GLMM models of the effects of MHC-I functional divergence on offspring recruitment in great reed warblers (Acrocephalus arundinaceus)

Jacob Roved 5/9/2019

This document presents the R code and output from the GLMM models behind the results on the effects of MHC-I functional divergence on offspring recruitment in 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:

```
Nest = Nest ID
Recruitment = Recruited
Total.no.alleles = No. MHC-I alleles
PdistPBR = P-distance PBR
PdistPSS = P-distance PSS
```

Rescale the variables PdistPBR, PdistPSS, and Total.no.alleles to avoid convergence issue in glmer.

```
dataMHC[,c(13,15,16)] <- scale(dataMHC[,c(13,15,16)],center=T,scale=T)
dataMHC.males <- subset(dataMHC,dataMHC$Sex==1)
dataMHC.females <- subset(dataMHC,dataMHC$Sex==2)
attach(dataMHC)</pre>
```

Load R packages.

```
library(lme4)

## Warning: package 'lme4' was built under R version 3.4.4

## Loading required package: Matrix

library(arm)

## Warning: package 'arm' was built under R version 3.4.4

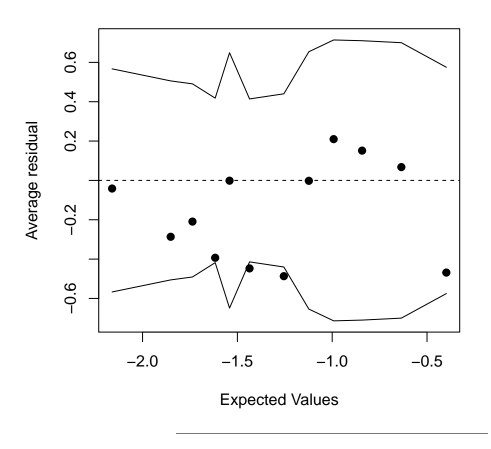
## Loading required package: MASS

##
## arm (Version 1.10-1, built: 2018-4-12)
```

Working directory is /Users/jacobroved/Desktop/LU/Data general/P_dist_statistics

1. GLMM model of offspring recruitment status versus the P-distance PBR with total no. alleles as covariate and nest ID as random factor

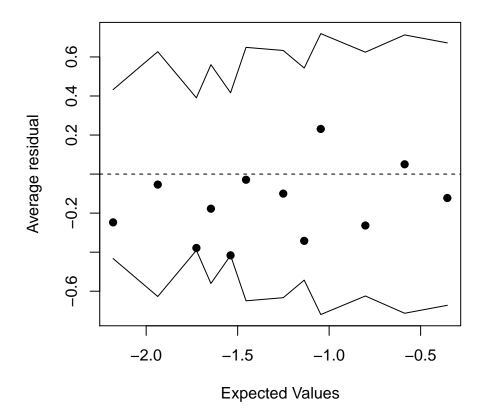
```
fm1 <- glmer(Recruitment~Nest+Total.no.alleles+PdistPBR+(1|Nest),family=binomial)</pre>
## boundary (singular) fit: see ?isSingular
summary(fm1)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
  Family: binomial (logit)
## Formula: Recruitment ~ Nest + Total.no.alleles + PdistPBR + (1 | Nest)
##
##
       AIC
                BIC
                      logLik deviance df.resid
##
      159.2
                       -74.6
                                 149.2
              174.1
##
## Scaled residuals:
##
      Min
                1Q Median
                                3Q
## -0.9343 -0.5746 -0.4492 -0.3211
                                   2.8172
##
## Random effects:
## Groups Name
                       Variance Std.Dev.
## Nest
          (Intercept) 0
## Number of obs: 145, groups: Nest, 34
##
## Fixed effects:
##
                    Estimate Std. Error z value Pr(>|z|)
                                0.36895 -1.555
## (Intercept)
                    -0.57372
                                                  0.1199
                    -0.04428
                                0.02189
                                        -2.023
                                                  0.0431 *
## Total.no.alleles -0.20584
                                0.24294
                                        -0.847
                                                  0.3968
## PdistPBR
                     0.10167
                                0.24300
                                         0.418
                                                  0.6757
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
               (Intr) Nest
##
                            Ttl.n.
## Nest
              -0.832
## Totl.n.llls 0.031 0.017
## PdistPBR
              -0.045 0.026 0.555
## convergence code: 0
## boundary (singular) fit: see ?isSingular
binnedplot(predict(fm1), resid(fm1), cex.pts=1, col.int="black")
```



2. GLMM model of offspring recruitment status versus the P-distance PSS with total no. alleles as covariate and nest ID as random factor

