Natural Populations Preliminary Results

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Summary results

Data cleaning

I did a series of PCAs using the variables that were measured continously between 2017 - 2019. I choose these years because 2014 - 2016 were already analysed by Sofia Carvajal. Although we can include these in further analysis (PTA) or selection estimates. I removed NAs, check for outliers and corrected some values from the raw data. The clean subset of data was saved as a second table called TribulusPC (https://github.com/Winer-DanielR/Tribulus-Natural-population/tree/master/Data/Processed) this new dataset was used for the following analysis.

General PCA

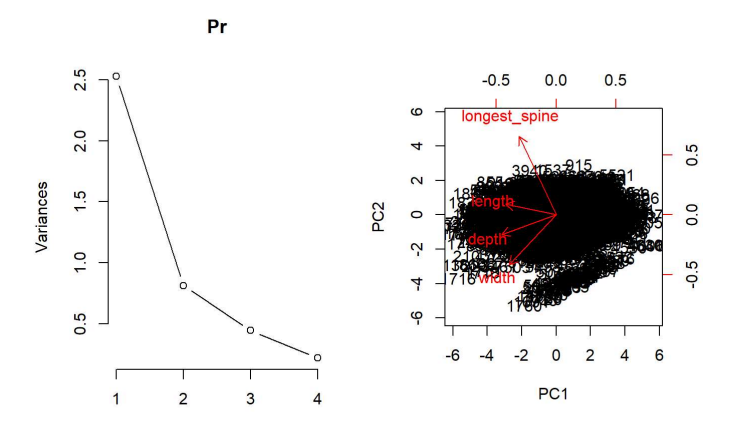
The PCA used the following variables:

- length
- width
- depth
- · longest spine

I did not include spine tip distance since I did not measure that trait in 2019 due to the measurements involving a lot of guessing given that most of the mericarps did not have completed spines.

PCA summary

```
## Importance of components:
##
                             PC1
                                    PC2
                                           PC3
                                                   PC4
## Standard deviation
                          1.5897 0.9011 0.6657 0.46669
## Proportion of Variance 0.6318 0.2030 0.1108 0.05445
## Cumulative Proportion 0.6318 0.8347 0.9456 1.00000
```

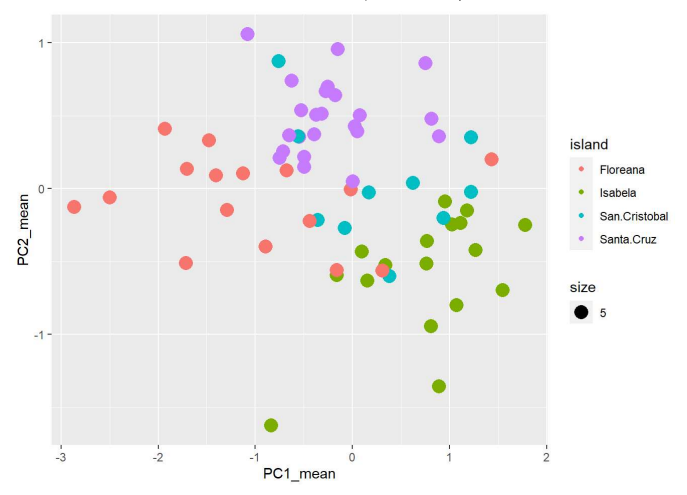


Left: Variance explained per PC. Rigth: Individual PCA as described above.

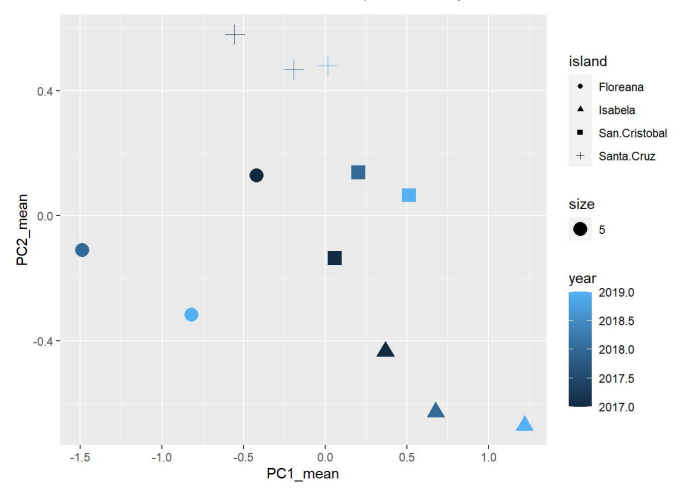
PCA means

Then I estimated the means of the PC scores obtained above and plot them in different ways to visualize the patters of variation.

