# Appendix: Modeling Wing Morphology

for 'paper\_title'; Bernat, AV, Cenzer, ML

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# Details of the Analyses

This document was generated by R Markdown on 2021-07-27 using R version 4.0.5. The document provides the step-by-step analytical methods used in the manuscript by Anastasia Bernat (AVB) and Meredith Cenzer (MLC). Multiple draft scripts were written by AVB between 2021-03-01 and 2021-07-26 until being distilled and complied by AVB and code reviewed by MLC at the University of Chicago into this comprehensive script. All draft scripts can be viewed in the GitHub repository, SBB-dispersal (https://github.com/mlcenzer/SBB-dispersal), within the directory avbernat\_working\_on > All\_Morphology > stats.

All code and output from the statistical analyses is shown. Code for data cleaning and the generation of plots is not displayed, but can be viewed in the wing\_summary.Rmd file and its accompanying three sourced scripts. To repeat analyses and the generation of plots, all data files and sourced scripts should follow the directory structure presented in the SBB-dispersal repository.

# Description of the Data

This document analyzes two main datasets, **raw\_data** and **data\_long**. The raw\_data set provides morphology measurements for each soapberry bug, *Jadera haematoloma*, collected and measured between the years 2013 and 2020. There are four morphology measurements: beak length, thorax width, wing length, and body length. The sex, wing morph (long-winged, shot-winged, or ambiguously-winged), and host plant the bug was collected from as well as the month and year each bug was collected in was recorded. The data\_long set provides the same recordings as the raw\_data set, but it has been filtered for only long-winged soapberry bugs.

### Abbreviations Used in the Data and Code

- sd standard deviation
- $\bullet$  se standard error
- **binom** a column name that ends in 'binom' is a column that has been recodified into binary data (0's and 1's). Example columns: sex\_binom, pophost\_binom, and wing\_morph\_binom
- **pophost** the host plant soapberry bugs were collected from, which was either *Koelreuteria* elegans or *Cardiospermum corindum*, occasionally called (and abbreviated) as goldenrain tree (GRT) or balloon vine (BV), respectively
- months since start proxy for year where the first collection occurred on May 2013
- month\_of\_year proxy for season where collections occurred only in months February, April, May, August, September, October, and September
- wing2body a computed and unitless column that calculates the wing length divided by the body length of a soapberry bug
- wing2thorax a computed and unitless column that calculates the wing length divided by the thorax width of a soapberry bug
- S short-winged morph
- L long-winged morph
- LS or SL ambiguous wing morph

# **Data Cleaning And Exploration**

# Read Libraries

The occurrence of long-wing morphology and the wing-to-body ratio of *J. haematoloma* were analyzed using multivariate, generalized linear modeling (GLM) as implemented in the R packages lme4 and binom. The dplyr package helped speed data manipulation processes by grouping data quickly. All plots, except the histograms, were generated using ggplot libraries and helper functions found in R packages ggformula and cowplot.

Additional R packages not shown below, but embedded in the sourced scripts are zoo and lubridate, which aid in data manipulation and datetime manipulation, respectively.

```
library(lme4) # fit regressions
library(dplyr) # data manipulation
library(ggformula) # ggplot plotting
library(cowplot) # ggplot helper functions to arrange multi-panel figures
library(binom) # binomial confidence intervals
```

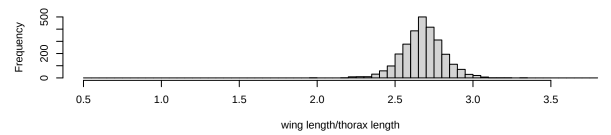
#### Read Source Files

#### Read the Data

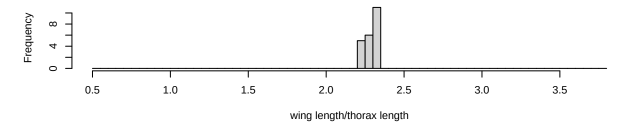
```
data_list = read_morph_data(paste0(dir,"All_Morphology/stats/data/allmorphology05.18.21.csv"))
## number of missing dates: 0
##
## morph types: L S NA LS SL
##
      recoding missing morph types...
      S if wing2thorax <=2.2, L if wing2thorax >=2.5
##
##
## ambiguous wing morph bug count:
## filtered out NA wing2body for data_long...
raw_data = data_list[[1]]
data_long = data_list[[2]] # long-wing bugs only
data_long = remove_torn_wings(data_long)
##
## number of bugs with torn wings: 193
```

