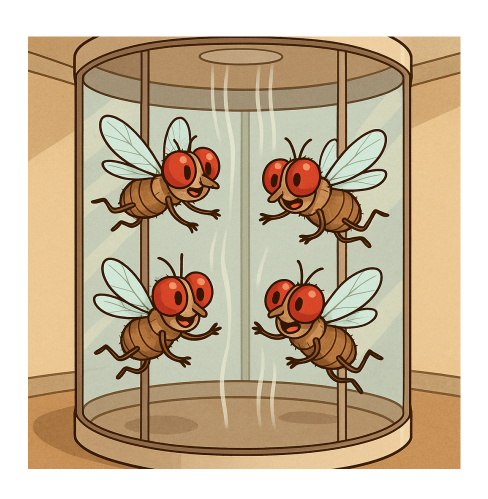
Statistical Analysis of Flight Assay Data in R Jun Ishigohoka



Environment

We recommend you use Rstudio.

- Download flight_assay_analysis.Rmd from https://github.com/PallaresLab/EMB L_Drosophila_course_2025/blob/main/scripts/flight_assay_analysis.Rmd, place it in the same directory as your flight assay data.
- Open flight assay analysis. Rmd in Rstudio.
- Follow the instructions

Or

- PDF: download from here
- HTML: access here

Preparation

If you don't have the packages, install them.

```
packages <- c("ggplot2", "car")
for (pkg in packages) {
   if (!requireNamespace(pkg, quietly = TRUE)) {
     install.packages(pkg)
   }
}</pre>
```

Load libraries.

```
library(ggplot2)
library(car) # Anova() and leveneTest()
```

Read data in two data frames.

Edit the file names accordingly.

```
d_1 <- read.csv("data/test_data/R4_female.csv")
d_2 <- read.csv("data/test_data/R4_male.csv")
head(d_1)</pre>
```

```
##
    X.1
                      Х
                             Y
           Area
## 1
       1 5.456 407.768 9.192
## 2
       2 6.666 343.943 10.074
       3 11.087 306.180 11.690
## 3
## 4
      4 4.932 376.921 16.866
       5 4.569 148.680 25.018
## 5
## 6
       6 5.389 330.855 26.006
```

Add a column for the factor you are comparing.

Edit the name of factor (sex) and each level of factor (female/male) accordingly.

```
d 1$sex <- "female"
d 2$sex <- "male"
```

```
What are the mean and standard deviation of landing height in each group?
mean(d 1$Y)
## [1] 200.7005
sd(d 1$Y)
## [1] 144.5707
mean(d 2\$Y)
## [1] 182.0229
sd(d_2\$Y)
## [1] 136.5356
```

Let's make these two data frame together. Concatenate the two data frames. Then set it as a factor.

Edit the name of factor (sex) and each level of the factor (female/male) accordingly.

```
d <- rbind(d 1, d 2)
d$sex <- factor(d$sex, levels = c("female", "male"))</pre>
```

Visualisation

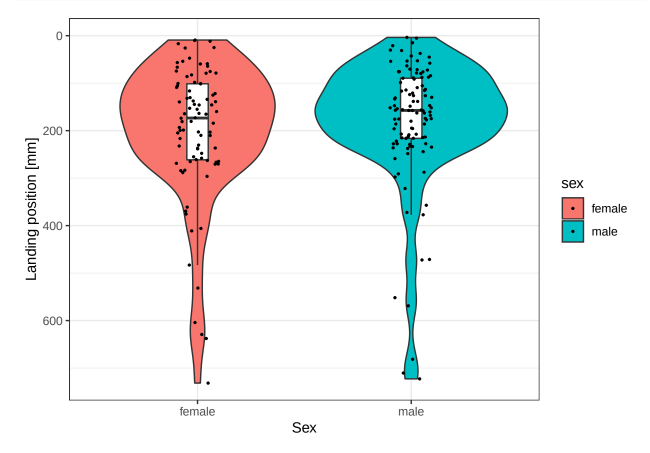
Let's visualise the distribution of landing height. Once you have visualised the distributions, try to describe them.

If you have ggplot2...

