## HR\_Va\_h2\_R\_2022

#### Helen Payne

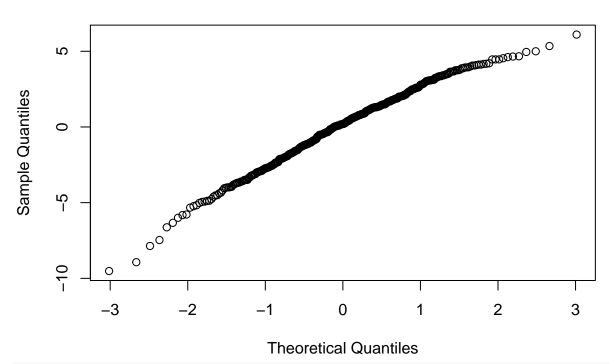
2024-06-18

```
#load pHRkages
library(lme4)
## Loading required package: Matrix
library(tidyverse)
## -- Attaching core tidyverse packages ----
                                                  ----- tidyverse 2.0.0 --
## v dplyr
             1.1.4
                                     2.1.5
                        v readr
             1.0.0
## v forcats
                        v stringr
                                     1.5.1
## v ggplot2 3.4.4
                       v tibble
                                     3.2.1
## v lubridate 1.9.3
                         v tidyr
                                     1.3.0
## v purrr
              1.0.2
## -- Conflicts ----- tidyverse_conflicts() --
## x tidyr::expand() masks Matrix::expand()
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                    masks stats::lag()
## x tidyr::pack() masks Matrix::pack()
## x tidyr::unpack() masks Matrix::unpack()
## i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become error
library(lmerTest)
##
## Attaching package: 'lmerTest'
## The following object is masked from 'package:lme4':
##
##
       lmer
## The following object is masked from 'package:stats':
##
##
       step
library(car)
## Loading required package: carData
##
## Attaching package: 'car'
## The following object is masked from 'package:dplyr':
##
##
       recode
##
```

```
## The following object is masked from 'package:purrr':
##
##
################################
Read in the data:
HR_22_fit <- read_csv(here::here("data_sheets", "compiled_sheets", "HR_mastersheet_Fitness-mains_2022.c
## Rows: 406 Columns: 57
## -- Column specification -------
## Delimiter: ","
         (5): Donor, Recipient, Gen, Replicated, Needed Area Redo
        (43): Year, Sequence, Cohort, Block, Transect, Plant_ID, F_plant_ID, fl...
## dbl
        (1): F_multi
## num
         (3): F_plant, Rep_FitP, any_FitP
## lgl
## date (5): Germ_Date, Sow_Date, Plant_Date, FFD, LFD
## i Use `spec()` to retrieve the full column specification for this data.
## i Specify the column types or set `show_col_types = FALSE` to quiet this message.
#log scale traits that are highly skewed
HR_22_fit$skel_dryweight_mg_SEG <- log(HR_22_fit$skel_dryweight_mg_SEG)
HR_22_fit$est_fecundity <- sqrt(HR_22_fit$est_fecundity + 1)</pre>
HR 22 fit$SLA SEG <- log(HR 22 fit$SLA SEG)
#mean center the traits of interest
traits <- c("corolla_diam_mm_SEG", "skel_dryweight_mg_SEG", "fl_duration", "est_fecundity", "msm_all",
# Mean center eHRh trait
for (trait in traits) {
 trait_mean <- mean(HR_22_fit[[trait]], na.rm = TRUE)</pre>
 HR_22_fit[[paste0(trait, "_centered")]] <- HR_22_fit[[trait]] - trait_mean</pre>
# Create the mixed model for corolla area
\#corolla\_model \leftarrow lmer(corolla\_diam\_mm\_SEG\_centered \sim (1 \mid Recipient) + (1 \mid Donor), data = HR\_22\_fit)
corolla_model <- lmer(corolla_diam_mm_SEG_centered ~ (1 | Donor), data = HR_22_fit)
rand(corolla_model)#including Donor and Transect as random effects significantly improves the models fi
## ANOVA-like table for random-effects: Single term deletions
##
## Model:
## corolla diam mm SEG centered ~ (1 | Donor)
##
              npar logLik
                              AIC LRT Df Pr(>Chisq)
## <none>
                 3 -949.59 1905.2
## (1 | Donor)
                 2 -952.50 1909.0 5.827 1
                                              0.01578 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# ExtrHRt residuals from the model
residuals <- resid(corolla_model)</pre>
```