# Histograms of Wing Morph Data

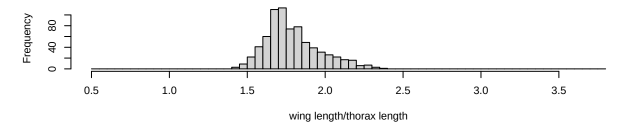
# Histogram of wing length/thorax length for long winged SBB



# Histogram of wing length/thorax length for SBB w/o recorded wing morph



# Histogram of wing length/thorax length for short winged SBB

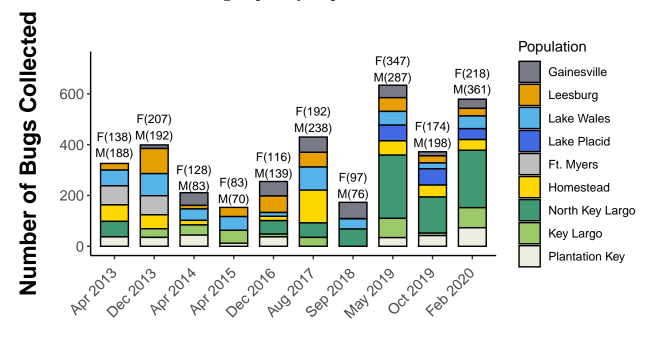


Notice that there are 30 bugs that are hard to identify as either S or L.

# **Barplots**

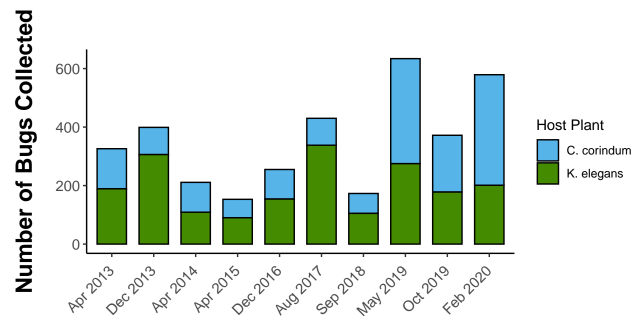
Bugs were collected on different years and months. These barplots show the bugs collected per **population**, **host plant**, and **sex** across the years and months.