Here I estimated the means per population and years for each island:



Here I estimated the means per year and sorted by island. -You can see there is a trend per year that goes in different directions-:

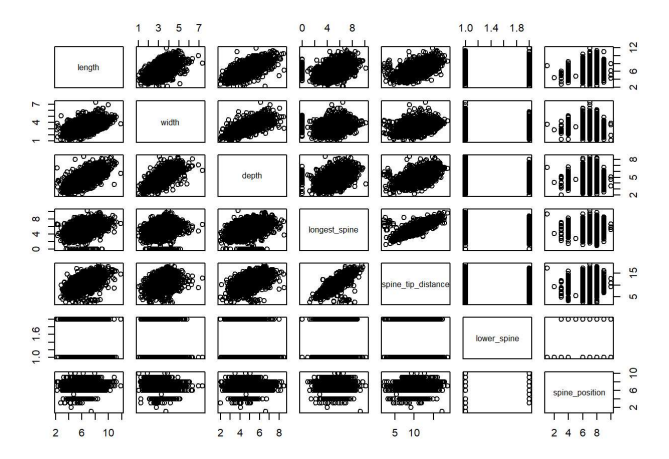


Variance Partition Analysis

I reviewed Sofia's code and use it as a reference to do the partition analysis. I ran a series of individual models per trait. I made sure that population was nested within islands and included island and year as random models. I edited the dataset to make all the other variables factors.

Descriptive data:

```
##
                         means
                                      sd
                                          min
## length
                      6.308864
                                1.359785 2.23 11.89
## width
                       3.22321 0.6389286 1.03
## depth
                      4.782531 0.8408234 2.01
                                               8.54
## longest_spine
                      5.372706
                                1.334417
                                            0 10.18
## spine tip distance 10.70068
                                2.451824 2.17
                                                 NA
```



Generalized linear model:

 $Imer(trait \sim 1 + (1|year) + (1|island) + (1|island:population), data = tribulusPC, REML = T)$

Where trait was each of the measured traits:

- length
- width
- depth
- · longest spine
- · spine tip distance

For lower spine I used a different model since is a binomial trait:

glmer(lower spine ~ 1 + (1|year) + (1|island) + (1|island:population), data = tribulusPC, na.action = na.omit, family = binomial)

I did not use spine position but I can include that as well I can do a similar analysis per island

For each trait I used those models I checked assumptions (as Sofia did) and looked at the variance explained by each trait and looked at the significance effect of each random factor: year, island and population within island.

Summary Results Length:

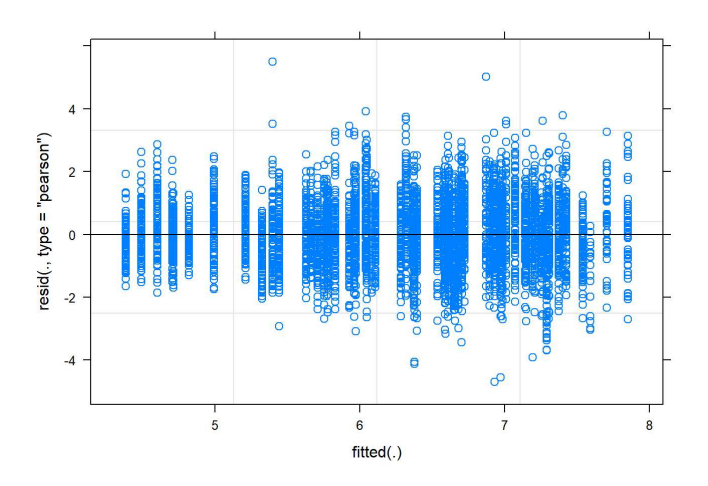
```
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: length \sim 1 + (1 \mid year) + (1 \mid island) + (1 \mid island:population)
      Data: tribulusPC
##
## REML criterion at convergence: 16457.2
##
## Scaled residuals:
##
      Min
               10 Median
                                3Q
                                       Max
  -4.4644 -0.6554 -0.0379 0.6406 5.2236
##
## Random effects:
   Groups
                                 Variance Std.Dev.
##
                      Name
##
   island:population (Intercept) 0.40583 0.6371
   island
                     (Intercept) 0.44624 0.6680
##
                      (Intercept) 0.02885 0.1698
##
   year
                                  1.10964 1.0534
##
   Residual
## Number of obs: 5552, groups: island:population, 25; island, 4; year, 3
##
## Fixed effects:
##
               Estimate Std. Error
                                       df t value Pr(>|t|)
                                           17.04 0.000172 ***
## (Intercept)
               6.3427
                            0.3723 3.4968
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Testing significance of each random effects:

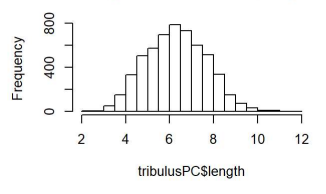
```
ranova(lm.length)
```

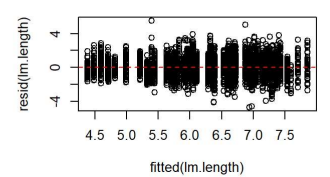
```
## ANOVA-like table for random-effects: Single term deletions
##
## Model:
## length \sim (1 | year) + (1 | island) + (1 | island:population)
##
                          npar logLik AIC
                                                LRT Df Pr(>Chisq)
## <none>
                             5 -8228.6 16467
## (1 | year)
                             4 -8274.1 16556
                                              90.92 1 < 2.2e-16 ***
## (1 | island)
                             4 -8233.0 16474
                                               8.84 1
                                                         0.002954 **
## (1 | island:population)
                            4 -8766.3 17541 1075.31 1 < 2.2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