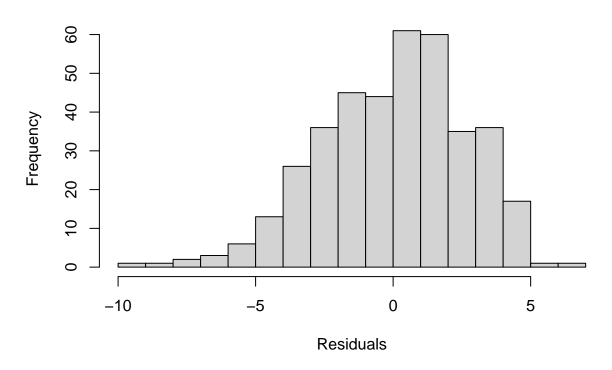
# Q-Q plot for normality
qqnorm(residuals) #looks good

# Normal Q-Q Plot



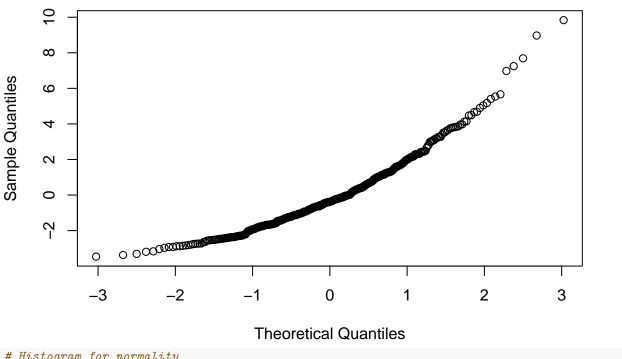
# Histogram for normality
hist(residuals, breaks = 20, main = "Histogram of Residuals", xlab = "Residuals") #normal

# **Histogram of Residuals**



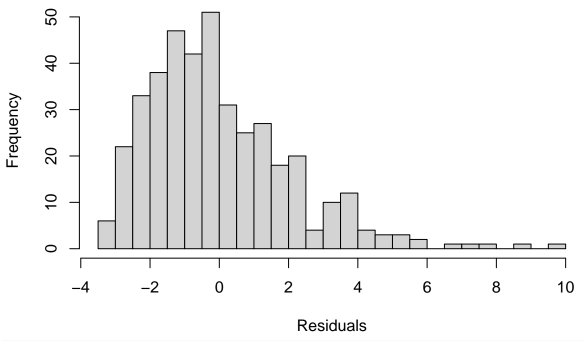
```
#use Model 2 (Donor)
# Create the mixed model for skeleton weight, with skeleton weight log transformed
\#skel\_model \leftarrow lmer((skel\_dryweight\_mg\_SEG\_centered) \sim (1 \mid Recipient) + (1 \mid Donor), data = HR\_22\_fit)
\#skel\_model \leftarrow lmer((skel\_dryweight\_mg\_SEG\_centered) \sim (1 \mid Donor), data = HR\_22\_fit) \#singular
# Create the mixed model for flowering duration
\#fl\_duration\_model \leftarrow lmer(fl\_duration\_centered \sim (1 \mid Recipient) + (1 \mid Donor), \ data = HR\_22\_fit) \ \#Sin(lmer)
# Create the mixed model for flowering duration
\#fl\_duration\_model \leftarrow lmer(fl\_duration\_centered \sim (1 \mid Donor), data = HR\_22\_fit) \#Singular
# Create the mixed model for estimated fecundity, sqrt transforming estimated fecundity
est_fecundity_model <- lmer((est_fecundity_centered) ~ (1 Transect) + (1 | Recipient) + (1 | Donor), da
rand(est_fecundity_model)
## ANOVA-like table for random-effects: Single term deletions
##
## Model:
## (est_fecundity_centered) ~ (1 | Transect) + (1 | Recipient) + (1 | Donor)
##
                    npar logLik
                                     AIC
                                              LRT Df Pr(>Chisq)
## <none>
                       5 -899.92 1809.8
## (1 | Transect)
                       4 -905.03 1818.1 10.2260 1
                                                        0.001385 **
                       4 -900.34 1808.7 0.8386 1
## (1 | Recipient)
                                                        0.359810
## (1 | Donor)
                       4 -899.92 1807.8 0.0063 1
                                                        0.936887
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# ExtrHRt residuals from the model
residuals <- resid(est_fecundity_model)</pre>
# Q-Q plot for normality
qqnorm(residuals) #good enough
```