Plot 1: Collection numbers grouped by Population and Datetime



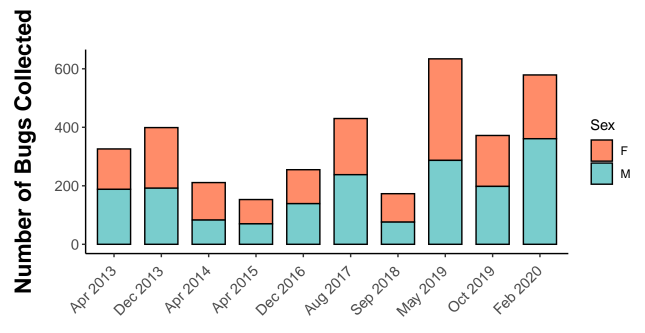
# **Field Collection Month**

Plot 2: Collection numbers grouped by Host Plant and Datetime



**Field Collection Month** 

Plot 3: Collection numbers grouped by Sex and Datetime



# **Field Collection Month**

# Regression Modeling

# Long-Wing Morph Frequency

data = data.frame(R=raw\_data\$wing\_morph\_binom,

How does sex, host plant, month, and/or year effect whether a soapberry bug is long-winged (wing\_morph\_binom=1) or short-winged (wing\_morph\_binom=0)?

```
A=raw_data$sex_binom,
                 B=raw_data$pophost_binom,
                  C=(raw_data$month_of_year),
                 D=raw_data$months_since_start)
model_script = pasteO(source_path, "generic models-binomial glm 4-FF.R")
model_comparisonsAIC(model_script)
##
                     [,2]
                                [,3]
          [,1]
                                          [,4]
                                                      [,5]
          3145.306
                                          3147.201
## AICs
                     3146.842
                               3147.157
                                                     3148.521
## models 98
                     110
                               84
                                          107
                                                      105
          0.2529382 0.1173602 0.1002697 0.09808583 0.05068685
##
       glm(formula = R \sim A * B + A * D + B * C + C * D, family = binomial,
## m98
       data = data)
##
## m110
            glm(formula = R \sim A * B + A * D + B * C + B * D + C * D, family = binomial,
       data = data)
        glm(formula = R \sim A * D + B * C + C * D, family = binomial, data = data)
## m84
            glm(formula = R \sim A * B + A * C + A * D + B * C + C * D, family = binomial,
## m107
##
       data = data)
## m105
            glm(formula = R \sim A * D + B * C + B * D + C * D, family = binomial,
##
       data = data)
```

```
anova(m98, m110, test="Chisq") # adding B*D does not improve fit
anova(m84, m98, test="Chisq") # adding A*B improves fit
anova(m63, m84, test="Chisq") # Adding C*D improves fit
anova(m51, m63, test="Chisq") # Adding B improves fit
## Analysis of Deviance Table
##
## Model 1: R ~ A * B + A * D + B * C + C * D
## Model 2: R ~ A * B + A * D + B * C + B * D + C * D
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          3461
                  3127.3
## 2
          3460
                  3126.8 1 0.46421
                                       0.4957
## Analysis of Deviance Table
## Model 1: R ~ A * D + B * C + C * D
## Model 2: R ~ A * B + A * D + B * C + C * D
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          3462
                  3131.2
## 2
          3461
                  3127.3 1
                              3.8506 0.04973 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Analysis of Deviance Table
## Model 1: R \sim A * D + C * D + B
## Model 2: R ~ A * D + B * C + C * D
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          3463
                  3137.3
## 2
          3462
                              6.1886 0.01286 *
                  3131.2 1
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Analysis of Deviance Table
##
## Model 1: R ~ A * D + C * D
## Model 2: R ~ A * D + C * D + B
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
                  3497.3
          3464
          3463
## 2
                  3137.3 1
                              359.93 < 2.2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

#### Best Fit