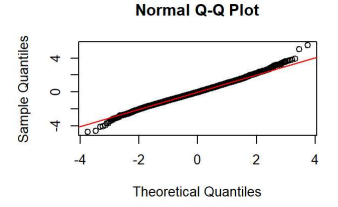
Checking assumptions *length*:

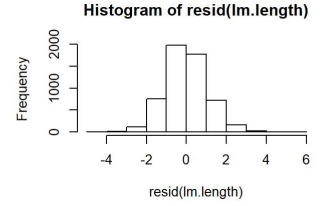












Summary Results Width

```
summary(lm.width)
```

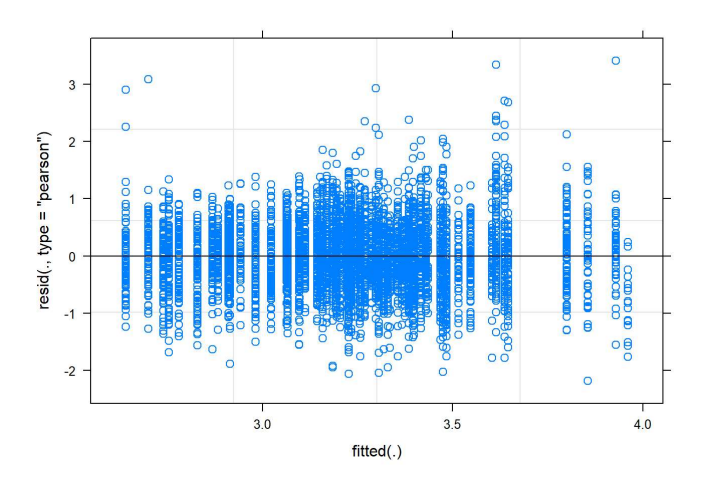
```
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: width ~ 1 + (1 | year) + (1 | island) + (1 | island:population)
      Data: tribulusPC
##
##
## REML criterion at convergence: 9820.7
##
## Scaled residuals:
##
      Min
               1Q Median
                               3Q
                                      Max
## -3.7678 -0.6146 -0.0292 0.6219 5.8806
##
## Random effects:
##
   Groups
                                 Variance Std.Dev.
                     Name
   island:population (Intercept) 0.085086 0.29169
                     (Intercept) 0.014726 0.12135
##
   year
                     (Intercept) 0.007487 0.08653
   Residual
                                 0.336490 0.58008
##
## Number of obs: 5552, groups: island:population, 25; island, 4; year, 3
##
## Fixed effects:
##
               Estimate Std. Error
                                       df t value Pr(>|t|)
## (Intercept) 3.24292 0.09881 4.73038
                                            32.82 9.17e-07 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Testing significance of each random effects:

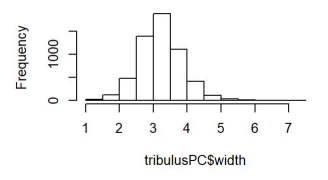
```
ranova(lm.width)
```

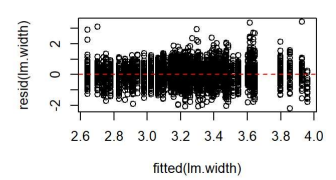
```
## ANOVA-like table for random-effects: Single term deletions
##
## Model:
## width \sim (1 | year) + (1 | island) + (1 | island:population)
##
                          npar logLik
                                           AIC
                                                  LRT Df Pr(>Chisq)
## <none>
                             5 -4910.3 9830.7
## (1 year)
                             4 -4947.8 9903.5 74.84 1
                                                            <2e-16 ***
## (1 | island)
                             4 -4910.8 9829.6 0.92 1
                                                            0.3367
## (1 | island:population) 4 -5236.4 10480.9 652.18 1
                                                           <2e-16 ***
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

