Modeling Wing Morphology Summary File

Data Cleaning And Exploration

Read Libraries

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

Read Source Files

Read the Data

```
data_list <- read_morph_data("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: 48

##

## filtered out NA wing2body for data_long...

raw_data = data_list[[1]]

data_long = data_list[[2]] # long-wing bugs only

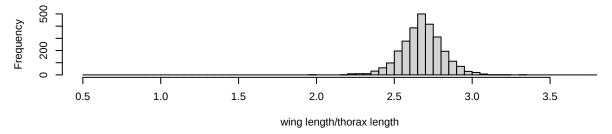
data_long = remove_torn_wings(data_long)

##

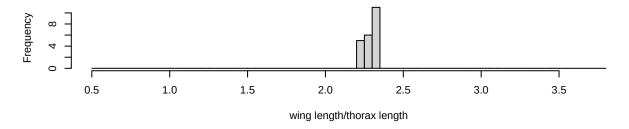
## number of bugs with torn wings: 193
```

Histograms of Wing Morph Data

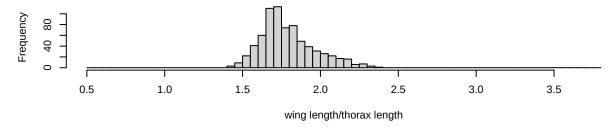
Histogram of wing length/thorax length for long winged SBB



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



Histogram of wing length/thorax length for short winged SBB

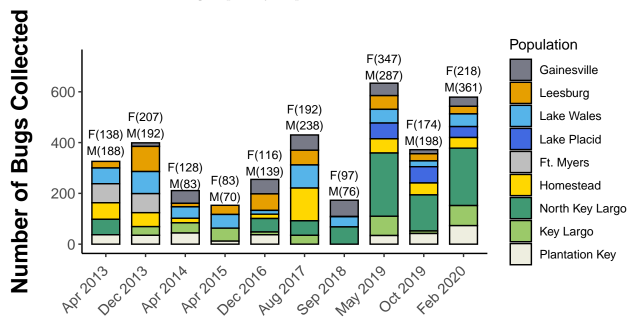


Notice that there are 30 bugs that are hard to identify as either S or L.

Barplots

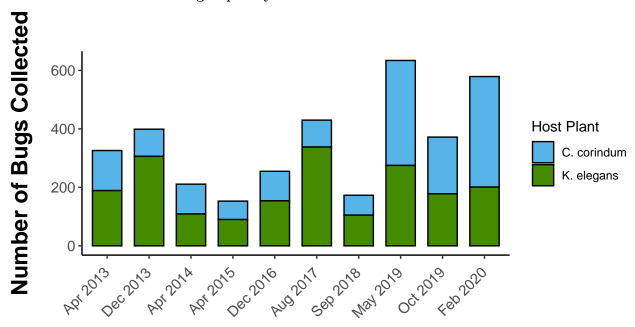
Bugs were collected on different years and months. These barplots show the bugs collected per **population**, **host plant**, and **sex** across the years and months.

Plot 1: Collection numbers grouped by Population and Datetime



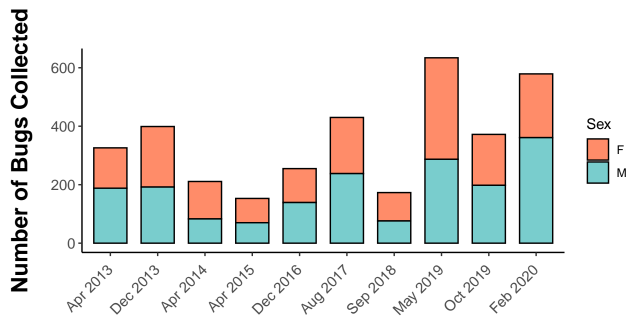
Field Collection Month

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

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)?

```
data<-data.frame(R=raw_data$wing_morph_binom,
                 A=raw_data$sex_binom,
                 B=raw_data$pophost_binom,
                 C=(raw_data$month_of_year),
                 D=raw_data$months_since_start)
model_script = paste0(source_path, "generic models-binomial glm 4-FF.R")
model_comparisonsAIC(model_script)
##
          [,1]
                     [,2]
                               [,3]
                                         [,4]
                                                     [,5]
## AICs
          3145.306 3146.842 3147.157
                                         3147.201
                                                     3148.521
## models 98
                    110
                               84
                                         107
                                                     105
```

```
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 \sim 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 ~ 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
                  3131.2 1 6.1886 0.01286 *
## ---
## 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
         3464
                  3497.3
## 2
         3463
                  3137.3 1 359.93 < 2.2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Best Fit

