=c(2,2))
plot(fm2_PSS)
```





GLM of life span on total number of alleles and P-distance PBR in both sexes

3Q

0.4280

## Deviance Residuals:

Median

-0.1750

1Q

-0.6573

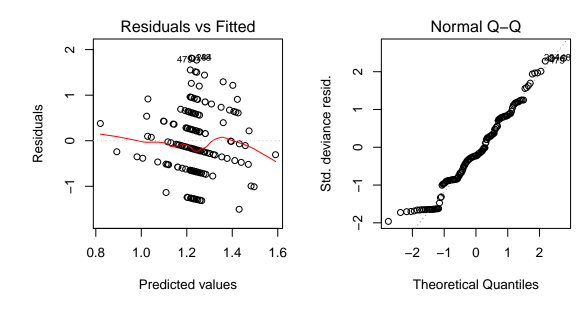
Min

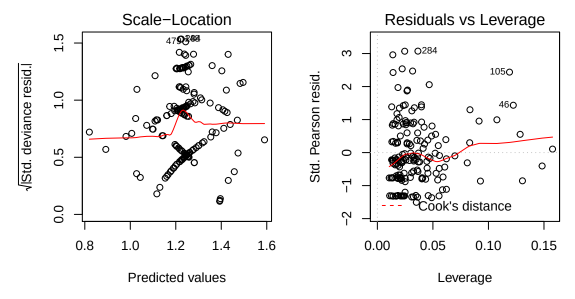
## -1.5050

##

Max

```
##
## Coefficients:
                           Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                          1.264632 1.539616 0.821 0.4126
## Total.no.alleles
                          0.005697 0.019714
                                               0.289 0.7730
## PdistPBR
                          -0.346263 4.627717 -0.075 0.9404
## Sexmale
                          -4.028310 2.322488 -1.734
                                                      0.0847 .
## Total.no.alleles:Sexmale 0.022109 0.031166 0.709 0.4791
## PdistPBR:Sexmale
                   12.009561 6.736310 1.783
                                                      0.0765 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for Negative Binomial(5.1463) family taken to be 0.6098127)
##
##
      Null deviance: 101.638 on 170 degrees of freedom
## Residual deviance: 97.999 on 165 degrees of freedom
## AIC: 708.28
##
## Number of Fisher Scoring iterations: 4
overdisp_fun(fm3_PBR)
        chisq
                    ratio
                                 rdf
                                              p
## 100.6190635
                0.6098125 165.0000000
                                     0.9999804
par(mfrow=c(2,2))
plot(fm3_PBR)
```



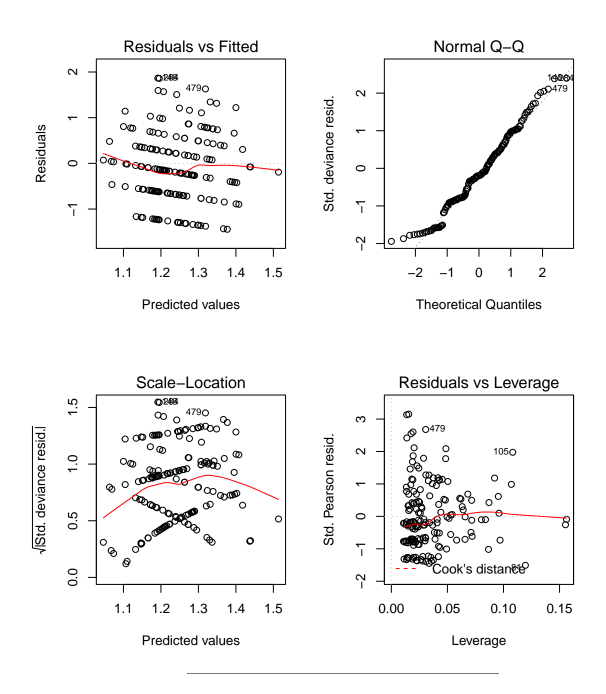


GLM of life span on total number of alleles and P-distance PSS in both sexes

```
fm3_PSS <- glm(LifeSpan~Total.no.alleles+PdistPSS+Sex+Total.no.alleles:Sex+PdistPSS:Sex,negative.binomi
s3_PSS <- summary(fm3_PSS)
s3_PSS</pre>
```

```
##
## Call:
  glm(formula = LifeSpan ~ Total.no.alleles + PdistPSS + Sex +
       Total.no.alleles:Sex + PdistPSS:Sex, family = negative.binomial(theta = 5.14632,
##
##
       link = "log"), data = data[!is.na(PdistPSS) & !is.na(ToSucLife_1014),
##
       ])
##
## Deviance Residuals:
##
       Min
                      Median
                                    3Q
                 1Q
                                            Max
## -1.4432
           -0.6218
                    -0.1523
                                0.3805
                                         1.8661
```

```
##
## Coefficients:
                          Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                          3.179939 1.780045 1.786 0.0759 .
## Total.no.alleles
                          -0.001591 0.019815 -0.080
                                                      0.9361
## PdistPSS
                          -5.357085 4.688965 -1.142 0.2549
## Sexmale
                          -5.492737 2.862678 -1.919 0.0567.
## Total.no.alleles:Sexmale 0.024473 0.032567 0.751
                                                        0.4534
## PdistPSS:Sexmale
                   14.375044 7.249065 1.983
                                                      0.0490 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for Negative Binomial(5.1463) family taken to be 0.6166023)
##
##
      Null deviance: 101.638 on 170 degrees of freedom
## Residual deviance: 99.039 on 165 degrees of freedom
## AIC: 709.32
##
## Number of Fisher Scoring iterations: 4
overdisp_fun(fm3_PSS)
        chisq
                   ratio
                                 rdf
## 101.7392809
                0.6166017 165.0000000
                                     0.9999717
par(mfrow=c(2,2))
plot(fm3_PSS)
```



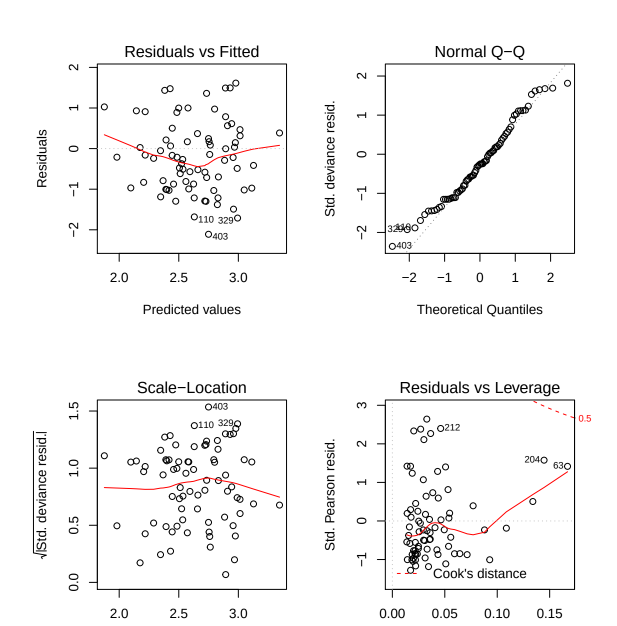
Modeling the effects of MHC-I functional divergence on lifetime number of fledgings

GLM of lifetime number of fledgings on total number of alleles and P-distance PBR in males only

```
fm4_PBR <- glm(ToSucLife_1014~Total.no.alleles+PdistPBR,negative.binomial(theta=1.656555,link="log"),da
s4_PBR <- summary(fm4_PBR)
s4_PBR
##
## Call:</pre>
```

## glm(formula = ToSucLife\_1014 ~ Total.no.alleles + PdistPBR, family = negative.binomial(theta = 1.656

```
link = "log"), data = data.males[!is.na(PdistPBR) & !is.na(ToSucLife_1014),
##
##
      1)
##
## Deviance Residuals:
##
      Min
                1Q
                    Median
                                  3Q
                                          Max
## -2.1104 -0.8979 -0.2296 0.3733
                                       1.6130
## Coefficients:
##
                   Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                   -4.77799
                              2.36035 -2.024 0.04660 *
## Total.no.alleles 0.04216
                               0.03285
                                       1.283 0.20344
## PdistPBR
                   21.95219
                               6.66919
                                        3.292 0.00154 **
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for Negative Binomial(1.6566) family taken to be 0.8162788)
##
##
      Null deviance: 71.194 on 75 degrees of freedom
## Residual deviance: 61.065 on 73 degrees of freedom
## AIC: 548.69
##
## Number of Fisher Scoring iterations: 5
overdisp_fun(fm4_PBR)
##
       chisq
                  ratio
                               rdf
## 59.5882720  0.8162777  73.0000000  0.8708043
par(mfrow=c(2,2))
plot(fm4_PBR)
```



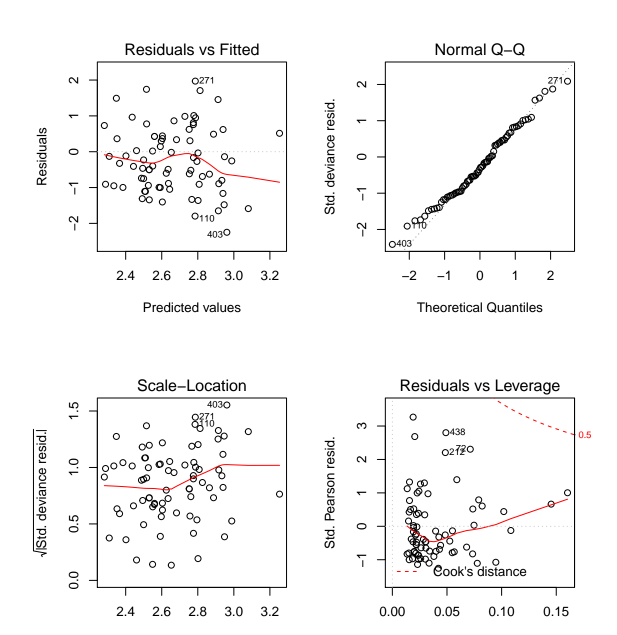
GLM of lifetime number of fledgings on total number of alleles and P-distance PSS in males only

Predicted values

