Lifetime fitness through female and male function: the influence of density and genetically effective population size

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Introduction

The following code performs fixed-effects aster analyses on data examining the effects of density and effective genetic population size (Ne) on female (seeds seet) fitness. LM analysis of biomass (above and below ground) and number of aborted ovules follows the aster analyses.

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Preliminaries

Set working directory and load data.

Examine data

```
fin<- read.csv("data/aster.dat.csv")</pre>
names(fin)
   [1] "Treat"
                      "plotID"
                                   "Den"
                                                "Gen"
                                                             "plantID"
                                   "flw"
                                                             "frt.2"
   [6] "familyID"
                     "surv"
                                                "frt"
## [11] "seeds"
                      "aborted"
                                   "rel.seeds" "seed.wt"
                                                             "mass.a"
## [16] "mass.b"
head(fin)
     Treat plotID Den Gen plantID familyID surv flw frt frt.2 seeds aborted
##
## 1 HDHG
                 5
                     Η
                        HG
                                            4
                                                           3
                                                                       33
                                                                               14
                                  1
## 2
      HDHG
                 5
                     H HG
                                  2
                                            6
                                                  1
                                                      1
                                                         21
                                                                 6
                                                                       69
                                                                               22
## 3
      HDHG
                 5
                                                                 7
                                                                      73
                     Η
                        HG
                                  3
                                           11
                                                      1
                                                         16
                                                                               19
                                                  1
      HDHG
                 5
                     Η
                        HG
                                                      1
                                                         18
                                                                      107
                                                                               23
## 4
                                  4
                                           14
                                                  1
                                  5
## 5
      HDHG
                     H HG
                                            6
                                                      1
                                                          1
                                                                       16
                                                                                4
                                                  1
                                                                 1
## 6 HDHG
                     H HG
                                  6
                                           14
                                                  1
                                                      1
                                                                        0
                                                                                0
     rel.seeds seed.wt mass.a mass.b
## 1 0.5714286
                  6.792
                            5.8
                                    0.8
## 2 1.1948052
                  6.995
                            1.9
                                    0.2
## 3 1.2640693
                  7.413
                           20.8
                                    1.3
## 4 1.8528139
                  8.554
                           18.2
                                    1.1
## 5 0.2770563
                  7.771
                           65.8
                                    6.1
## 6 0.000000
                  0.000
                            6.6
                                    1.2
```

Make sure Den (density treatment), Gen (Ne treatment), plotID, plantID, and familyID are all classified as factors. Otherwise

```
fin$Den<- as.factor(fin$Den)</pre>
fin$Gen<- as.factor(fin$Gen)</pre>
fin$plotID<- as.factor(fin$plotID)</pre>
```

```
fin$plantID<- as.factor(fin$plantID)
fin$familyID<- as.factor(fin$familyID)</pre>
```

Setting up for aster analyses

Load aster package (make sure you have most current version)

```
library(aster)
```

```
## Loading required package: trust
```

Begin by naming variables that will be used in the graphical model of the aster analyses:

flw - total number of flowers produced frt - total number of fruits produced frt.2 - subsetted number of fruits collected seeds - total number of seeds collected from subsetted fruits

```
vars<- c( "flw", "frt", "frt.2", "seeds")</pre>
```

Reshape the data so that all response variables are located in a single vector, in a new data set called "redata"

```
redata <- reshape(fin, varying = list(vars), direction = "long", timevar = "varb", times = as.factor(var
```

Designate the terminal fitness variable "seeds" (make it numberic), and then add it to the reshaped data

```
fit <- grepl("seeds", as.character(redata$varb))
fit<- as.numeric(fit)
redata$fit <- fit</pre>
```

Check

```
with(redata, sort(unique(as.character(varb)[fit == 0])))
## [1] "flw" "frt" "frt.2"
```

```
with(redata, sort(unique(as.character(varb)[fit == 1])))
```

[1] "seeds"

Add a variable "root" to redata, where value is 1. This is the "starting point" of the aster graphical model (i.e. a seed planted)

```
redata<- data.frame(redata, root=1)</pre>
```

Set up the graphical model and designate the statistical distribution for each node. This graphical model has four nodes (in object pred) described earlier. Statistical family for each node is described by object fam.

```
pred<- c(0,1,2,3)
fam<- c(1,2,2,2)
```

Show distribution family for each node

```
sapply(fam.default(), as.character)[fam]
```

```
## [1] "bernoulli" "poisson" "poisson" "poisson"
```