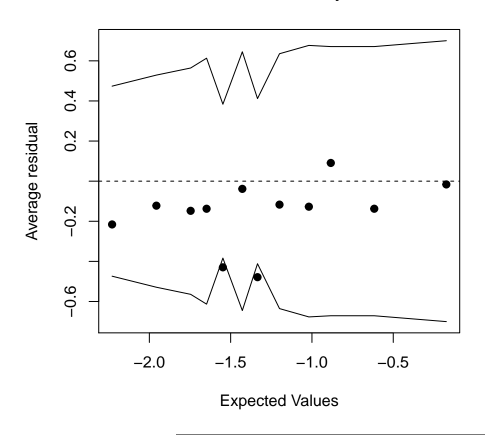
```
fm2 <- glmer(Recruitment~Nest+Total.no.alleles+PdistPSS+(1|Nest),family=binomial)</pre>
## boundary (singular) fit: see ?isSingular
summary(fm2)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
    Family: binomial (logit)
## Formula: Recruitment ~ Nest + Total.no.alleles + PdistPSS + (1 | Nest)
##
##
                       logLik deviance df.resid
        AIC
                 BIC
      158.5
                        -74.3
##
               173.4
                                  148.5
##
  Scaled residuals:
                1Q Median
##
       Min
                                 3Q
                                        Max
   -1.0125 -0.5631 -0.4428 -0.3192
##
##
## Random effects:
                       Variance Std.Dev.
  Groups Name
  Nest
           (Intercept) 0
```

```
## Number of obs: 145, groups: Nest, 34
##
## Fixed effects:
##
                    Estimate Std. Error z value Pr(>|z|)
                                0.36922 -1.576
## (Intercept)
                    -0.58199
                                                  0.1150
## Nest
                    -0.04416
                                0.02184 -2.022
                                                  0.0432 *
## Total.no.alleles -0.14806
                                0.23602 -0.627
                                                  0.5305
## PdistPSS
                                0.23584
                                                  0.3545
                     0.21838
                                          0.926
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
               (Intr) Nest
               -0.831
## Nest
## Totl.n.llls 0.023 0.011
               -0.058 0.010 0.523
## PdistPSS
## convergence code: 0
## boundary (singular) fit: see ?isSingular
binnedplot(predict(fm2), resid(fm2), cex.pts=1, col.int="black")
```



3. GLMM model of offspring recruitment status versus the P-distance PBR with total no. alleles as covariate, sex as a fixed factor, and nest ID as random factor

```
fm3 <- glmer(Recruitment~Nest+Total.no.alleles+PdistPBR+Sex+Total.no.alleles:Sex</pre>
             +PdistPBR:Sex+(1|Nest),family=binomial)
## boundary (singular) fit: see ?isSingular
summary(fm3)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
  Family: binomial (logit)
## Formula:
## Recruitment ~ Nest + Total.no.alleles + PdistPBR + Sex + Total.no.alleles:Sex +
##
      PdistPBR:Sex + (1 | Nest)
##
##
                       logLik deviance df.resid
       AIC
                BIC
      163.5
##
              187.3
                       -73.8
                                147.5
##
## Scaled residuals:
              10 Median
##
      Min
                               3Q
                                       Max
## -1.1101 -0.5431 -0.4505 -0.3267
##
## Random effects:
## Groups Name
                       Variance Std.Dev.
## Nest
           (Intercept) 0
                               0
## Number of obs: 145, groups: Nest, 34
## Fixed effects:
##
                       Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                    0.71719
                                            0.157
                                                       0.875
                        0.11252
                                    0.02194 -2.064
## Nest
                        -0.04529
                                                       0.039 *
                                    0.73636 -0.492
## Total.no.alleles
                       -0.36193
                                                       0.623
## PdistPBR
                        0.27721
                                    0.73005
                                              0.380
                                                       0.704
                       -0.48833
                                    0.42947 -1.137
                                                       0.256
## Total.no.alleles:Sex 0.11392
                                                       0.825
                                    0.51600
                                              0.221
## PdistPBR:Sex
                       -0.15047
                                    0.52319 -0.288
                                                       0.774
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
               (Intr) Nest
                            Ttl.n. PdsPBR Sex
              -0.491
## Nest
## Totl.n.llls 0.117 -0.092
## PdistPBR
              -0.006 -0.064 0.547
              -0.855 0.072 -0.077 0.004
## Ttl.n.111:S -0.109 0.101 -0.943 -0.529 0.072
## PdistPBR:Sx -0.029 0.074 -0.518 -0.943 0.031 0.562
## convergence code: 0
## boundary (singular) fit: see ?isSingular
```



4. GLMM model of offspring recruitment status versus the P-distance PSS with total no. alleles as covariate, sex as a fixed factor, and nest ID as random factor

```
fm4 <- glmer(Recruitment~Nest+Total.no.alleles+PdistPSS+Sex+Total.no.alleles:Sex</pre>
             +PdistPSS:Sex+(1|Nest),family=binomial)
## boundary (singular) fit: see ?isSingular
summary(fm4)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
##
   Family: binomial (logit)
## Formula:
##
  Recruitment ~ Nest + Total.no.alleles + PdistPSS + Sex + Total.no.alleles:Sex +
       PdistPSS:Sex + (1 | Nest)
##
##
##
        AIC
                 BIC
                       logLik deviance df.resid
      162.9
               186.7
                        -73.5
                                  146.9
                                             137
##
## Scaled residuals:
##
       Min
                1Q Median
                                 3Q
                                        Max
```

```
## -1.1370 -0.5422 -0.4522 -0.2866 2.6490
##
## Random effects:
                     Variance Std.Dev.
## Groups Name
## Nest (Intercept) 0
## Number of obs: 145, groups: Nest, 34
## Fixed effects:
                      Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                     0.096661 0.718282 0.135
                                                 0.893
## Nest
                     -0.044211 0.022051 -2.005
                                                 0.045 *
## Total.no.alleles
                     -0.480208 0.721181 -0.666 0.505
                     -0.008826 0.722379 -0.012 0.990
## PdistPSS
## Sex
                     -0.487721 0.430271 -1.134 0.257
## Total.no.alleles:Sex 0.250167 0.499615 0.501 0.617
                      0.162552 0.546985 0.297 0.766
## PdistPSS:Sex
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
             (Intr) Nest Ttl.n. PdsPSS Sex Tt..:S
             -0.494
## Totl.n.llls 0.111 -0.116
## PdistPSS -0.016 -0.115 0.526
## Sex
             -0.854 0.072 -0.041 0.073
## Ttl.n.lll:S -0.090 0.124 -0.944 -0.523 0.015
## PdistPSS:Sx 0.006 0.118 -0.479 -0.946 -0.072 0.534
## convergence code: 0
## boundary (singular) fit: see ?isSingular
binnedplot(predict(fm4), resid(fm4), cex.pts=1, col.int="black")
```