```
M1 = glm(wing_morph_binom ~ sex_binom * pophost_binom + sex_binom * months_since_start +
          pophost_binom * month_of_year + month_of_year * months_since_start,
         data=raw_data, family="binomial")
summary(M1)
##
## Call:
## glm(formula = wing_morph_binom ~ sex_binom * pophost_binom +
       sex_binom * months_since_start + pophost_binom * month_of_year +
##
##
       month_of_year * months_since_start, family = "binomial",
       data = raw_data)
##
##
## Deviance Residuals:
##
      Min
                     Median
                                          Max
                 10
                                  3Q
## -2.3803
                     0.4321
                                        1.2552
             0.3597
                               0.8450
## Coefficients:
##
                                     Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                     0.7516501 0.1841942 4.081 4.49e-05 ***
## sex_binom
                                    -0.2597900 0.0902673 -2.878 0.004002 **
                                     1.1256358 0.1142931 9.849 < 2e-16 ***
## pophost_binom
## months_since_start
                                     0.0107239 0.0029582 3.625 0.000289 ***
## month_of_year
                                    0.0995560 0.0255307 3.899 9.64e-05 ***
## sex_binom:pophost_binom
                                    0.0973323  0.0495811  1.963  0.049635 *
## sex_binom:months_since_start
                                    0.0037212 0.0015337 2.426 0.015254 *
## pophost_binom:month_of_year
                                    -0.0379395
                                                0.0150617 -2.519 0.011771 *
## months_since_start:month_of_year -0.0014557 0.0004553 -3.198 0.001386 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 3562.3 on 3469 degrees of freedom
## Residual deviance: 3127.3 on 3461
                                       degrees of freedom
     (62 observations deleted due to missingness)
## AIC: 3145.3
## Number of Fisher Scoring iterations: 5
```

### Long-Wing Morph Variance

SE = function(x){sd(x)/sqrt(length(x))}

In addition to modeling how sex, host plant, month, or year effects whether a soapberry bug is long-winged or short-winged, we modeled how those same factors affected the variance of wing morph.

data=raw\_data,FUN=SE)\$wing\_morph\_binom
wmorph\_table\$n = aggregate(wing\_morph\_binom~sex\_binom\*pophost\_binom\*month\_of\_year\*months\_since

wmorph\_table\$se = aggregate(wing\_morph\_binom~sex\_binom\*pophost\_binom\*month\_of\_year\*months\_sinc

```
data=raw_data,FUN=length) $wing_morph_binom
```

```
data = wmorph_table
data = data.frame(R=data$sd,
                 A=data$sex_binom,
                 B=data$pophost_binom,
                 C=(data$month_of_year),
                 D=data$months since start)
model_script = paste0(source_path, "generic models-gaussian glm 4-FF.R")
model_comparisonsAIC(model_script)
##
          [,1]
                    [,2]
                              [,3]
                                         [,4]
## AICs
          -92.39855 -90.95292 -90.75898 -90.41465
## models 2
                    5
                              8
                                         9
## probs 0.183788 0.0892081 0.08096352 0.06815837
##
## m2
        glm(formula = R ~ B, family = gaussian, data = data)
## m5
        glm(formula = R ~ A + B, family = gaussian, data = data)
## m8
        glm(formula = R ~ B + C, family = gaussian, data = data)
## m9
        glm(formula = R ~ B + D, family = gaussian, data = data)
anova(m2, m5, test="Chisq") # Adding A does not improve fit
anova(m2, m8, test="Chisq") # Adding C does not improve fit
anova(m2, m9, test="Chisq") # Adding D does not improve fit
anova(m0, m2, test="Chisq") # Adding B improves fit
## Analysis of Deviance Table
##
## Model 1: R ~ B
## Model 2: R ~ A + B
     Resid. Df Resid. Dev Df Deviance Pr(>Chi)
                  0.20009
## 1
            38
## 2
            37
                  0.19734 1 0.0027541
                                         0.4724
## Analysis of Deviance Table
## Model 1: R ~ B
## Model 2: R ~ B + C
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
            38
                  0.20009
            37
                  0.19830 1 0.0017949
                                         0.5628
## Analysis of Deviance Table
## Model 1: R ~ B
## Model 2: R ~ B + D
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
            38
                  0.20009
## 2
            37
                  0.20001 1 8.0534e-05
## Analysis of Deviance Table
##
## Model 1: R ~ 1
## Model 2: R ~ B
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
```

```
39
## 1
                 0.62439
## 2
           38
                 0.20010 1 0.42429 < 2.2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Best Fit
M2 = glm(sd ~ pophost binom, data=wmorph table, family="gaussian")
summary(M2)
##
## Call:
## glm(formula = sd ~ pophost_binom, family = "gaussian", data = wmorph_table)
## Deviance Residuals:
##
        Min
                    1Q
                           Median
                                          3Q
                                                    Max
## -0.249168 -0.041487
                         0.005877
                                    0.041147
                                               0.171269
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                 0.35216
                            0.01147 30.693 < 2e-16 ***
## pophost_binom -0.10299
                            0.01147 -8.976 6.28e-11 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for gaussian family taken to be 0.005265667)
      Null deviance: 0.62439 on 39
                                     degrees of freedom
```

### Wing-to-Body Ratio

## AIC: -92.399

##

How does sex, host plant, month, and/or year effect the wing-to-body ratio of long-winged soapberry bugs?

## Residual deviance: 0.20010 on 38 degrees of freedom

## Number of Fisher Scoring iterations: 2