```
M2 = 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 (M2)
##
## 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
                 10
                      Median
                                   30
                                           Max
##
## -2.3803
                      0.4321
                                        1.2552
            0.3597
                               0.8450
##
## Coefficients:
                                      Estimate Std. Error z value Pr(>|z|)
##
                                                          4.081 4.49e-05 ***
## (Intercept)
                                     0.7516501 0.1841942
## sex binom
                                    -0.2597900 0.0902673 -2.878 0.004002 **
## pophost_binom
                                     1.1256358   0.1142931   9.849   < 2e-16 ***
## months_since_start
                                     0.0107239 0.0029582
                                                            3.625 0.000289 ***
## month_of_year
                                               0.0255307
                                                            3.899 9.64e-05 ***
                                     0.0995560
## 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 *
                                                0.0150617 -2.519 0.011771 *
## pophost_binom:month_of_year
                                    -0.0379395
## 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
```

Modeling Variance

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 = 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]
         -92.39855 -90.95292 -90.75898 -90.41465
## AICs
## models 2
                   5
                              8
## 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 {\it 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)
## 1
           38 0.20009
           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
                 0.20001 1 8.0534e-05 0.9029
           37
## 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

```
M3 = glm(sd ~ pophost_binom, data=wmorph_table, family="gaussian")
summary(M3)
##
## Call:
## glm(formula = sd ~ pophost_binom, family = "gaussian", data = wmorph_table)
##
## Deviance Residuals:
##
        Min
                     1Q
                            Median
                                           ЗQ
                                                     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
##
## Residual deviance: 0.20010 on 38 degrees of freedom
## AIC: -92.399
## Number of Fisher Scoring iterations: 2
```

Wing-to-body Ratio

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

```
##
          [,1]
                     [,2]
                                [,3]
                                           [,4]
                                                       [,5]
                                                                  [,6]
## AICs
          -9722.301 -9721.371 -9720.852
                                           -9720.339
                                                      -9720.331
                                                                  -9719.674
                                                       97
                                                                  76
## models 88
                     99
                               58
                                           92
## 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(m22, m42, test="Chisq") # adding B*C does not improve fit
anova(m16, m22, test="Chisq") # adding C does improve fit
anova(m22, m34, test="Chisq") # adding D does not improve fit
anova(m34, m58, test="Chisq") # Adding B*D marginally improves fit
## Analysis of Deviance Table
## Model 1: R ~ A * B + C
## Model 2: R ~ A * B + B * C
    Resid. Df Resid. Dev Df
                              Deviance Pr(>Chi)
## 1
         1898
                 0.67065
## 2
         1897
                 0.67061 1 4.1833e-05
                                         0.7308
## Analysis of Deviance Table
## Model 1: R ~ A * B
## Model 2: R ~ A * B + C
   Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
         1899
                 0.68181
## 2
         1898
                 0.67065 1 0.011157 1.919e-08 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Analysis of Deviance Table
##
## Model 1: R ~ A * B + C
## Model 2: R ~ A * B + C + D
   Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
         1898
                 0.67065
## 2
         1897
                 0.67063 1 2.1817e-05 0.8038
## 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
M4 = glm(wing2body_c ~ sex_binom*pophost_binom + month_of_year_c, data=data_long, family=gaussian)
summary(M4)
##
## Call:
```

```
## glm(formula = wing2body_c ~ sex_binom * pophost_binom + month_of_year_c,
##
       family = gaussian, data = data_long)
##
## Deviance Residuals:
##
        Min
                           Median
                                          3Q
                                                    Max
  -0.070223 -0.010939 -0.000117
                                    0.010773
                                               0.114608
##
##
## Coefficients:
##
                            Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                          -0.0007445 0.0004473 -1.664
                                                          0.0962 .
## sex_binom
                          -0.0017478
                                      0.0004468 -3.912 9.48e-05 ***
## pophost_binom
                                                  9.762 < 2e-16 ***
                           0.0043900 0.0004497
## month_of_year_c
                           0.0007628 0.0001358
                                                  5.619 2.20e-08 ***
## sex_binom:pophost_binom 0.0018351 0.0004468
                                                  4.108 4.17e-05 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
  (Dispersion parameter for gaussian family taken to be 0.0003533471)
##
##
      Null deviance: 0.72538 on 1902 degrees of freedom
## Residual deviance: 0.67065 on 1898 degrees of freedom
## AIC: -9717.7
##
## Number of Fisher Scoring iterations: 2
```

Modeling 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.