```
fm4_PSS <- glm(ToSucLife_1014~Total.no.alleles+PdistPSS,negative.binomial(theta=1.656555,link="log"),da
s4_PSS <- summary(fm4_PSS)</pre>
s4_PSS
##
   glm(formula = ToSucLife_1014 ~ Total.no.alleles + PdistPSS, family = negative.binomial(theta = 1.656
##
       link = "log"), data = data.males[!is.na(PdistPSS) & !is.na(ToSucLife_1014),
##
       ])
##
## Deviance Residuals:
##
                      Median
       Min
                 1Q
                                    3Q
                                            Max
## -2.2487
            -0.9171
                     -0.2981
                                0.4302
                                          1.9706
```

Leverage

```
##
## Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                 -3.72706 3.21410 -1.160 0.2500
                              0.03699 0.685
## Total.no.alleles 0.02535
                                               0.4953
## PdistPSS
                 16.73262
                            7.94661
                                       2.106 0.0387 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for Negative Binomial(1.6566) family taken to be 0.9075688)
##
      Null deviance: 71.194 on 75 degrees of freedom
##
## Residual deviance: 66.568 on 73 degrees of freedom
## AIC: 554.19
##
## Number of Fisher Scoring iterations: 6
overdisp_fun(fm4_PSS)
##
       chisq
                  ratio
                              rdf
## 66.2526731 0.9075709 73.0000000 0.6988037
par(mfrow=c(2,2))
plot(fm4_PSS)
```



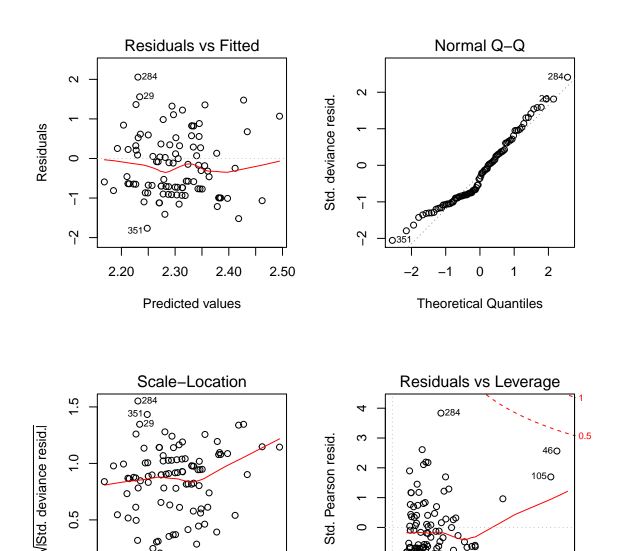
GLM of lifetime number of fledgings on total number of alleles and P-distance PBR in females only

Predicted values

```
fm5_PBR <- glm(ToSucLife_1014~Total.no.alleles+PdistPBR,negative.binomial(theta=1.656555,link="log"),da
s5_PBR <- summary(fm5_PBR)
s5_PBR
##
   glm(formula = ToSucLife_1014 ~ Total.no.alleles + PdistPBR, family = negative.binomial(theta = 1.656
       link = "log"), data = data.females[!is.na(PdistPBR) & !is.na(ToSucLife_1014),
##
##
       ])
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                            Max
## -1.7646
           -0.7670
                     -0.3059
                               0.3335
                                         2.0569
```

Leverage

```
##
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
                  3.19732 2.06239 1.550
## (Intercept)
                                               0.125
                            0.02644 0.447
## Total.no.alleles 0.01181
                                                0.656
## PdistPBR
                 -3.38634 6.19942 -0.546
                                                0.586
## (Dispersion parameter for Negative Binomial(1.6566) family taken to be 0.7562186)
##
##
      Null deviance: 63.653 on 94 degrees of freedom
## Residual deviance: 63.063 on 92 degrees of freedom
## AIC: 618.72
## Number of Fisher Scoring iterations: 6
overdisp_fun(fm5_PBR)
       chisq
                  ratio
                              rdf
## 69.5720925 0.7562184 92.0000000 0.9608471
par(mfrow=c(2,2))
plot(fm5_PBR)
```



GLM of lifetime number of fledgings on total number of alleles and P-distance PSS in females only

2.40

2.50

2.30

Predicted values

0.0

2.20

```
fm5_PSS <- glm(ToSucLife_1014~Total.no.alleles+PdistPSS,negative.binomial(theta=1.656555,link="log"),da
s5_PSS <- summary(fm5_PSS)</pre>
s5_PSS
##
   glm(formula = ToSucLife_1014 ~ Total.no.alleles + PdistPSS, family = negative.binomial(theta = 1.656
       link = "log"), data = data.females[!is.na(PdistPSS) & !is.na(ToSucLife_1014),
##
##
       ])
##
## Deviance Residuals:
##
       Min
                      Median
                 1Q
                                    3Q
                                            Max
## -1.7586
           -0.7825
                     -0.3324
                                0.4298
                                          2.0345
```

0.00

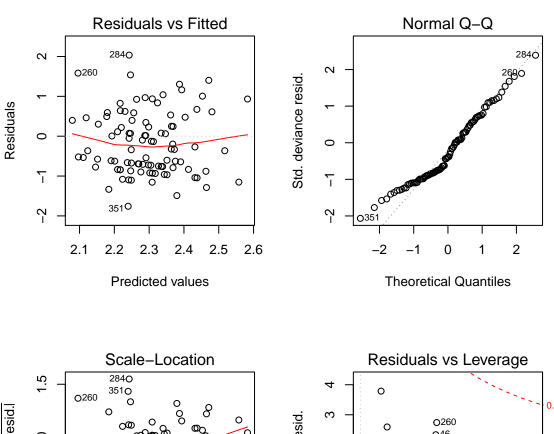
ook's distance

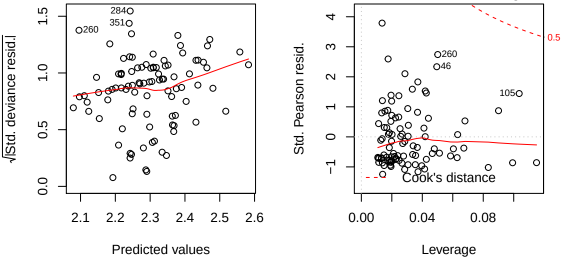
Leverage

0.08

0.12

```
##
## Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                  5.08582 2.35264 2.162 0.0332 *
## Total.no.alleles 0.00407
                              0.02605 0.156
                                               0.8762
## PdistPSS
                 -7.90094
                              6.19254 -1.276 0.2052
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for Negative Binomial(1.6566) family taken to be 0.7343895)
##
      Null deviance: 63.653 on 94 degrees of freedom
##
## Residual deviance: 62.076 on 92 degrees of freedom
## AIC: 617.73
##
## Number of Fisher Scoring iterations: 5
overdisp_fun(fm5_PSS)
##
       chisq
                  ratio
                              rdf
## 67.5638913 0.7343901 92.0000000 0.9738731
par(mfrow=c(2,2))
plot(fm5_PSS)
```

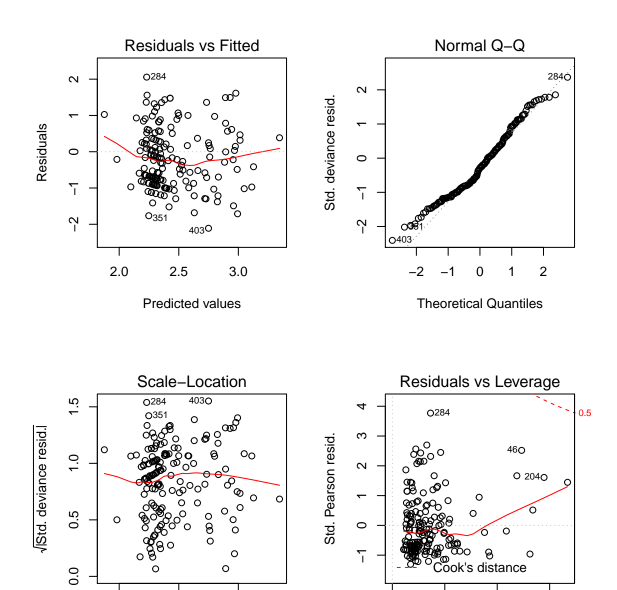




GLM of lifetime number of fledgings on total number of alleles and P-distance PBR in both sexes