```
data = data.frame(R=data_long$wing2body_c, # centered
                 A=data_long$sex_binom,
                 B=data_long$pophost_binom,
                 C=data_long$month_of_year_c, # centered
                 D=data_long$months_since_start_c) # centered
model_script = paste0(source_path, "generic models-gaussian glm 4-FF.R")
model_comparisonsAIC(model_script)
##
                               [,3]
          [,1]
                     [,2]
                                          [,4]
                                                      [,5]
                                                                 [,6]
## AICs
          -9722.301 -9721.371 -9720.852
                                          -9720.339
                                                     -9720.331
                                                                 -9719.674
## models 88
                    99
                               58
                                          92
                                                      97
                                                                 76
```

## probs 0.1948772 0.1224324 0.09441271 0.07306166 0.07277994 0.05239229

## m88 glm(formula =  $R \sim A * B + A * D + B * D + C$ , family = gaussian,

```
##
       data = data)
       glm(formula = R \sim A * B + A * D + B * D + C * D, family = gaussian,
## m99
##
       data = data)
## m58 glm(formula = R \sim A * B + B * D + C, family = gaussian, data = data)
## m92 glm(formula = R \sim A * B + A * C + A * D + B * D, family = gaussian,
      data = data)
## m97
       glm(formula = R \sim A * B + A * D + B * C + B * D, family = gaussian,
##
       data = data)
## m76 glm(formula = R \sim A * B + B * D + C * D, family = gaussian, data = data)
anova(m88, m58, test="Chisq") # Adding A*D marginally improves fit
anova(m76, m58, test="Chisq") # Adding C*D does not improve fit
anova(m88, m99, test="Chisq") # adding C*D does not improve fit
anova(m34, m58, test="Chisq") # Adding B*D improves fit
## Analysis of Deviance Table
## Model 1: R ~ A * B + A * D + B * D + C
## Model 2: R ~ A * B + B * D + C
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          1895
               0.66692
## 2
          1896
                  0.66813 -1 -0.00121 0.06371 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Analysis of Deviance Table
##
## Model 1: R ~ A * B + B * D + C * D
## Model 2: R ~ A * B + B * D + C
##
     Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          1895
                  0.66784
## 2
          1896
                  0.66813 -1 -0.0002886
                                          0.3655
## Analysis of Deviance Table
## Model 1: R ~ A * B + A * D + B * D + C
## Model 2: R ~ A * B + A * D + B * D + C * D
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          1895
                  0.66692
          1894
                  0.66655 1 0.00037502
## Analysis of Deviance Table
##
## Model 1: R ~ A * B + C + D
## Model 2: R ~ A * B + B * D + C
     Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          1897
                  0.67063
## 2
          1896
                  0.66813 1 0.0024994 0.00774 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