### Normal Q-Q Plot



# Histogram for normality
hist(residuals, breaks = 20, main = "Histogram of Residuals", xlab = "Residuals") #normal-ish

## **Histogram of Residuals**

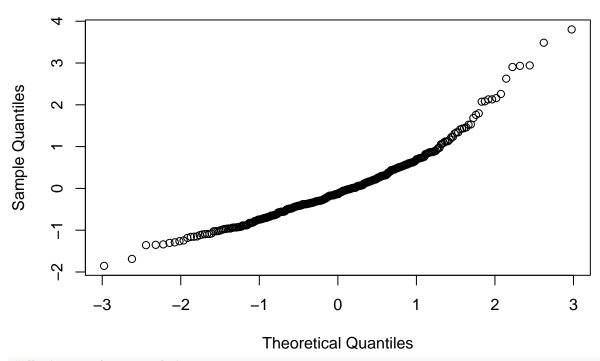


# Create the mixed model for mean seed mass, log transformed mean seed mass
msm\_model <- lmer((msm\_all\_centered) ~ (1| Transect) + (1 | Recipient) + (1 | Donor), data = HR\_22\_fit)</pre>

```
# ExtrHRt residuals from the model
residuals <- resid(msm_model)

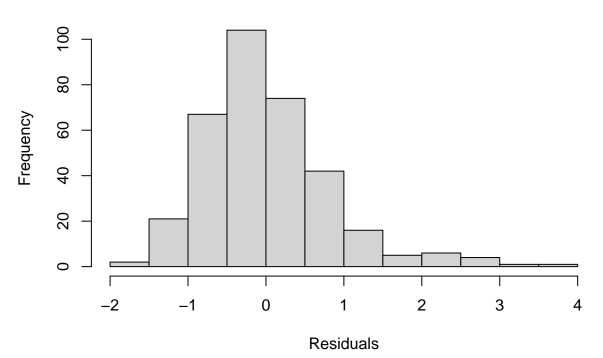
# Q-Q plot for normality
qqnorm(residuals) #good enough</pre>
```