(Edit the name of factor (sex), each level of the factor (female/male), and unit (mm) accordingly.)

```
ggplot(data = d, # data to be plotted
         mapping = aes(x=sex, y=Y, fill = sex)) + # columns in the data frame to be used
          geom_violin() + # violin plot
3
          geom_boxplot(
                            # box plot
            width = 0.1,
                           # width of the box plot
5
            fill = "white", # colour inside the box plot
                            # whether to have points for outliers.
            outliers = F
                            # FALSE because of geom_fitter() below
```

```
# each data point with "jitter" along x axis
           geom_jitter(
10
                              # size of points
             size = 0.5,
11
             width = 0.1
                              # the amount of jitter
^{12}
             ) +
13
           scale_y_reverse( # to make the plot same direction as the tapes
14
             limits = c(NA, 0) # so that the lowest Y is 0
15
16
     labs(
                              # Add labels
17
       x = "Sex",
18
       y = "Landing position [mm]") +
19
     theme_bw()
20
```



If you prefer the base plot or do not have ggplot2 installed....

(Edit the name of factor (sex), each level of the factor (female/male), and unit (mm) accordingly.)

```
boxplot(d$Y ~ as.factor(d$sex),

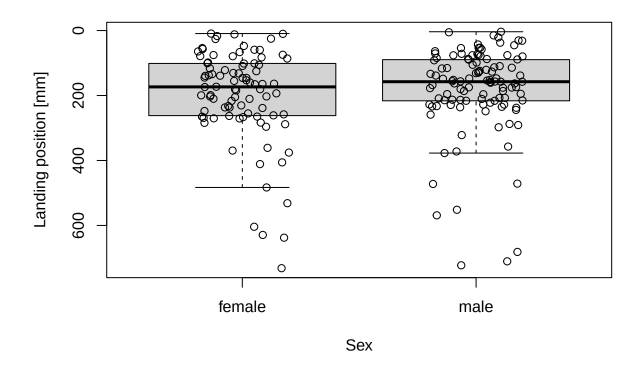
pch = NA,

xlab = "Sex",

ylab = "Landing position [mm]",

ylim = rev(range(d$Y))
```

```
6 )
7 points(jitter(as.numeric(as.factor(d$sex))), d$Y)
```



Statistical analysis (hypothesis testing)

Are your data normally distributed?

Many parametric tests assume normality. Let's run Shapiro-Wilk test for each group to test normality.

Edit the name of factor (sex), each level of the factor (female/male) accordingly.

```
shapiro.test(subset(d, sex == "female")$Y)

##

## Shapiro-Wilk normality test

##

## data: subset(d, sex == "female")$Y

## W = 0.87191, p-value = 3.497e-07

shapiro.test(subset(d, sex == "male")$Y)
```

```
##
## Shapiro-Wilk normality test
##
## data: subset(d, sex == "male")$Y
## W = 0.80816, p-value = 1.326e-10
```

Does the result fit your description of the distributions? If they are not normally distributed, think why. Do you expect normally distributed data from this experiment?

Does your data have equal variance?

Many parametric tests assume equal variance between groups.

Let's run two tests for homogeneity: F-test and Brown-Forsythe test.

Run F-test.

Edit the name of factor (sex) accordingly.

```
##
## F test to compare two variances
##
## data: Y by sex
## F = 1.1212, num df = 87, denom df = 108, p-value = 0.5699
## alternative hypothesis: true ratio of variances is not equal to 1
## 95 percent confidence interval:
## 0.7535657 1.6828055
## sample estimates:
## ratio of variances
## 1.121162
```

Run Brown-Forsythe test (car needs to be installed).

Edit the name of factor (sex) accordingly.

```
leveneTest(Y ~ sex, d, center = median)

## Levene's Test for Homogeneity of Variance (center = median)

## Df F value Pr(>F)

## group 1 0.8082 0.3698

## 195
```

If you do not have car installed, you can run Brown-Forsythe test manually. Compute absolute deviance from median for each sex manually, and run ANOVA.

Edit the name of factor (sex) accordingly.

```
d$z <- NA
for (sex in levels(d$sex)) {
```

```
idx <- which(d$sex == sex)
    d[idx,]$z <- abs(d[idx, ]$Y - median(d[idx, ]$Y))
  }
5
6
  summary(aov(d$z ~ d$sex))
                       Sum Sq Mean Sq F value Pr(>F)
  ##
                   Df
  ## d$sex
                          9114
                                  9114
                                         0.808
                    1
  ## Residuals
                  195 2199050
                                 11277
```

Is the mean landing distance different between groups?

If the data are normally distributed and have equal variance, student t-test

182.0229

If data are normally distributed but not with equal variance, Welch's t-test

mean in group male

Edit the name of factor (sex) accordingly.

200.7005

mean in group female

##

```
## -21.20353 58.55868

## sample estimates:

## mean in group female mean in group male

## 200.7005 182.0229
```

If data are not normally distributed, you can run non-parametric tests or permutation test.

Option 1: Non-parametric test

Let's run Mann-Whitney U test.