```
fm6_PBR <- glm(ToSucLife_1014~Total.no.alleles+PdistPBR+Sex+Total.no.alleles:Sex+PdistPBR:Sex,negative.
s6_PBR <- summary(fm6_PBR)
s6_PBR
##
## Call:
   glm(formula = ToSucLife_1014 ~ Total.no.alleles + PdistPBR +
##
       Sex + Total.no.alleles:Sex + PdistPBR:Sex, family = negative.binomial(theta = 1.656555,
##
##
       link = "log"), data = data[!is.na(PdistPBR) & !is.na(ToSucLife_1014),
       ])
##
##
  Deviance Residuals:
##
       Min
##
                 1Q
                      Median
                                    3Q
                                            Max
```

```
## -2.1104 -0.8215 -0.2475 0.3559
                                    2.0569
##
## Coefficients:
##
                          Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                           3.19732
                                     2.09831 1.524 0.12948
## Total.no.alleles
                          0.01181
                                   0.02690 0.439 0.66129
## PdistPBR
                          -3.38634 6.30739 -0.537 0.59207
## Sexmale
                                     3.12179 -2.555 0.01153 *
                          -7.97518
## Total.no.alleles:Sexmale 0.03035
                                     0.04194 0.724 0.47029
## PdistPBR:Sexmale
                          25.33822
                                     9.07946 2.791 0.00588 **
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for Negative Binomial(1.6566) family taken to be 0.7827894)
##
##
      Null deviance: 143.78 on 170 degrees of freedom
## Residual deviance: 124.13 on 165 degrees of freedom
## AIC: 1167.4
## Number of Fisher Scoring iterations: 6
overdisp_fun(fm6_PBR)
##
        chisq
                                 rdf
                    ratio
                                              р
## 129.1602445
                0.7827894 165.0000000
                                     0.9821231
par(mfrow=c(2,2))
plot(fm6_PBR)
```



GLM of lifetime number of fledgings on total number of alleles and P-distance PSS in both sexes

2.0

2.5

Predicted values

3.0

```
fm6_PSS <- glm(ToSucLife_1014~Total.no.alleles+PdistPSS+Sex+Total.no.alleles:Sex+PdistPSS:Sex,negative.
s6_PSS <- summary(fm6_PSS)
s6_PSS
##
## Call:
   glm(formula = ToSucLife_1014 ~ Total.no.alleles + PdistPSS +
##
       Sex + Total.no.alleles:Sex + PdistPSS:Sex, family = negative.binomial(theta = 1.656555,
##
       link = "log"), data = data[!is.na(PdistPBR) & !is.na(ToSucLife_1014),
##
       ])
##
##
  Deviance Residuals:
##
       Min
                      Median
##
                 1Q
                                    3Q
                                            Max
```

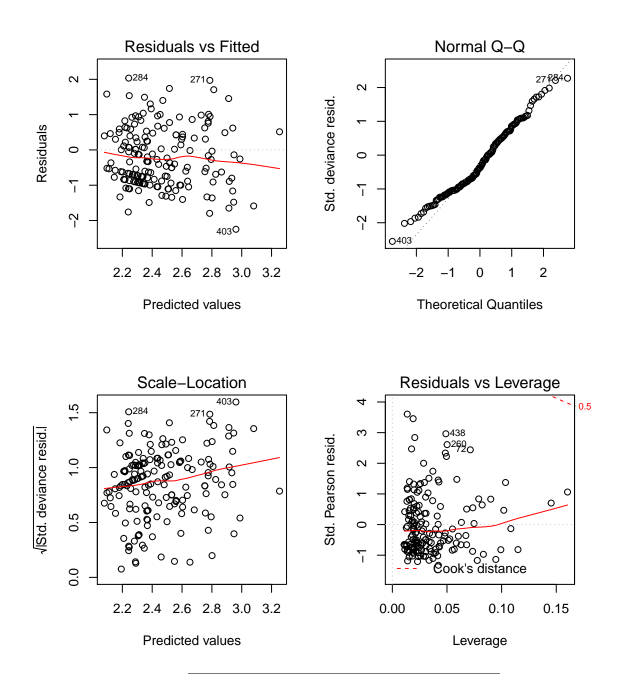
0.00

0.05

0.10

Leverage

```
## -2.2487 -0.8420 -0.3253 0.4339
                                    2.0345
##
## Coefficients:
##
                           Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                           5.085983 2.472325 2.057 0.0412 *
## Total.no.alleles
                          0.004071 0.027371 0.149
                                                      0.8819
## PdistPSS
                          -7.901424 6.507563 -1.214
                                                      0.2264
                                     3.917111 -2.250
## Sexmale
                          -8.813039
                                                      0.0258 *
## Total.no.alleles:Sexmale 0.021282
                                     0.044408 0.479
                                                       0.6324
## PdistPSS:Sexmale
                          24.634040 9.938732 2.479 0.0142 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for Negative Binomial(1.6566) family taken to be 0.8110086)
##
##
      Null deviance: 143.78 on 170 degrees of freedom
## Residual deviance: 128.64 on 165 degrees of freedom
## AIC: 1171.9
## Number of Fisher Scoring iterations: 6
overdisp_fun(fm6_PSS)
##
        chisq
                                 rdf
                   ratio
                                              р
## 133.8165637
               0.8110095 165.0000000
                                     0.9641613
par(mfrow=c(2,2))
plot(fm6_PSS)
```



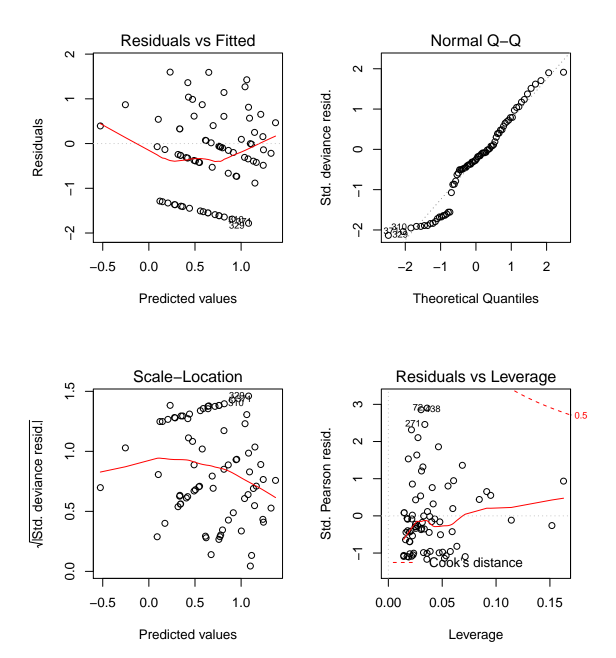
Modeling the effects of MHC-I functional divergence on lifetime number of recruits

GLM of lifetime number of recruits on total number of alleles and P-distance PBR in males only

```
fm7_PBR <- glm(RecruLife_1014~Total.no.alleles+PdistPBR,negative.binomial(theta=1.40551,link="log"),dat
s7_PBR <- summary(fm7_PBR)
s7_PBR

##
## Call:
## glm(formula = RecruLife_1014 ~ Total.no.alleles + PdistPBR, family = negative.binomial(theta = 1.405</pre>
```

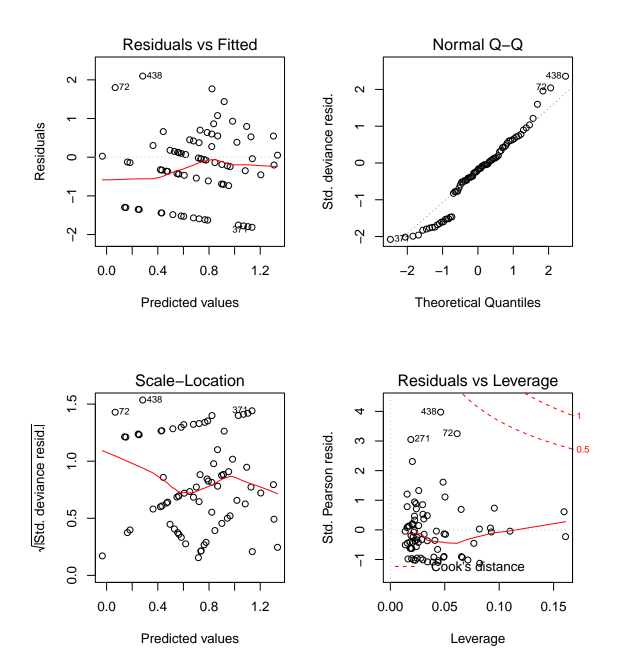
```
link = "log"), data = data.males[!is.na(PdistPBR) & !is.na(ToSucLife_1014),
##
##
      1)
##
## Deviance Residuals:
##
      Min
                1Q
                    Median
                                 3Q
                                         Max
## -1.7826 -0.7714 -0.2091 0.3456
                                       1.5977
## Coefficients:
##
                   Estimate Std. Error t value Pr(>|t|)
                  -9.69571
                               3.11922 -3.108 0.00268 **
## (Intercept)
## Total.no.alleles 0.09970
                               0.04266
                                        2.337 0.02219 *
## PdistPBR
                  28.94496
                               8.73674
                                        3.313 0.00144 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for Negative Binomial(1.4055) family taken to be 0.7232984)
##
##
      Null deviance: 73.051 on 75 degrees of freedom
## Residual deviance: 64.076 on 73 degrees of freedom
## AIC: 290.62
##
## Number of Fisher Scoring iterations: 5
overdisp_fun(fm7_PBR)
##
       chisq
                  ratio
                              rdf
## 52.8009289 0.7233004 73.0000000 0.9640887
par(mfrow=c(2,2))
plot(fm7_PBR)
```



GLM of lifetime number of recruits on total number of alleles and P-distance PSS in males only