#### Best Fit

```
M3 = glm(wing2body_c ~ sex_binom*pophost_binom + pophost_binom*months_since_start_c
         + month_of_year_c, data=data_long, family=gaussian)
summary(M3)
##
## Call:
## glm(formula = wing2body_c ~ sex_binom * pophost_binom + pophost_binom *
       months_since_start_c + month_of_year_c, family = gaussian,
       data = data_long)
##
##
## Deviance Residuals:
        Min
                    1Q
                           Median
                                          30
                                                    Max
## -0.070837 -0.010794 -0.000093
                                    0.010596
                                               0.113993
##
## Coefficients:
                                       Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                                     -4.542e-04 4.601e-04 -0.987 0.32368
                                     -1.787e-03 4.467e-04 -4.001 6.55e-05 ***
## sex_binom
## pophost_binom
                                      4.289e-03 4.613e-04 9.297 < 2e-16 ***
                                     -1.727e-05 2.225e-05 -0.776 0.43763
## months since start c
## month_of_year_c
                                      7.155e-04 1.379e-04 5.188 2.35e-07 ***
                                      1.804e-03 4.466e-04 4.038 5.60e-05 ***
## sex_binom:pophost_binom
## pophost_binom:months_since_start_c 5.904e-05 2.217e-05 2.663 0.00781 **
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for gaussian family taken to be 0.0003523901)
##
       Null deviance: 0.72538 on 1902
                                       degrees of freedom
## Residual deviance: 0.66813
                              on 1896
                                       degrees of freedom
## AIC: -9720.9
##
## Number of Fisher Scoring iterations: 2
```

### Wing-to-Body Ratio Variance

In addition to modeling how sex, host plant, month, or year effects the wing-to-body ratio of a soapberry bug, we modeled how those same factors affected the variance of wing-to-body ratio.

```
model_comparisonsAIC(model_script)
##
          [,1]
                    [,2]
                              [,3]
          -280.1872 -279.8713 -279.4758
## AICs
## models 8
                    19
                              2
## probs 0.1198675 0.1023577 0.08398967
##
## m8
        glm(formula = R ~ B + C, family = gaussian, data = data)
## m19 glm(formula = R ~ B * C, family = gaussian, data = data)
        glm(formula = R ~ B, family = gaussian, data = data)
## m2
anova(m8, m19, test="Chisq") # Adding B*C does not improve fit
anova(m2, m8, test="Chisq") # Adding C does not improve fit
anova(m0, m2, test="Chisq") # Adding B improves fit
## Analysis of Deviance Table
##
## Model 1: R ~ B + C
## Model 2: R ~ B * C
     Resid. Df Resid. Dev Df Deviance Pr(>Chi)
            33 0.00070342
## 1
            32 0.00067127 1 3.215e-05
## 2
## Analysis of Deviance Table
##
## Model 1: R ~ B
## Model 2: R ~ B + C
    Resid. Df Resid. Dev Df
                               Deviance Pr(>Chi)
##
## 1
           34 0.00075844
            33 0.00070342 1 5.5025e-05 0.1081
## Analysis of Deviance Table
##
## Model 1: R ~ 1
## Model 2: R ~ B
    Resid. Df Resid. Dev Df
                               Deviance Pr(>Chi)
            35 0.00087733
## 1
## 2
            34 0.00075844 1 0.00011888 0.02097 *
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
Best Fit
M4 = glm(sd ~ pophost_binom, data=w2b_table, family=gaussian)
summary(M4)
##
## Call:
## glm(formula = sd ~ pophost_binom, family = gaussian, data = w2b_table)
##
## Deviance Residuals:
                       1Q
                               Median
          Min
                                               3Q
                                                          Max
## -0.0059374 -0.0033018 -0.0006274
                                        0.0022332
                                                    0.0147212
##
```

model\_script = paste0(source\_path, "generic models-gaussian glm 4-FF.R")

```
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                0.0165999 0.0007872 21.088
                                               <2e-16 ***
## pophost_binom 0.0018172 0.0007872
                                       2.309
                                               0.0272 *
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for gaussian family taken to be 2.230719e-05)
##
##
       Null deviance: 0.00087733 on 35 degrees of freedom
## Residual deviance: 0.00075844 on 34 degrees of freedom
## AIC: -279.48
##
## Number of Fisher Scoring iterations: 2
```