## Normal Q-Q Plot



# Histogram for normality
hist(residuals, breaks = 20, main = "Histogram of Residuals", xlab = "Residuals") #normal-ish

### **Histogram of Residuals**

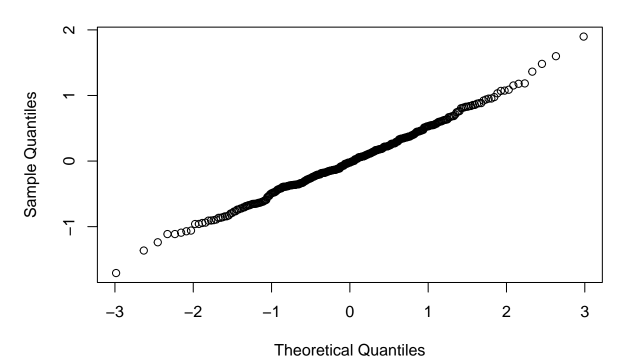


# Test the significance of the random effects
rand(msm\_model) #none of these random effects significantly improve the model's fit

```
## ANOVA-like table for random-effects: Single term deletions
##
## Model:
## (msm_all_centered) ~ (1 | Transect) + (1 | Recipient) + (1 | Donor)
##
                                                   npar logLik
                                                                                                AIC
                                                                                                                      LRT Df Pr(>Chisq)
## <none>
                                                            5 -477.78 965.55
                                                            4 -483.77 975.53 11.9810 1 0.0005375 ***
## (1 | Transect)
                                                            4 -478.07 964.13 0.5823
                                                                                                                                           0.4454056
## (1 | Recipient)
                                                                                                                                   1
## (1 | Donor)
                                                            4 -481.21 970.42 6.8727
                                                                                                                                 1 0.0087524 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# Create the mixed model for SLA
\#SLA\_model \leftarrow lmer((SLA\_SEG\_centered) \sim (1/Transect) + (1 | Recipient) + (1 | Donor), data = HR\_22\_fit)
\#SLA\_model \leftarrow lmer((SLA\_SEG\_centered) \sim (1 \mid Donor), data = HR\_22\_fit) \#Singular
# Create the mixed model for mean seed mass, log transformed mean seed mass
\#LMA\_model \leftarrow lmer(LMA\_SEG\_centered \sim (1/Transect) + (1 | Recipient) + (1 | Donor), data = HR\_22\_fit) \#LMA\_model \leftarrow lmer(LMA\_SEG\_centered \sim (1/Transect) + (1 | Recipient) + (1 | Donor), data = HR\_22\_fit) \#LMA\_model \leftarrow lmer(LMA\_SEG\_centered \sim (1/Transect) + (1 | Recipient) + (1 | Donor), data = HR\_22\_fit) \#LMA\_model \leftarrow lmer(LMA\_SEG\_centered \sim (1/Transect) + (1 | Recipient) + (1 | Donor), data = HR\_22\_fit) \#LMA\_model \leftarrow lmer(LMA\_SEG\_centered \sim (1/Transect) + (1 | Recipient) + (1 | Donor), data = HR\_22\_fit) \#LMA\_model \leftarrow lmer(LMA\_SEG\_centered \sim (1/Transect) + (1 | Recipient) + (1 | Donor), data = HR\_22\_fit) \#LMA\_model \leftarrow lmer(LMA\_SEG\_centered \sim (1/Transect) + (1 | Recipient) + (1 | Donor), data = HR\_22\_fit) \#LMA\_model \leftarrow lmer(LMA\_SEG\_centered \sim (1/Transect) + (1 | Recipient) + (1 | Donor), data = HR\_22\_fit) \#LMA\_model \leftarrow lmer(LMA\_SEG\_centered \sim (1/Transect) + (1 | Recipient) + (1 | Recipient
# Create the mixed model for mean seed mass, log transformed mean seed mass
\#LMA\_model \leftarrow lmer(LMA\_SEG\_centered \sim (1 \mid Donor), data = HR\_22\_fit) \#Singular
# Create the mixed model for d13C, log transformed mean seed mass
d13C_model <- lmer(d13C_SEG_centered ~ (1|Transect) + (1 | Recipient) + (1 | Donor), data = HR_22_fit)
# ExtrHRt residuals from the model
residuals <- resid(d13C_model)</pre>
```

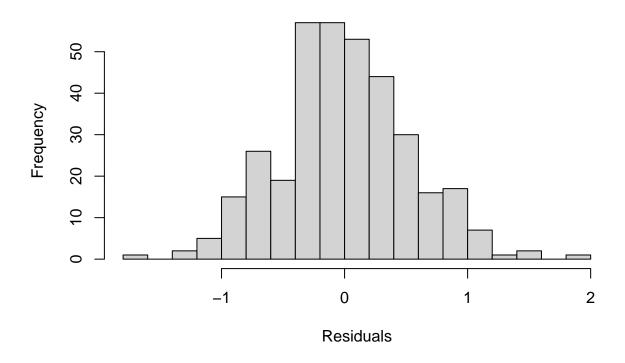
# Q-Q plot for normality
qqnorm(residuals) #good enough