```
fm7_PSS <- glm(RecruLife_1014~Total.no.alleles+PdistPSS,negative.binomial(theta=1.40551,link="log"),dat
s7_PSS <- summary(fm7_PSS)</pre>
s7_PSS
##
   glm(formula = RecruLife_1014 ~ Total.no.alleles + PdistPSS, family = negative.binomial(theta = 1.405
       link = "log"), data = data.males[!is.na(PdistPSS) & !is.na(ToSucLife_1014),
##
##
       ])
##
## Deviance Residuals:
##
       Min
                      Median
                 1Q
                                    3Q
                                            Max
## -1.8120
            -0.7114
                     -0.1635
                                0.3208
                                          2.0971
```

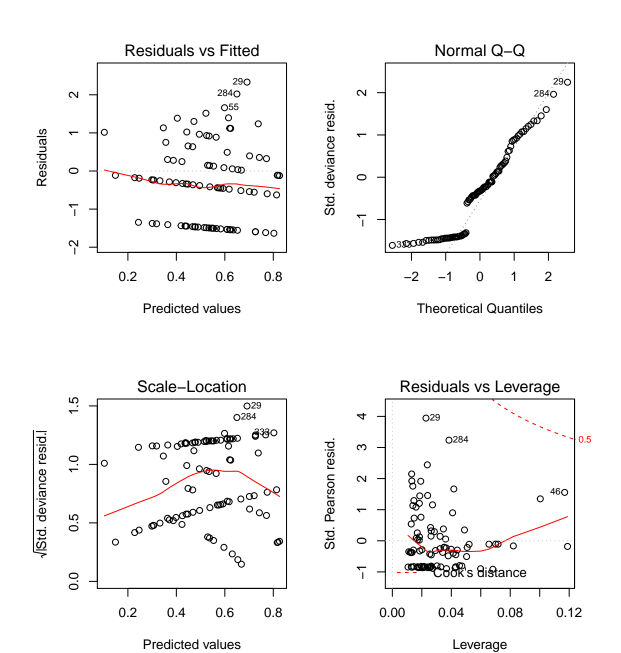
```
##
## Coefficients:
                   Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) -10.26910 4.20981 -2.439 0.0171 *
                                        1.942
## Total.no.alleles 0.09372
                               0.04826
                                                0.0560 .
## PdistPSS
                   26.86065 10.33874 2.598 0.0113 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for Negative Binomial(1.4055) family taken to be 0.8291243)
##
      Null deviance: 73.051 on 75 degrees of freedom
## Residual deviance: 67.128 on 73 degrees of freedom
## AIC: 293.67
##
## Number of Fisher Scoring iterations: 5
overdisp_fun(fm7_PSS)
##
       chisq
                  ratio
                              rdf
## 60.5258756  0.8291216  73.0000000  0.8512463
par(mfrow=c(2,2))
plot(fm7_PSS)
```



GLM of lifetime number of recruits on total number of alleles and P-distance PBR in females only

```
fm8_PBR <- glm(RecruLife_1014~Total.no.alleles+PdistPBR,negative.binomial(theta=1.40551,link="log"),dat
s8_PBR <- summary(fm8_PBR)
s8_PBR
##
   glm(formula = RecruLife_1014 ~ Total.no.alleles + PdistPBR, family = negative.binomial(theta = 1.405
##
       link = "log"), data = data.females[!is.na(PdistPBR) & !is.na(ToSucLife_1014),
##
       ])
##
## Deviance Residuals:
##
       Min
                      Median
                 1Q
                                    3Q
                                            Max
## -1.6347
           -1.4625
                     -0.3396
                               0.3135
                                         2.3327
```

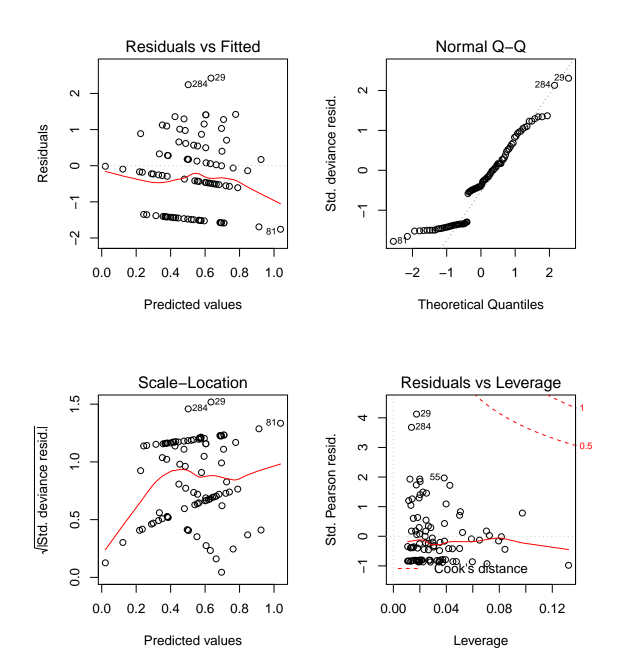
```
##
## Coefficients:
##
                   Estimate Std. Error t value Pr(>|t|)
                 -0.53598 3.44587 -0.156
## (Intercept)
                                               0.877
                              0.04371 -0.952
## Total.no.alleles -0.04161
                                                 0.344
## PdistPBR
                    5.22055 10.36608 0.504
                                                 0.616
## (Dispersion parameter for Negative Binomial(1.4055) family taken to be 1.102901)
##
##
      Null deviance: 113.2 on 94 degrees of freedom
## Residual deviance: 111.5 on 92 degrees of freedom
## AIC: 345.37
## Number of Fisher Scoring iterations: 5
overdisp_fun(fm8_PBR)
        chisq
                    ratio
                                  rdf
## 101.4667300
                1.1028992 92.0000000
                                      0.2345127
par(mfrow=c(2,2))
plot(fm8_PBR)
```



GLM of lifetime number of recruits on total number of alleles and P-distance PSS in females only

```
fm8_PSS <- glm(RecruLife_1014~Total.no.alleles+PdistPSS,negative.binomial(theta=1.40551,link="log"),dat
s8_PSS <- summary(fm8_PSS)</pre>
s8_PSS
##
   glm(formula = RecruLife_1014 ~ Total.no.alleles + PdistPSS, family = negative.binomial(theta = 1.405
       link = "log"), data = data.females[!is.na(PdistPSS) & !is.na(ToSucLife_1014),
##
##
       ])
##
## Deviance Residuals:
##
       Min
                      Median
                 1Q
                                    3Q
                                            Max
## -1.7596
           -1.4213
                     -0.4168
                                0.3066
                                          2.4245
```

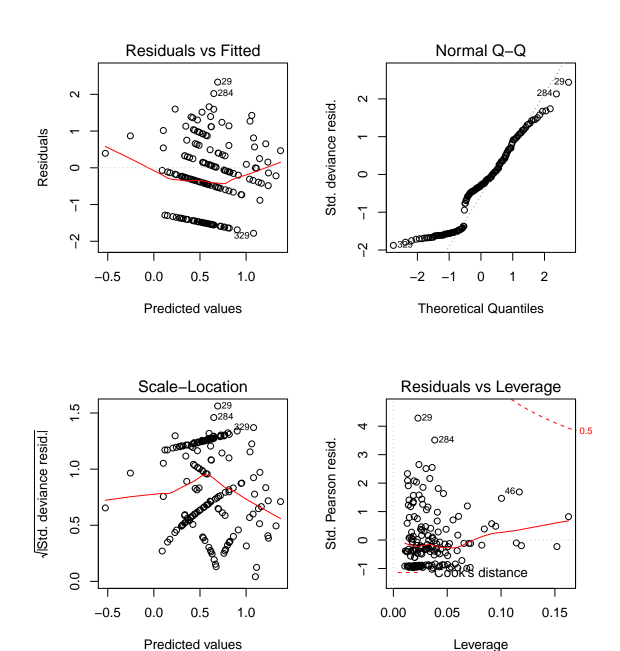
```
##
## Coefficients:
##
                   Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                  4.10393 3.92662 1.045
                                               0.299
## Total.no.alleles -0.06025
                             0.04425 -1.362
                                                 0.177
## PdistPSS
                   -7.68950 10.33549 -0.744
                                                 0.459
## (Dispersion parameter for Negative Binomial(1.4055) family taken to be 1.125134)
##
##
      Null deviance: 113.20 on 94 degrees of freedom
## Residual deviance: 111.23 on 92 degrees of freedom
## AIC: 345.1
## Number of Fisher Scoring iterations: 5
overdisp_fun(fm8_PSS)
        chisq
                    ratio
                                  rdf
                                               p
## 103.5124441
                1.1251353 92.0000000
                                      0.1936933
par(mfrow=c(2,2))
plot(fm8_PSS)
```



GLM of lifetime number of recruits on total number of alleles and P-distance PBR in both sexes