Aster Analyses

First aster analysis with only fitness data. Note, aster reads the redata version of the data.

```
aout<- aster(resp~varb , pred, fam, varb, id, root, data=redata)
summary(aout, show.graph=T)</pre>
```

```
##
## Call:
##
  aster.formula(formula = resp ~ varb, pred = pred, fam = fam,
       varvar = varb, idvar = id, root = root, data = redata)
##
##
##
## Graphical Model:
    variable predecessor family
##
##
   flw
             root
                         bernoulli
##
  frt
             flw
                         poisson
  frt.2
             frt
                         poisson
##
    seeds
             frt.2
                         poisson
##
##
               Estimate Std. Error z value Pr(>|z|)
## (Intercept) -44.6476
                             0.5507
                                     -81.08
                                              <2e-16 ***
## varbfrt
                49.1734
                             0.5599
                                      87.82
                                              <2e-16 ***
## varbfrt.2
                34.2048
                             0.5538
                                      61.76
                                              <2e-16 ***
## varbseeds
                46.9877
                             0.5507
                                      85.32
                                              <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Add density to the model as a fixed effect. Note that we follow the "no naked predictors" rule with fit: (Den)
aout.d<- aster(resp~varb + fit:(Den), pred, fam, varb, id, root, data=redata)
summary(aout.d, show.graph=T)
##
## aster.formula(formula = resp ~ varb + fit:(Den), pred = pred,
##
       fam = fam, varvar = varb, idvar = id, root = root, data = redata)
##
##
## Graphical Model:
   variable predecessor family
##
  flw
             root
                         bernoulli
##
  frt.
             flw
                         poisson
##
   frt.2
             frt
                         poisson
##
    seeds
             frt.2
                         poisson
##
##
                 Estimate Std. Error z value Pr(>|z|)
## (Intercept) -39.755770
                             0.597861 -66.497
                                                <2e-16 ***
## varbfrt
                44.287436
                             0.606131 73.066
                                                <2e-16 ***
## varbfrt.2
                29.362530
                                       49.053
                             0.598590
                                                <2e-16 ***
                42.077712
## varbseeds
                             0.597597 70.412
                                                <2e-16 ***
                 0.041854
## fit:DenH
                             0.003121 13.411
                                                <2e-16 ***
## fit:DenL
                -0.039658
                             0.004752 -8.346
                                                <2e-16 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Original predictor variables dropped (aliased)
##
        fit:DenM
```

Do not try to determine the significance <code>Density</code> from the <code>summary</code> statement. These are not reliable. Instead, perform a liklihood ratio test. The results of the liklihood ratio test show that the model with <code>Density</code> explains significantly more variation than the model without.

```
anova(aout, aout.d)
## Analysis of Deviance Table
##
## Model 1: resp ~ varb
## Model 2: resp ~ varb + fit:(Den)
   Model Df Model Dev Df Deviance P(>|Chi|)
## 1
           4
                 132606
## 2
            6
                 133112 2
                             505.61 < 2.2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Make Up Data
Simulate regression data, and do the regression.
n <- 50
x \leftarrow seq(1, n)
a.true <- 3
b.true <- 1.5
y.true <- a.true + b.true * x
s.true <- 17.3
y <- y.true + s.true * rnorm(n)
out1 <- lm(y \sim x)
summary(out1)
##
## Call:
## lm(formula = y \sim x)
## Residuals:
##
       Min
                1Q Median
                                3Q
## -39.321 -12.201 0.689 11.360 35.922
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) -2.1804 4.8487
                                    -0.45
                                             0.655
## x
                 1.6729
                           0.1655
                                    10.11 1.77e-13 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 16.89 on 48 degrees of freedom
## Multiple R-squared: 0.6804, Adjusted R-squared: 0.6737
## F-statistic: 102.2 on 1 and 48 DF, p-value: 1.775e-13
```

Figure with Code to Make It Shown

The following figure is produced by the following code

```
mydata <- data.frame(x, y)
plot(mydata)</pre>
```

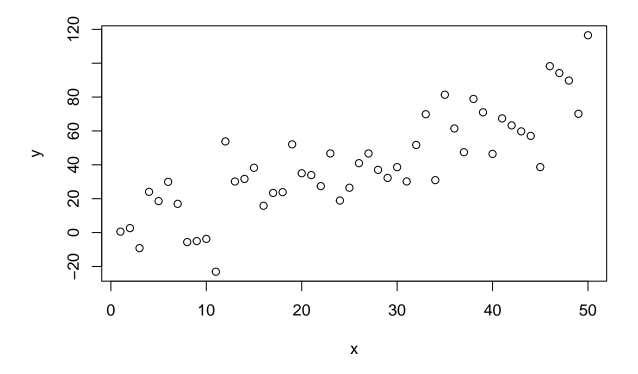


Figure 1: Simple Linear Regression