# LOESS & Linear Regression Plots

## Wing Morph Frequency

Group significant elements

```
# function to calculate 95% confidence interval (CI).
CI_95 = function(x) \{qnorm(0.975)*sd(x)/sqrt(length(x))\}
CI_95_binom_upper = function(y) {binom.confint(x=sum(y, na.rm=TRUE), n=length(y[!is.na(y)]),
                                                 conf.level=0.95, methods='exact')$upper}
CI_95_binom_lower = function(y) {binom.confint(x=sum(y, na.rm=TRUE), n=length(y[!is.na(y)]),
                                                 conf.level=0.95, methods='exact')$lower}
# aggregate the full data and compute standard error (SE), upper and lower CI, & sample size (
w_morph_summary = aggregate(wing_morph_binom~sex*pophost*month_of_year*months_since_start,
                           data=raw_data, FUN=mean)
w_morph_summary$se = aggregate(wing_morph_binom~sex*pophost*month_of_year*months_since_start,
                              data=raw_data, FUN=SE) $wing_morph_binom
w_morph_summary$upper = aggregate(wing_morph_binom~sex*pophost*month_of_year*months_since_star
                                 data=raw_data, FUN=CI_95_binom_upper)$wing_morph_binom
w_morph_summary$lower = aggregate(wing_morph_binom~sex*pophost*month_of_year*months_since_star
                                 data=raw_data, FUN=CI_95_binom_lower) $\sinc \text{wing_morph_binom}$
w_morph_summary$n = aggregate(wing_morph_binom~sex*pophost*month_of_year*months_since_start,
                             data=raw_data, FUN=length) $\sing_morph_binom
jitter = runif(n=nrow(w morph summary), min=-0.1, max=0.1) # jitter points slightly
w_morph_summary$dates = w_morph_summary$month_of_year + jitter
dd = w_morph_summary
```

### Check for LOESS Residuals

```
plot_lowess_residuals = function(lfit, x, y) {
    lfun = approxfun(lfit)
    fitted = lfun(x)
    resid = y-fitted
    plot(fitted,resid)
```

```
abline(h=0,col=8)
}
11 = lowess(dd$dates, dd$wing_morph_binom, f=0.4) # f = alpha, the smoother span
par(mfrow=c(1,2))
plot(dd$dates, dd$wing_morph_binom)
lines(11, type = "1")
plot_lowess_residuals(l1, dd$dates, dd$wing_morph_binom)
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                 4
                            8
                                10
                                      12
                                                           0.65
                                                                    0.75
                                                                              0.85
                      6
                     dd$dates
                                                                      fitted
```

Figure: Panels A, B, C, D (long-wing morph freq with month) & E (long-wing morph freq with year)

Note: Single-variate models or simpler models substituted best fit slope calculations below in order to avoid multiple interaction terms found in the best fit model. This led to cleaner glm line plotting.

# Panels A and B Regression Computations:

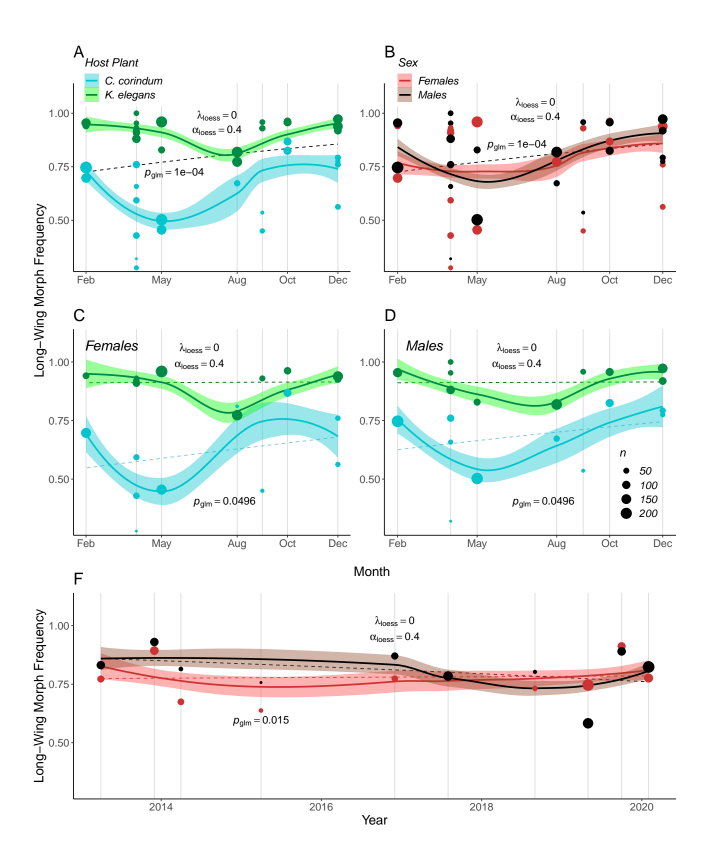
```
# single-variate model of month predicting wing morph
fit = glm(wing_morph_binom ~ month_of_year, family="binomial", data=raw_data)
xmonth = seq(2,12, 0.01)
wing_probs = predict(fit, list(month_of_year=xmonth), type="response")

# extract pvalue from best fit regression model
fit_pvalue = round(summary(M1)$coeff[,"Pr(>|z|)"][5],5)
pvalue = paste0("italic(p)[glm]==", fit_pvalue)
```