```
fm9_PBR <- glm(RecruLife_1014~Total.no.alleles+PdistPBR+Sex+Total.no.alleles:Sex+PdistPBR:Sex,negative.
s9_PBR <- summary(fm9_PBR)</pre>
s9_PBR
##
## Call:
   glm(formula = RecruLife_1014 ~ Total.no.alleles + PdistPBR +
##
       Sex + Total.no.alleles:Sex + PdistPBR:Sex, family = negative.binomial(theta = 1.40551,
##
##
       link = "log"), data = data[!is.na(PdistPBR) & !is.na(ToSucLife_1014),
       ])
##
##
  Deviance Residuals:
##
##
       Min
                 1Q
                      Median
                                    3Q
                                            Max
```

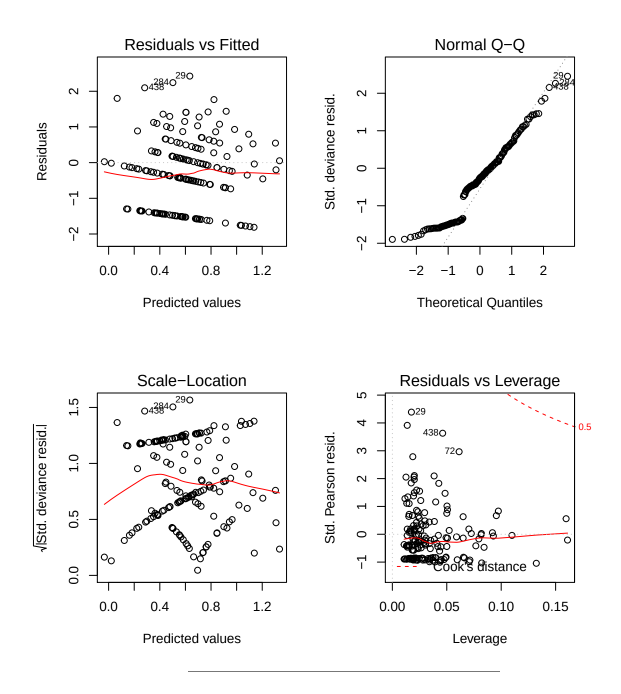
```
## -1.7826 -1.3926 -0.2654 0.3280
                                    2.3327
##
## Coefficients:
##
                          Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                          -0.53598
                                     3.17268 -0.169
                                                       0.8661
## Total.no.alleles
                          -0.04161
                                   0.04024 -1.034
                                                      0.3026
## PdistPBR
                          5.22055 9.54425 0.547
                                                       0.5851
                                     4.75842 -1.925
## Sexmale
                          -9.15973
                                                       0.0560 .
## Total.no.alleles:Sexmale 0.14132
                                   0.06303 2.242
                                                       0.0263 *
## PdistPBR:Sexmale
                          23.72440 13.77533 1.722
                                                      0.0869 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for Negative Binomial(1.4055) family taken to be 0.9349556)
##
##
      Null deviance: 187.69 on 170 degrees of freedom
## Residual deviance: 175.58 on 165 degrees of freedom
## AIC: 635.99
## Number of Fisher Scoring iterations: 5
overdisp_fun(fm9_PBR)
##
        chisq
                                 rdf
                    ratio
## 154.2676589
                0.9349555 165.0000000
                                     0.7147116
par(mfrow=c(2,2))
plot(fm9_PBR)
```



GLM of lifetime number of recruits on total number of alleles and P-distance PSS in both sexes

```
fm9_PSS <- glm(RecruLife_1014~Total.no.alleles+PdistPSS+Sex+Total.no.alleles:Sex+PdistPSS:Sex,negative.
s9_PSS <- summary(fm9_PSS)</pre>
s9_PSS
##
## Call:
   glm(formula = RecruLife_1014 ~ Total.no.alleles + PdistPSS +
##
       Sex + Total.no.alleles:Sex + PdistPSS:Sex, family = negative.binomial(theta = 1.40551,
##
##
       link = "log"), data = data[!is.na(PdistPSS) & !is.na(ToSucLife_1014),
       ])
##
##
##
  Deviance Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                            Max
```

```
## -1.8120 -1.3934 -0.2542 0.3154 2.4245
##
## Coefficients:
##
                           Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                           4.10393
                                      3.69103 1.112 0.2678
## Total.no.alleles
                           -0.06025
                                     0.04159 -1.449
                                                      0.1493
## PdistPSS
                           -7.68950
                                    9.71538 -0.791
                                                      0.4298
                                     5.90543 -2.434
## Sexmale
                          -14.37303
                                                       0.0160 *
## Total.no.alleles:Sexmale
                           0.15397
                                     0.06725 2.290
                                                       0.0233 *
## PdistPSS:Sexmale
                           34.55016 14.91831 2.316
                                                      0.0218 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for Negative Binomial(1.4055) family taken to be 0.9941722)
##
##
      Null deviance: 187.69 on 170 degrees of freedom
## Residual deviance: 178.36 on 165 degrees of freedom
## AIC: 638.77
## Number of Fisher Scoring iterations: 5
overdisp_fun(fm9_PSS)
##
        chisq
                                 rdf
                    ratio
                                              р
## 164.0383197
                0.9941716 165.0000000
                                     0.5065083
par(mfrow=c(2,2))
plot(fm9_PSS)
```



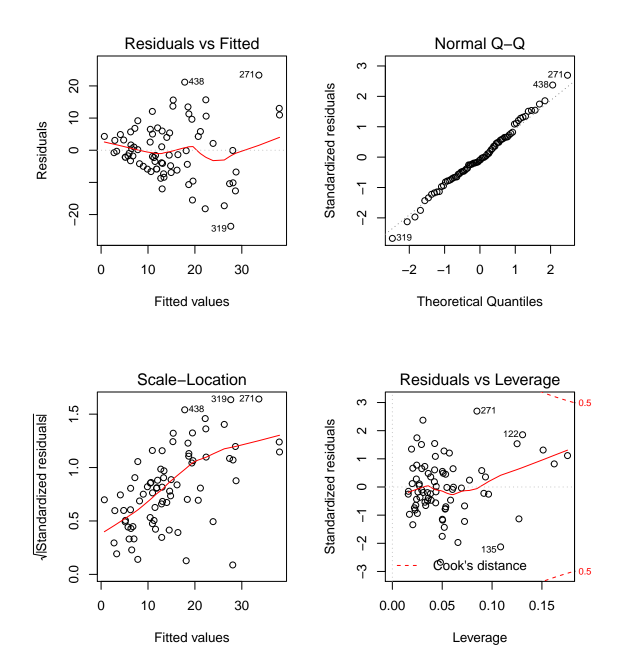
## Modeling the effects of MHC-I functional divergence on offspring fledging success

Linear model of lifetime number of fledgings on total number of alleles and P-distance PBR with lifespan as covariate in males only

```
fm10_PBR <- lm(ToSucLife_1014~LifeSpan+Total.no.alleles+PdistPBR,data=data.males[!is.na(PdistPBR)&!is.n
s10_PBR <- summary(fm10_PBR)
s10_PBR
##
## Call:</pre>
```

## lm(formula = ToSucLife\_1014 ~ LifeSpan + Total.no.alleles + PdistPBR,
## data = data.males[!is.na(PdistPBR) & !is.na(ToSucLife\_1014),

```
##
           ])
##
## Residuals:
##
       Min
                 1Q Median
                                    3Q
                                           Max
## -23.6438 -5.8328 -0.8704 5.4632 23.3595
##
## Coefficients:
                   Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                    -52.5044
                               29.4221 -1.785
                                                 0.0786 .
                                0.7195
## LifeSpan
                     4.7575
                                          6.612 5.68e-09 ***
## Total.no.alleles
                     0.2099
                                0.4034
                                          0.520
                                                  0.6044
## PdistPBR
                   152.3550
                               85.6042
                                          1.780
                                                  0.0793 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
\mbox{\tt \#\#} Residual standard error: 9.061 on 72 degrees of freedom
## Multiple R-squared: 0.4644, Adjusted R-squared: 0.4421
## F-statistic: 20.81 on 3 and 72 DF, p-value: 8.194e-10
par(mfrow=c(2,2))
plot(fm10_PBR)
```



Linear model of lifetime number of fledgings on total number of alleles and P-distance PSS with lifespan as covariate in males only

24.1169

Max

ЗQ

5.1370

##

##

## Residuals:

-23.6097

Min

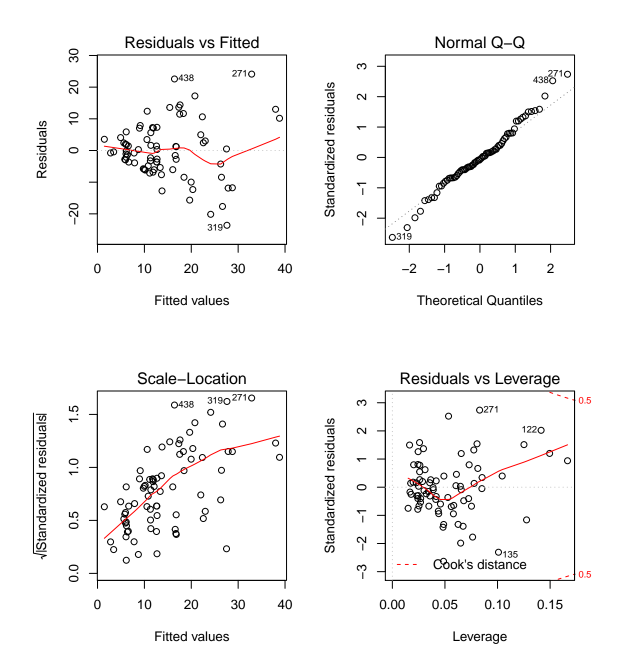
Median

-0.6145

1Q

-5.4712