```
w2b_table<-aggregate(wing2body~sex_binom*pophost_binom*month_of_year*months_since_start,
                     data=data_long, FUN=mean)
w2b_table$sd<-aggregate(wing2body~sex_binom*pophost_binom*month_of_year*months_since_start,
                           data=data_long, FUN=sd) $wing2body
w2b_table$se<-aggregate(wing2body~sex_binom*pophost_binom*month_of_year*months_since_start,
                           data=data_long, FUN=SE) $wing2body
data = w2b 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]
## AICs
          -280.1872 -279.8713 -279.4758
## models 8
                    19
                               2
## probs 0.1198675 0.1023577 0.08398967
##
## m8
        glm(formula = R ~ B + C, family = gaussian, data = data)
        glm(formula = R ~ B * C, family = gaussian, data = data)
```

```
glm(formula = R ~ B, family = gaussian, data = data)
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 0.2157
## 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)
## 1
           35 0.00087733
## 2
           34 0.00075844 1 0.00011888 0.02097 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Best Fit
M6 = glm(sd ~ pophost_binom, data=w2b_table, family=gaussian)
summary(M6)
##
## glm(formula = sd ~ pophost_binom, family = gaussian, data = w2b_table)
## Deviance Residuals:
         Min
                      1Q
                              Median
                                              3Q
                                                         Max
## -0.0059374 -0.0033018 -0.0006274
                                       0.0022332
                                                   0.0147212
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
                0.0165999 0.0007872 21.088
## (Intercept)
                                               <2e-16 ***
## pophost_binom 0.0018172 0.0007872
                                       2.309
                                               0.0272 *
## Signif. codes: 0 '***' 0.001 '**' 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

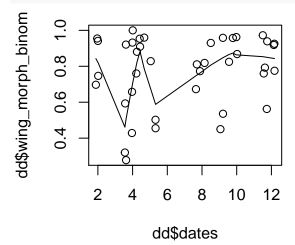
Group significant elements

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)
}

l1 = 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(l1, type = "l")
    plot_lowess_residuals(l1, dd$dates, dd$wing_morph_binom)</pre>
```



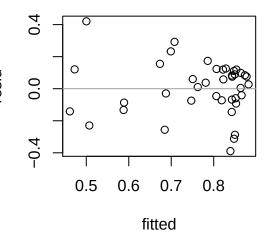
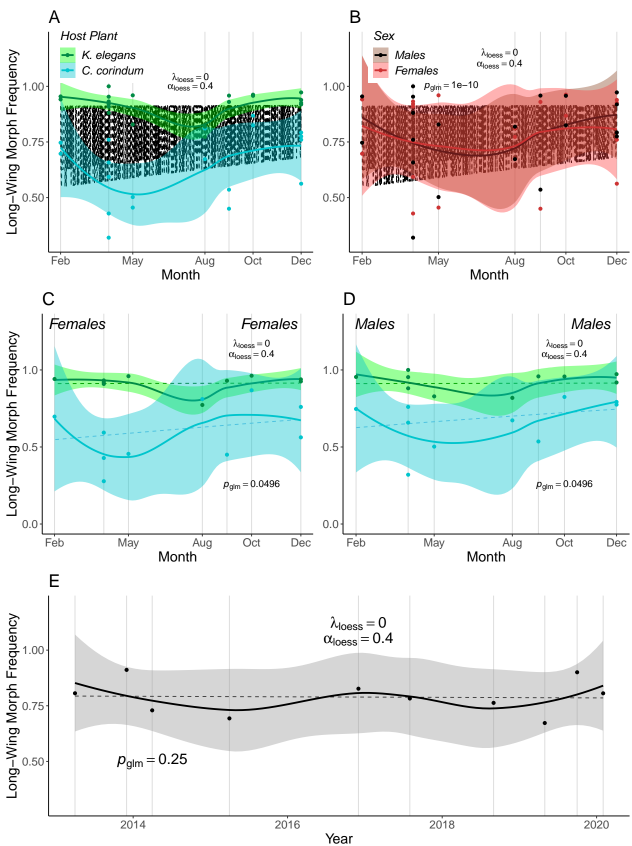


Figure: Panels A, B, C, D (long-wing morph freq with month) & E (long-wing morph freq with year)



Wing-to-body Ratio

Check for LOESS Residuals