# Normal Q-Q Plot



# Histogram for normality
hist(residuals, breaks = 20, main = "Histogram of Residuals", xlab = "Residuals") #normal-ish

# **Histogram of Residuals**



```
rand(d13C_model) #all Random effects significantly improve the models fit!
## ANOVA-like table for random-effects: Single term deletions
##
## Model:
## d13C_SEG_centered ~ (1 | Transect) + (1 | Recipient) + (1 | Donor)
##
                   npar logLik
                                    AIC
                                            LRT Df Pr(>Chisq)
## <none>
                      5 -344.48 698.96
## (1 | Transect)
                      4 -355.89 719.79 22.8310 1 1.769e-06 ***
## (1 | Recipient)
                      4 -344.70 697.39 0.4387 1
                                                       0.5077
                      4 -354.53 717.05 20.0954 1 7.367e-06 ***
## (1 | Donor)
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# Function to extrHRt variance components and calculate required values
calculate_variances <- function(model, trait_name) {</pre>
  var_components <- as.data.frame(VarCorr(model))</pre>
  # Initialize variables
  Va mat <- NA
  Va_sd_mat <- NA
  Va_pat <- NA
  Va_sd_pat <- NA
  res_var <- NA
  Vp <- NA
  Vp_sd <- NA
  h2 <- NA
  # Check if "Recipient" is included as a random effect
  if ("Recipient" %in% var_components$grp) {
    Va_mat <- (var_components$vcov[var_components$grp == "Recipient"]) * 4</pre>
    Va_sd_mat <- sqrt(Va_mat)</pre>
  }
  # Calculate other variance components
  if ("Donor" %in% var_components$grp) {
    Va_pat <- (var_components$vcov[var_components$grp == "Donor"]) * 4</pre>
    Va_sd_pat <- sqrt(Va_pat)</pre>
  if ("Residual" %in% var components$grp) {
    res_var <- (var_components$vcov[var_components$grp == "Residual"])</pre>
  # Calculate total phenotypic variance and narrow-sense heritability if components are available
  if (!is.na(Va_mat) & !is.na(Va_pat) & !is.na(res_var)) {
    Vp <- Va_mat + Va_pat + res_var</pre>
    Vp_sd <- sqrt(Vp)</pre>
    h2 <- (Va_pat) / Vp # assumed calculation
  }
  # Create the dataframe and add the traits column
  df <- data.frame(traits = trait_name, Va_mat, Va_sd_mat, Va_pat, Va_sd_pat, Vp, Vp_sd, h2)
```

# Test the significance of the random effects

```
return(df)
}
# Calculate variances for eHRh model and add trait names
corolla_variances <- calculate_variances(corolla_model, 'corolla_diameter')</pre>
#skel_variances <- calculate_variances(skel_model, "skel_biomass_mg")</pre>
est_fecundity_variances <- calculate_variances(est_fecundity_model, "estimated_fecundity")</pre>
msm variances <- calculate variances(msm model, "mean seed mass")</pre>
#SLA variances <- calculate variances(SLA model, "SLA")
#LMA_variances <- calculate_variances(LMA_model, "LMA")</pre>
d13C_variances <- calculate_variances(d13C_model, "delta_C_13")</pre>
# Combine the results into a single dataframe
variance_HR_2022 <- rbind(</pre>
  corolla_variances,
  d13C_variances,
  est_fecundity_variances,
  msm_variances
)
# Print the dataframe
print(variance_HR_2022)
                  traits
                              Va_mat Va_sd_mat
                                                    Va_pat Va_sd_pat
                                                                             Vр
## 1
                                             NA 2.28574571 1.5118683
        corolla_diameter
                                  NA
                                                                             NA
              delta_C_13 0.06168231 0.2483592 0.38907052 0.6237552 0.7761152
## 3 estimated_fecundity 1.06091705 1.0300083 0.05140342 0.2267232 5.7984106
          mean_seed_mass 0.19023736 0.4361621 0.48682339 0.6977273 1.4638778
##
        Vp_sd
                        h2
## 1
           NA
## 2 0.880974 0.501305080
## 3 2.407989 0.008865088
## 4 1.209908 0.332557388
#Save the csv file if you want
write_csv(x = variance_HR_2022, here::here("data_sheets", "compiled_sheets", "HR_Va_h2_R_2022.csv"))
```