```
##
## Coefficients:
##
                  Estimate Std. Error t value Pr(>|t|)
                -38.09164 38.25115 -0.996
## (Intercept)
                                               0.323
                            0.70723 7.107
## LifeSpan
                                                7e-10 ***
                   5.02654
## Total.no.alleles 0.07344 0.43684 0.168
                                                0.867
## PdistPSS
                   94.07153 96.00763 0.980
                                                0.330
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 9.198 on 72 degrees of freedom
## Multiple R-squared: 0.4482, Adjusted R-squared: 0.4252
## F-statistic: 19.49 on 3 and 72 DF, p-value: 2.356e-09
par(mfrow=c(2,2))
plot(fm10_PSS)
```



Linear model of lifetime number of fledgings on total number of alleles and P-distance PBR with lifespan as covariate in females only

```
fm11_PBR <- lm(ToSucLife_1014~LifeSpan+Total.no.alleles+PdistPBR,data=data.females[!is.na(PdistPBR)&!is
s11_PBR <- summary(fm11_PBR)
s11_PBR

##
## Call:
## lm(formula = ToSucLife_1014 ~ LifeSpan + Total.no.alleles + PdistPBR,
## data = data.females[!is.na(PdistPBR) & !is.na(ToSucLife_1014),
## ])</pre>
```

12.2748

Max

ЗQ

1.6074

##

##

## Residuals:

Min

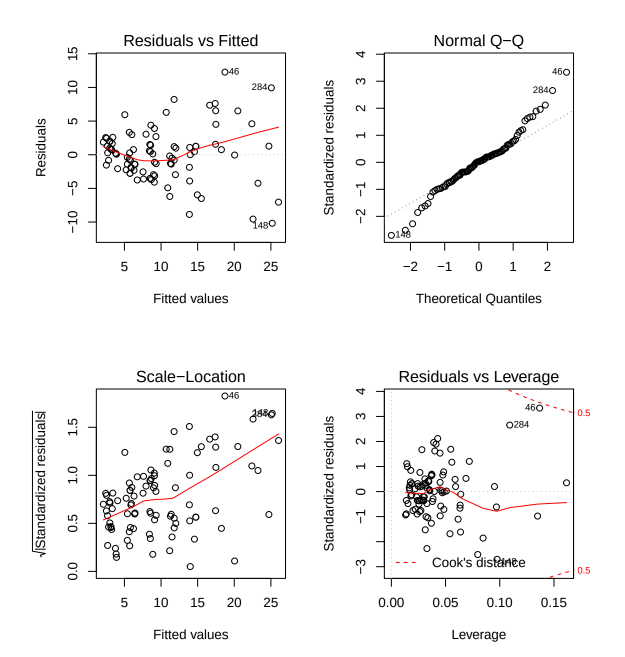
## -10.1845 -2.1607

Median

0.1229

1Q

```
##
## Coefficients:
##
                   Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                   11.11888 11.25593 0.988
                                               0.326
## LifeSpan
                            0.19055 14.901
                                              <2e-16 ***
                    2.83950
## Total.no.alleles 0.06114
                            0.14379 0.425
                                              0.672
## PdistPBR -37.50437 33.77333 -1.110
                                                0.270
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 3.965 on 91 degrees of freedom
## Multiple R-squared: 0.7121, Adjusted R-squared: 0.7026
## F-statistic: 75.02 on 3 and 91 DF, \, p-value: < 2.2e-16
par(mfrow=c(2,2))
plot(fm11_PBR)
```



Linear model of lifetime number of fledgings on total number of alleles and P-distance PSS with lifespan as covariate in females only

```
fm11_PSS <- lm(ToSucLife_1014~LifeSpan+Total.no.alleles+PdistPSS,data=data.females[!is.na(PdistPSS)&!is
s11_PSS <- summary(fm11_PSS)
s11_PSS

##
## Call:
## Call:
## lm(formula = ToSucLife_1014 ~ LifeSpan + Total.no.alleles + PdistPSS,
## data = data.females[!is.na(PdistPSS) & !is.na(ToSucLife_1014),
## ])
##</pre>
```

13.0347

Max

ЗQ

1.6705

## Residuals:

Min

## -10.3651 -2.0775

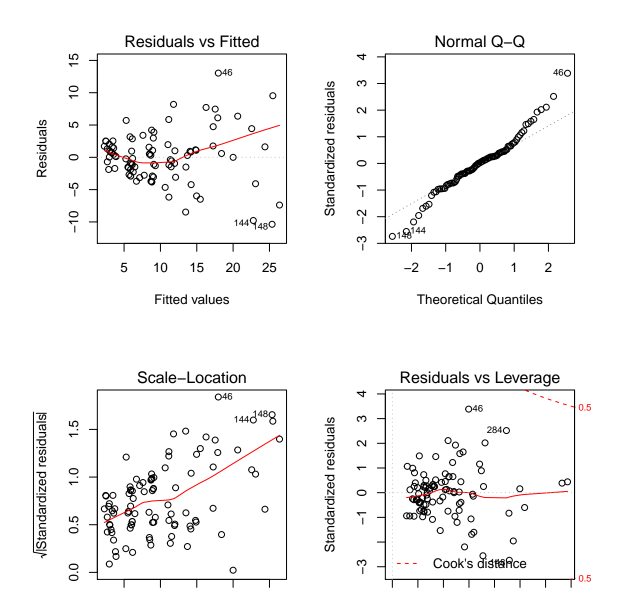
Median

0.1081

1Q

##

```
##
## Coefficients:
##
                   Estimate Std. Error t value Pr(>|t|)
                   11.40047 13.17409 0.865
                                                0.389
## (Intercept)
                             0.19194 14.700
## LifeSpan
                                               <2e-16 ***
                    2.82151
## Total.no.alleles 0.06757
                             0.14389 0.470
                                                0.640
## PdistPSS -33.30141 34.45899 -0.966
                                                0.336
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 3.971 on 91 degrees of freedom
## Multiple R-squared: 0.7111, Adjusted R-squared: 0.7016
## F-statistic: 74.68 on 3 and 91 DF, p-value: < 2.2e-16
par(mfrow=c(2,2))
plot(fm11_PSS)
```



Linear model of lifetime number of fledgings on total number of alleles and P-distance PBR with lifespan as covariate in both sexes

5

##

##

## Residuals:

-20.6629

Min

10

20

25

15

Fitted values

Median

-0.4787

1Q

-3.5429

```
fm12_PBR <- lm(ToSucLife_1014~LifeSpan+Total.no.alleles+PdistPBR+Sex+Total.no.alleles:Sex+PdistPBR:Sex,
s12_PBR <- summary(fm12_PBR)
s12_PBR

##
## Call:
## Call:
## lm(formula = ToSucLife_1014 ~ LifeSpan + Total.no.alleles + PdistPBR +
## Sex + Total.no.alleles:Sex + PdistPBR:Sex, data = data[!is.na(PdistPBR) &
## !is.na(ToSucLife_1014), ])</pre>
```

0.00

0.04

0.08

Leverage

0.12

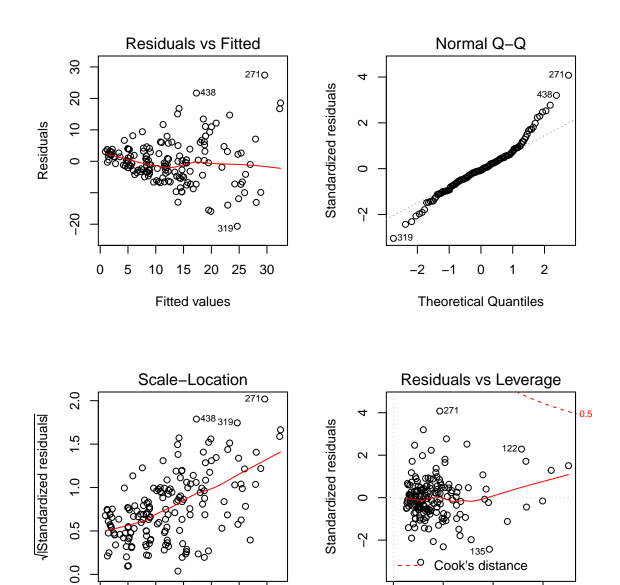
27.4032

Max

3Q

3.1341

```
##
## Coefficients:
##
                          Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                          9.26681 19.53120 0.474 0.6358
                                   0.28300 11.851
                                                    <2e-16 ***
## LifeSpan
                           3.35383
## Total.no.alleles
                          0.05105 0.24961 0.205 0.8382
## PdistPBR
                         -36.79178 58.63195 -0.628 0.5312
                         -76.15279
                                    29.39188 -2.591 0.0104 *
## Sexmale
## Total.no.alleles:Sexmale 0.29404 0.39225 0.750 0.4546
## PdistPBR:Sexmale
                         245.28201 85.58624 2.866 0.0047 **
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 6.883 on 164 degrees of freedom
## Multiple R-squared: 0.5405, Adjusted R-squared: 0.5236
## F-statistic: 32.15 on 6 and 164 DF, p-value: < 2.2e-16
par(mfrow=c(2,2))
plot(fm12_PBR)
```



Linear model of lifetime number of fledgings on total number of alleles and P-distance PSS with lifespan as covariate in both sexes

0

## Residuals:

Min

## -20.0382 -3.5805

##

5

10

20

Median

-0.2141

1Q

15

Fitted values

25

30