Edit the name of factor (sex) accordingly.

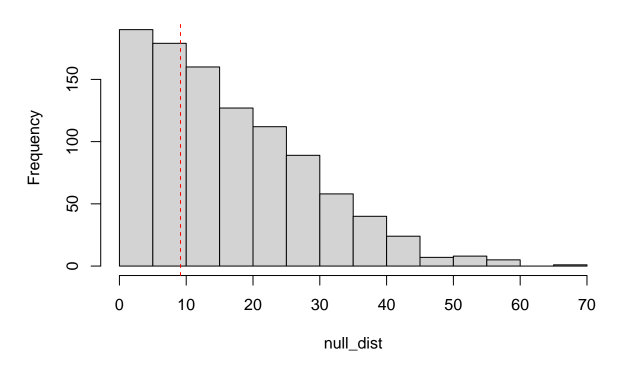
Option 2: Permutation test

Let's write a function to run a permutation test. Briefly, we shuffle the label, compute difference between groups. We repeat this process many times to get a null distribution. We compute p-value based on the position of the observed difference between groups in the null distribution.

Edit the name of factor (sex) and each level of the factor (female/male) accordingly.

```
# Null distribution of abs(mean(Y | female) - mean(Y | male))
   null dist <- sapply(1:1000, # repeat 1000 times
          function(x){
3
            d tmp <- d
4
            d tmp$sex <- sample(d$sex) # shuffle label
5
            return(abs(mean(d tmp[d tmp$sex == "female","Y"])
                     - mean(d tmp[d tmp$sex == "male","Y"]))
              )
          }
9
          )
10
11
   # Observed value of mean(female) - mean(male)
12
   obs <- abs(mean(d[d$sex == "female", "Y"] - d[d$sex == "male", "Y"]))
13
   # p value as the rank of observation in the null distribution
15
```

p_val: 0.657



Optional: Statistical modeling

Here, we conduct statistical modeling of landing distance using a generalised linear model (GLM) approach with the exponential distribution, considering landing events as a Poisson process with a constant "landing rate". Because the waiting time of a Poisson process is an exponentially distributed random variable, we can use a GLM with exponential distribution. We compare the result with a linear model (LM), which assumes a normal distribution as the underlying distribution.

GLM

Fit the data to an exponential GLM.

Edit the name of factor (sex) accordingly.

```
glm_1 <- glm(Y ~ sex,
data = d,
family = Gamma(link = "log") # This is how to specify exponential
)</pre>
```

According to the fitted GLM, do the two groups have different landing rate?

If you have car installed

```
1 Anova(glm 1)
  ## Analysis of Deviance Table (Type II tests)
  ##
  ## Response: Y
         LR Chisq Df Pr(>Chisq)
  ## sex 0.85803 1
                         0.3543
 summary(glm_1, dispersion = 1) # exponential is a Gamma with dispersion = 1
  ##
  ## Call:
  ## glm(formula = Y ~ sex, family = Gamma(link = "log"), data = d)
  ## Coefficients:
                 Estimate Std. Error z value Pr(>|z|)
  ## (Intercept) 5.30181
                             0.10660 49.735
                                               <2e-16 ***
                 -0.09768
                                                0.495
  ## sexmale
                             0.14331
                                     -0.682
  ## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
  ## (Dispersion parameter for Gamma family taken to be 1)
  ##
  ##
         Null deviance: 111.53 on 196 degrees of freedom
  ## Residual deviance: 111.07 on 195 degrees of freedom
  ## AIC: 2427.1
  ##
  ## Number of Fisher Scoring iterations: 5
  If you do not have car installed
drop1(glm_1, test = "Chi")
  ## Single term deletions
  ##
  ## Model:
  ## Y ~ sex
                           AIC scaled dev. Pr(>Chi)
            Df Deviance
                 111.07 2427.1
  ## <none>
```

```
## sex 1 111.53 2425.9 0.85803 0.3543
```

LM

Fit the data to a linear model.

```
Edit the name of factor (sex) accordingly.
 lm 1 \leftarrow lm(Y \sim sex,
               data = d
  If you do not have car installed
drop1(lm_1, test = "Chi")
  ## Single term deletions
  ##
  ## Model:
  ## Y ~ sex
            Df Sum of Sq
                              RSS
                                     AIC Pr(>Chi)
                          3831693 1949.5
  ## <none>
  ## sex
                    16986 3848679 1948.4
                                            0.3506
  summary(lm 1)
  ##
  ## Call:
  ## lm(formula = Y ~ sex, data = d)
  ##
  ## Residuals:
  ##
         Min
                   1Q Median
                                    3Q
                                           Max
  ## -191.51 -94.30 -25.02 45.70 540.72
  ##
  ## Coefficients:
                  Estimate Std. Error t value Pr(>|t|)
  ## (Intercept)
                    200.70
                                14.94
                                         13.43
                                                 <2e-16 ***
  ## sexmale
                   -18.68
                                20.09 -0.93
                                                  0.354
  ## ---
```