### Panels C and D Regression Computations:

### Panel F Regression Computations:

```
# multi-variate model with year, sex, and host plant predicting wing morph
fit3 = glm(wing_morph_binom ~ sex_binom * dates, family = "binomial", data = raw_data)
xyr = seq(sort(unique(dd$dates))[1],sort(unique(dd$dates))[10], 1)
set.seed(194842)
bsex = sample(c(-1,1), replace=TRUE, size=length(xyr))
bhost = sample(c(-1,1), replace=TRUE, size=length(xyr))
wprobs = predict(fit3, list(sex_binom = bsex,
                                pophost_binom = bhost,
                                dates = xyr), type="response")
pred = cbind(xyr, bsex, bhost, wprobs)
pred = as.data.frame(pred)
pred$xyr = as.Date.numeric(pred$xyr)
predF = pred[pred$bsex==1,]
predM = pred[pred$bsex==-1,]
# extract pualue from best fit regression model
fit_pvalue = round(summary(M1)$coeff[,"Pr(>|z|)"][7],3)
pvalue = paste0("italic(p)[glm]==", fit_pvalue)
```



### Wing-to-Body Ratio

### Group significant elements

### Check for LOESS Residuals

```
11 = lowess(d$dates, d$wing2body, f=0.4)

par(mfrow=c(1,2))
plot(d$dates, d$wing2body)
lines(l1, type = "l")
plot_lowess_residuals(l1, d$dates, d$wing2body)
```

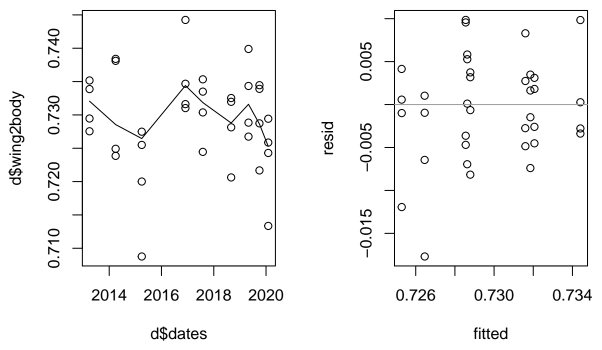


Figure: Panels A, B, (wing-to-body ratio with month) & C (wing-to-body ratio with year)

### Panel C Regression Computations:

```
pred = cbind(xyr, bhost, wprobs)
pred = as.data.frame(pred)
pred$xyr = as.Date.numeric(pred$xyr)

predK = pred[pred$bhost==1,]
predC = pred[pred$bhost==-1,]

# extract pvalue from best fit regression model
fit_pvalue = round(summary(M3)$coeff[,"Pr(>|t|)"][7],3)
pvalue = paste0("italic(p)[glm]==", fit_pvalue)
```