```
fm12_PSS <- lm(ToSucLife_1014~LifeSpan+Total.no.alleles+PdistPSS+Sex+Total.no.alleles:Sex+PdistPSS:Sex,
s12_PSS <- summary(fm12_PSS)
s12_PSS

##
## Call:
## lm(formula = ToSucLife_1014 ~ LifeSpan + Total.no.alleles + PdistPSS +
## Sex + Total.no.alleles:Sex + PdistPSS:Sex, data = data[!is.na(PdistPSS) &
## !is.na(ToSucLife_1014), ])
##</pre>
```

0.00

0.05

0.10

Leverage

0.15

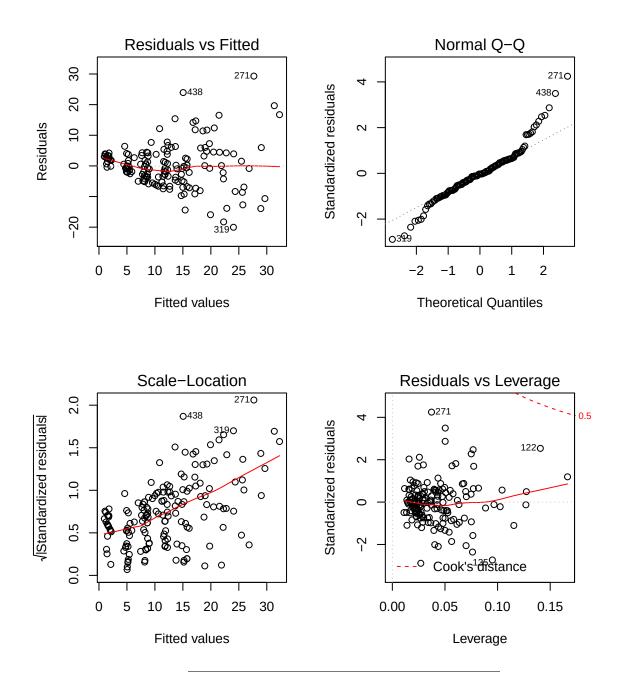
29.3120

Max

3Q

3.2492

```
##
## Coefficients:
##
                         Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                         4.96890 23.26213 0.214 0.831
                                   0.28785 11.971
## LifeSpan
                                                   <2e-16 ***
                          3.44599
## Total.no.alleles
                         0.06869 0.25489 0.269 0.788
## PdistPSS
                        -21.42623 60.94379 -0.352 0.726
## Sexmale
                                   37.35539 -1.532
                         -57.24369
                                                   0.127
                                                   0.755
## Total.no.alleles:Sexmale 0.13088
                                  0.41867 0.313
## PdistPSS:Sexmale 165.45598 94.84718 1.744
                                                   0.083 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 7.034 on 164 degrees of freedom
## Multiple R-squared: 0.52, Adjusted R-squared: 0.5024
## F-statistic: 29.61 on 6 and 164 DF, p-value: < 2.2e-16
par(mfrow=c(2,2))
plot(fm12_PSS)
```



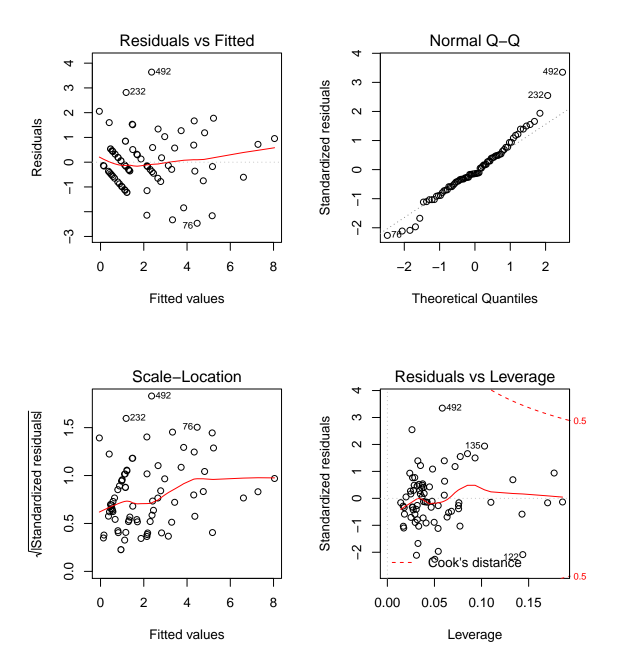
Modeling the effects of MHC-I functional divergence on offspring recruitment success

Linear model of lifetime number of recruits on total number of alleles and P-distance PBR with lifetime number of fledglings as covariate in males only

```
fm13_PBR <- lm(RecruLife_1014~ToSucLife_1014+Total.no.alleles+PdistPBR,data=data.males[!is.na(PdistPBR)]
s13_PBR <- summary(fm13_PBR)
s13_PBR</pre>
```

```
##
## Call:
## lm(formula = RecruLife_1014 ~ ToSucLife_1014 + Total.no.alleles +
```

```
PdistPBR, data = data.males[!is.na(PdistPBR) & !is.na(ToSucLife_1014),
##
##
      ])
##
## Residuals:
##
               1Q Median
                               3Q
                                      Max
## -2.4679 -0.6180 -0.1508 0.5467 3.6382
## Coefficients:
##
                   Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                   -5.48675
                               3.70735 -1.480
                                                 0.1432
## ToSucLife_1014
                    0.13848
                               0.01149 12.054
                                                 <2e-16 ***
## Total.no.alleles 0.11761
                               0.04971
                                         2.366
                                                 0.0207 *
                                         1.195
## PdistPBR
                   12.80035
                              10.71313
                                                 0.2361
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.12 on 72 degrees of freedom
## Multiple R-squared: 0.7178, Adjusted R-squared: 0.706
## F-statistic: 61.04 on 3 and 72 DF, p-value: < 2.2e-16
par(mfrow=c(2,2))
plot(fm13_PBR)
```



Linear model of lifetime number of recruits on total number of alleles and P-distance PSS with lifetime number of fledglings as covariate in males only

```
fm13_PSS <- lm(RecruLife_1014~ToSucLife_1014+Total.no.alleles+PdistPSS,data=data.males[!is.na(PdistPSS)]
s13_PSS <- summary(fm13_PSS)
s13_PSS

##
## Call:
## lm(formula = RecruLife_1014 ~ ToSucLife_1014 + Total.no.alleles +
## PdistPSS, data = data.males[!is.na(PdistPSS) & !is.na(ToSucLife_1014),
## ])
##
## Residuals:</pre>
```

Max

##

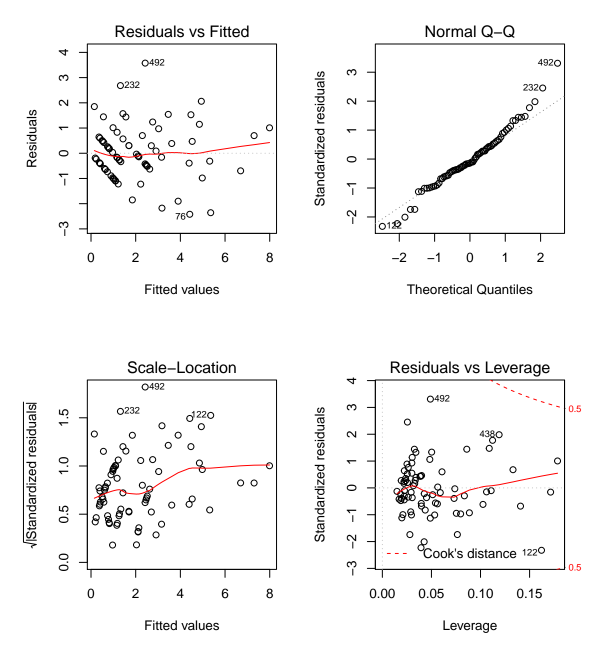
Min

1Q Median

## -2.4205 -0.6302 -0.1484 0.5762

3Q

```
##
## Coefficients:
##
                  Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                  -9.13968 4.63216 -1.973 0.05232 .
                           0.01088 12.775 < 2e-16 ***
## ToSucLife_1014 0.13900
## Total.no.alleles 0.14205
                           0.05242 2.710 0.00841 **
## PdistPSS
                20.22049
                           11.57660 1.747 0.08496 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.108 on 72 degrees of freedom
## Multiple R-squared: 0.7239, Adjusted R-squared: 0.7124
## F-statistic: 62.92 on 3 and 72 DF, p-value: < 2.2e-16
par(mfrow=c(2,2))
plot(fm13_PSS)
```



Linear model of lifetime number of recruits on total number of alleles and P-distance PBR with lifetime number of fledglings as covariate in females only

```
fm14_PBR <- lm(RecruLife_1014~ToSucLife_1014+Total.no.alleles+PdistPBR,data=data.females[!is.na(PdistPBE
s14_PBR <- summary(fm14_PBR)
s14_PBR