```
11 = lowess(d$dates, d$wing2body, f=0.4)
par(mfrow=c(1,2))
plot(d$dates, d$wing2body)
lines(11, type = "1")
plot_lowess_residuals(11, d$dates, d$wing2body)
                                                                                               \overline{\circ}
      0.740
                                                                                      0
                                       0
                 0
d$wing2body
      0.725
                                                                                               0
                                                                            8
                 8
                                0
                                                                     0
                                         0
                     0
                                                           .015
      0.710
                                           0
                      2016
                                                                 0.726
                                                                              0.730
                                                                                           0.734
              2014
                               2018
                                        2020
                        d$dates
                                                                              fitted
```

Figure: Panel A & B (wing-to-body with month)

```
w2b_summary<-aggregate(wing2body~sex*pophost*month_of_year, data=data_long, FUN=mean)
df = w2b_summary
df$pophost = factor(df$pophost, levels = c("K. elegans", "C. corindum") )
df$`Host Plant` = df$pophost
df$sex[df$sex=="F"]<-"Females"
df$sex[df$sex=="M"]<-"Males"</pre>
df$sex = factor(df$sex, levels = c("Males", "Females") )
df$`Sex` = df$sex
p1 = ggplot() +
  ggtitle("A") + xlab("Month") + ylab("Wing-to-Body Ratio") +
  geom vline(xintercept = xlab allmonths, color="gainsboro") +
  geom_smooth(data=data_long, method="glm", se=FALSE, linetype = "dashed",
              mapping = aes(x = month_of_year, y = wing2body), colour="black", lwd=0.5) +
  geom_smooth(data=df, method="loess",
              mapping = aes(x = month_of_year, y = wing2body, colour=`Host Plant`, fill=`Host Plant`))
  geom_point(data=df, mapping = aes(x = month_of_year, y = wing2body, colour=`Host Plant`)) +
```

```
ylim(0.71, 0.75) +
  customPlot +
  scale_color_manual(values=c("C. corindum" = "turquoise3", "K. elegans" = "springgreen4")) +
  scale_fill_manual(values = c("C. corindum" = "turquoise3", "K. elegans" = "green")) +
  scale_x_continuous(breaks=xlab_months, labels=month_labs)
p2 = ggplot() +
  ggtitle("B") + xlab("Month") + ylab("Wing-to-Body Ratio") +
  geom_vline(xintercept = xlab_allmonths, color="gainsboro") +
  geom_smooth(data=data_long, method="glm", se=FALSE, linetype = "dashed",
             mapping = aes(x = month_of_year, y = wing2body), colour="black", lwd=0.5) +
  geom_smooth(data=df, method="loess",
             mapping = aes(x = month_of_year, y = wing2body, colour=Sex, fill=Sex)) +
  geom_point(data=df, mapping = aes(x = month_of_year, y = wing2body, colour=Sex)) +
  ylim(0.71, 0.75) +
  customPlot +
  scale_color_manual(values=c("Females" = "brown3", "Males" = "black")) +
  scale_fill_manual(values = c("Females" = "brown1", "Males" = "sienna4")) +
  scale_x_continuous(breaks=xlab_months, labels= month_labs) + theme(axis.title.y = element_blank())
# Extract from LOESS and linear regressions
alpha = paste("alpha[loess] == ", ggplot_build(p1)$data[[3]]$alpha[1])
## `geom_smooth()` using formula 'y ~ x'
## Warning: Removed 473 rows containing non-finite values (stat_smooth).
## `geom_smooth()` using formula 'y ~ x'
degree="lambda[loess] == 0"
mlinear = glm(wing2body ~ month_of_year, data=df, family=gaussian) # aggregated df
summary(M4) # individual bugs df (best fit model)
##
## Call:
## glm(formula = wing2body_c ~ sex_binom * pophost_binom + month_of_year_c,
##
      family = gaussian, data = data_long)
##
## Deviance Residuals:
##
        Min
                           Median
                                         30
                                                   Max
                    1Q
## -0.070223 -0.010939 -0.000117
                                   0.010773
                                              0.114608
##
## Coefficients:
                            Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                          -0.0007445 0.0004473 -1.664
                                                         0.0962 .
                          ## sex binom
## pophost_binom
                           0.0043900 0.0004497
                                                 9.762 < 2e-16 ***
## month_of_year_c
                           ## sex_binom:pophost_binom 0.0018351 0.0004468 4.108 4.17e-05 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for gaussian family taken to be 0.0003533471)
##
##
      Null deviance: 0.72538 on 1902 degrees of freedom
## Residual deviance: 0.67065 on 1898 degrees of freedom
```

```
## AIC: -9717.7
##
## Number of Fisher Scoring iterations: 2

M4_beta = summary(M4)$coeff[,"Estimate"][4]
M4_pvalue = round(summary(M4)$coeff[,"Pr(>|t|)"][4],10)
pvalue = paste0("italic(p)[glm]==", M4_pvalue)
```