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

Multiple R-squared: 0.004413, Adjusted R-squared: -0.0006922

Residual standard error: 140.2 on 195 degrees of freedom

F-statistic: 0.8644 on 1 and 195 DF, p-value: 0.3537

GLM vs LM

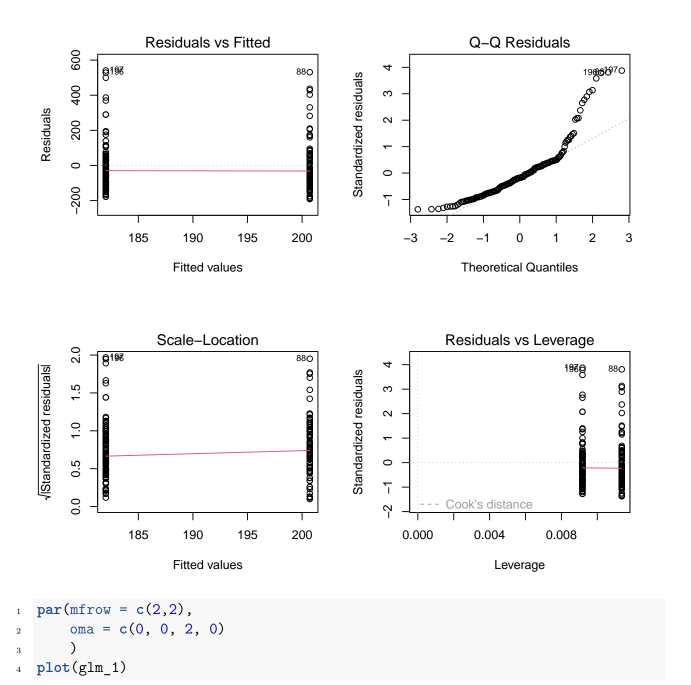
Compare the diagnostic plots¹ between glm_1 and lm_1. Is using LM for landing distance data worse than exponential GLM?

- Is the linearity assumption met?
- Is the data homoscedastic?
- Are residuals distributed normally?
- Are there outliers?

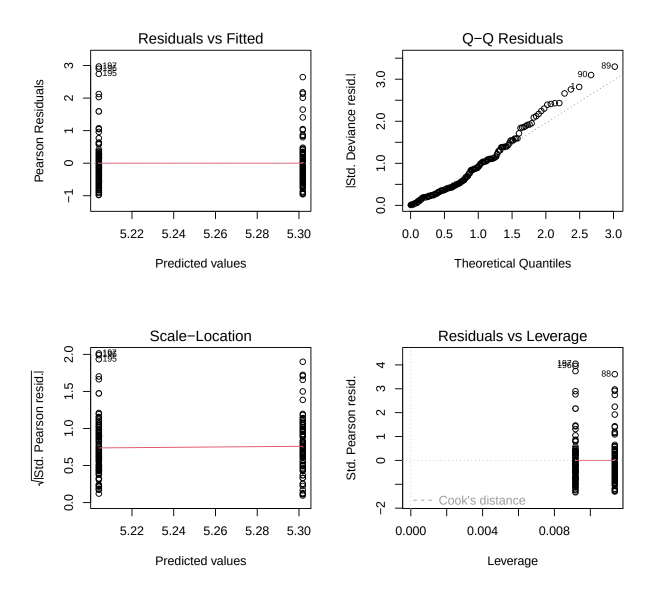
```
par(mfrow = c(2,2),
    oma = c(0, 0, 2, 0)
    )
plot(lm_1)
```

 $^{^{1}\}mbox{What do the diagnostic plots mean? https://www.sthda.com/english/articles/39-regression-model-diagnostics/161-linear-regression-assumptions-and-diagnostics-in-r-essentials/}$

$Im(Y \sim sex)$



$glm(Y \sim sex)$



What's next?

In real life, we have multiple biological and technical replicates. How do we account for such random effects?

Further reading

- Whitlock & Schluter. The Analysis of Biological Data 3rd edition. Macmillan Learning. 2020
- Zuur et al. Mixed Effects Models and Extensions in Ecology with R. Springer. 2009.