##
## Call:
## lm(formula = RecruLife_1014 ~ ToSucLife_1014 + Total.no.alleles +
## PdistPBR, data = data.females[!is.na(PdistPBR) & !is.na(ToSucLife_1014),
## ])
##</pre>
```

Max

5.2017

## Residuals:

Min

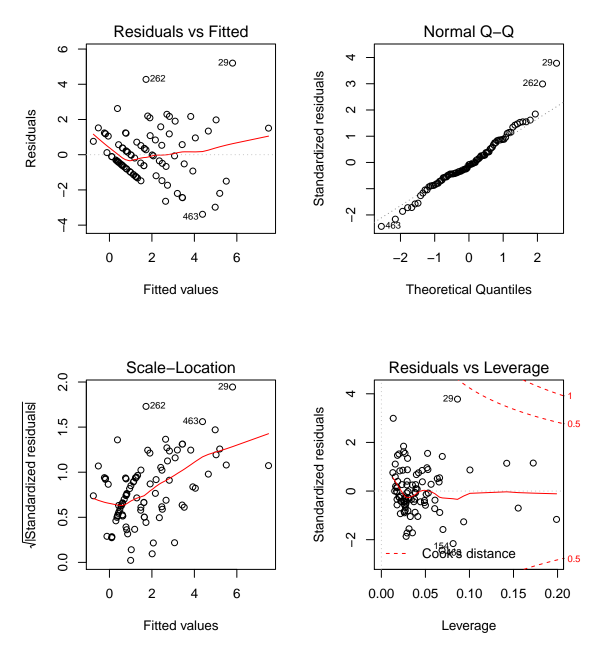
1Q Median

## -3.3836 -0.7834 -0.1138 0.7974

3Q

##

```
##
## Coefficients:
##
                  Estimate Std. Error t value Pr(>|t|)
                  -5.31908 4.10214 -1.297
## (Intercept)
                                               0.198
                            0.02052 10.446
## ToSucLife_1014 0.21433
                                              <2e-16 ***
## Total.no.alleles -0.08760
                            0.05224 -1.677
                                                0.097 .
## PdistPBR
                 19.49656
                            12.28996
                                      1.586
                                                0.116
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.439 on 91 degrees of freedom
## Multiple R-squared: 0.5521, Adjusted R-squared: 0.5373
## F-statistic: 37.39 on 3 and 91 DF, p-value: 7.743e-16
par(mfrow=c(2,2))
plot(fm14_PBR)
```



Linear model of lifetime number of recruits on total number of alleles and P-distance PSS with lifetime number of fledglings as covariate in females only

```
fm14_PSS <- lm(RecruLife_1014~ToSucLife_1014+Total.no.alleles+PdistPSS,data=data.females[!is.na(PdistPS
s14_PSS <- summary(fm14_PSS)
s14_PSS

##
## Call:
## lm(formula = RecruLife_1014 ~ ToSucLife_1014 + Total.no.alleles +
## PdistPSS, data = data.females[!is.na(PdistPSS) & !is.na(ToSucLife_1014),
## ])
##</pre>
```

Max

5.3625

## Residuals:

Min

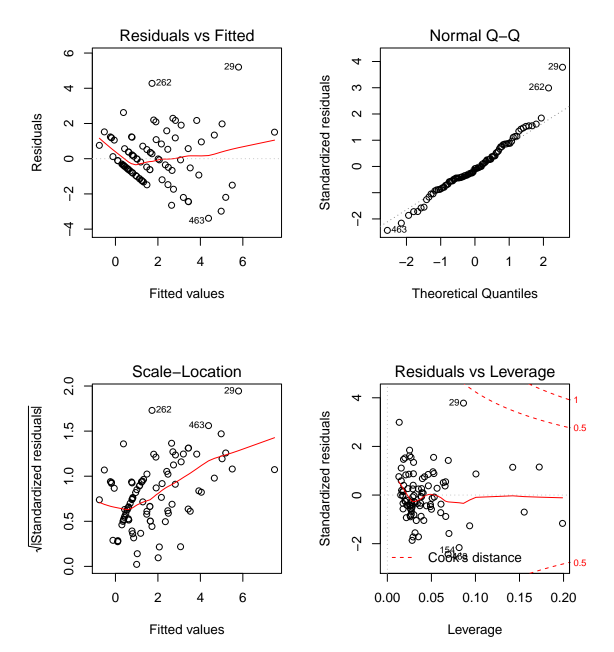
1Q Median

## -3.5868 -0.7087 -0.1396 0.7950

3Q

##

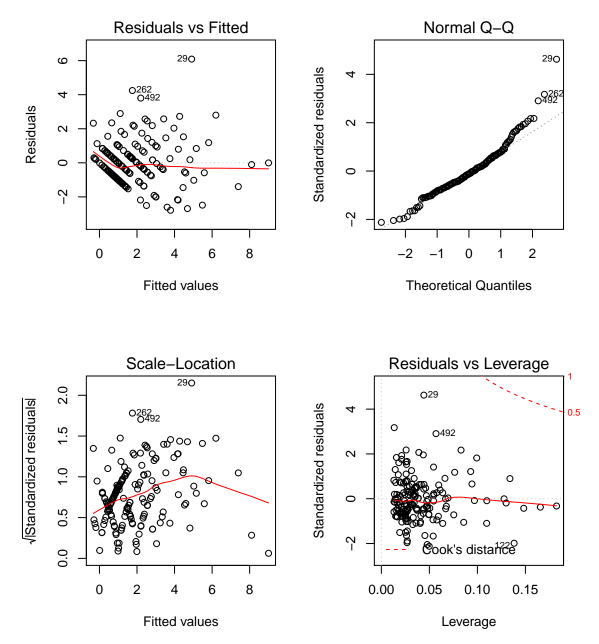
```
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
                -1.52332 4.85235 -0.314 0.7543
## (Intercept)
## ToSucLife_1014 0.21371
                         0.02093 10.210
                                          <2e-16 ***
0.0535 .
## PdistPSS
                 6.92181 12.70041
                                  0.545
                                          0.5871
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.457 on 91 degrees of freedom
## Multiple R-squared: 0.5412, Adjusted R-squared: 0.5261
## F-statistic: 35.78 on 3 and 91 DF, p-value: 2.288e-15
par(mfrow=c(2,2))
plot(fm14_PBR)
```



Linear model of lifetime number of recruits on total number of alleles and P-distance PBR with lifetime number of fledglings as covariate in both sexes

```
fm15_PBR <- lm(RecruLife_1014~ToSucLife_1014+Total.no.alleles+PdistPBR+Sex+Total.no.alleles:Sex+PdistPB
s15_PBR <- summary(fm15_PBR)</pre>
s15_PBR
##
  lm(formula = RecruLife_1014 ~ ToSucLife_1014 + Total.no.alleles +
##
       PdistPBR + Sex + Total.no.alleles:Sex + PdistPBR:Sex, data = data[!is.na(PdistPBR) &
##
##
       !is.na(ToSucLife_1014), ])
##
## Residuals:
##
       Min
                1Q Median
                                 3Q
                                        Max
## -2.7865 -0.8432 -0.1574 0.6730
                                    6.0886
```

```
##
## Coefficients:
                         Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                         -4.25247 3.82396 -1.112 0.2677
                                   0.01121 14.656 <2e-16 ***
## ToSucLife_1014
                          0.16436
## Total.no.alleles
                         -0.08175 0.04885 -1.674 0.0961 .
## PdistPBR
                         17.42576 11.48167 1.518 0.1310
## Sexmale
                          1.38591
                                   5.86322 0.236 0.8134
## Total.no.alleles:Sexmale 0.18208
                                   0.07688 2.368 0.0190 *
## PdistPBR:Sexmale -13.49193 17.13914 -0.787 0.4323
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.347 on 164 degrees of freedom
## Multiple R-squared: 0.602, Adjusted R-squared: 0.5874
## F-statistic: 41.34 on 6 and 164 DF, p-value: < 2.2e-16
par(mfrow=c(2,2))
plot(fm15_PBR)
```



Linear model of lifetime number of recruits on total number of alleles and P-distance PSS with lifetime number of fledglings as covariate in both sexes

```
fm15_PSS <- lm(RecruLife_1014~ToSucLife_1014+Total.no.alleles+PdistPSS+Sex+Total.no.alleles:Sex+PdistPS
s15_PSS <- summary(fm15_PSS)</pre>
s15_PSS
##
  lm(formula = RecruLife_1014 ~ ToSucLife_1014 + Total.no.alleles +
##
       PdistPSS + Sex + Total.no.alleles:Sex + PdistPSS:Sex, data = data[!is.na(PdistPSS) &
##
##
       !is.na(ToSucLife_1014), ])
##
## Residuals:
##
       Min
                1Q Median
                                 3Q
                                        Max
## -2.9970 -0.8266 -0.1215 0.6371
                                    6.2583
```

```
##
## Coefficients:
                         Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                         0.53677 4.45724 0.120 0.90429
                                  0.01097 14.846 < 2e-16 ***
## ToSucLife_1014
                          0.16279
## Total.no.alleles
                        -0.10010 0.04900 -2.043 0.04268 *
## PdistPSS
                         2.49426 11.70647 0.213 0.83154
                         -7.69698 7.22931 -1.065 0.28858
## Sexmale
## Total.no.alleles:Sexmale 0.23085 0.08048 2.869 0.00467 **
## PdistPSS:Sexmale 11.70793 18.39705 0.636 0.52540
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.352 on 164 degrees of freedom
## Multiple R-squared: 0.5987, Adjusted R-squared: 0.5841
## F-statistic: 40.79 on 6 and 164 DF, p-value: < 2.2e-16
par(mfrow=c(2,2))
plot(fm15_PSS)
```