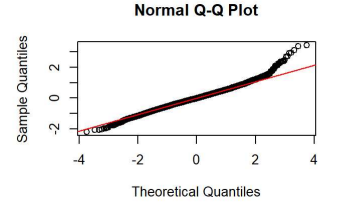
Checking assumptions width:

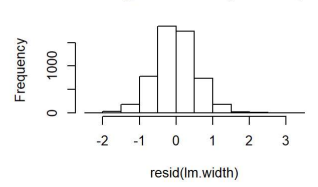


Histogram of tribulusPC\$width









Histogram of resid(Im.width)

Summary Results Depth

```
summary(lm.depth)
```

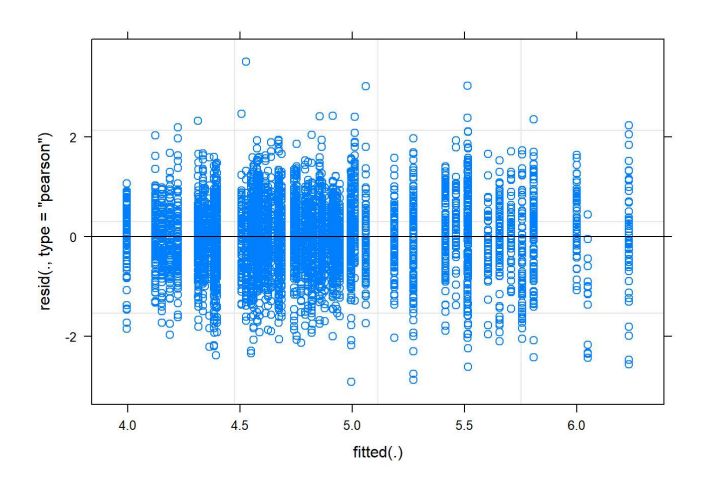
```
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: depth \sim 1 + (1 \mid year) + (1 \mid island) + (1 \mid island:population)
      Data: tribulusPC
##
##
## REML criterion at convergence: 12043.5
##
## Scaled residuals:
##
      Min
                10 Median
                                3Q
                                       Max
## -4.1191 -0.6260 0.0180 0.6526 4.9630
##
## Random effects:
##
   Groups
                                  Variance Std.Dev.
                      Name
   island:population (Intercept) 0.17119 0.4137
                      (Intercept) 0.14782 0.3845
##
   year
                      (Intercept) 0.01652 0.1285
   Residual
                                  0.50120 0.7080
##
## Number of obs: 5552, groups: island:population, 25; island, 4; year, 3
##
## Fixed effects:
##
               Estimate Std. Error
                                       df t value Pr(>|t|)
## (Intercept) 4.8287 0.2232 3.7845
                                           21.64 4.17e-05 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Testing significance of each random effects:

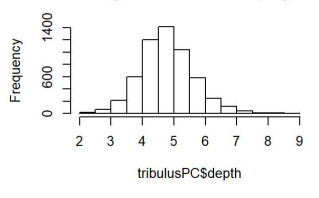
```
ranova(lm.depth)
```

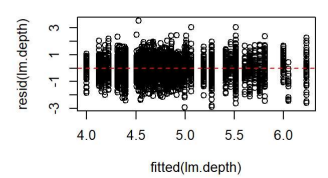
```
## ANOVA-like table for random-effects: Single term deletions
##
## Model:
## depth \sim (1 | year) + (1 | island) + (1 | island:population)
##
                          npar logLik AIC
                                               LRT Df Pr(>Chisq)
                             5 -6021.8 12054
## <none>
## (1 | year)
                             4 -6078.3 12164 113.02 1 < 2.2e-16 ***
## (1 | island)
                             4 -6025.3 12059
                                              7.06 1 0.007879 **
## (1 | island:population) 4 -6396.2 12800 748.78 1 < 2.2e-16 ***
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```

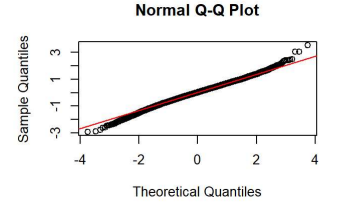
Checking assumptions *depth*:

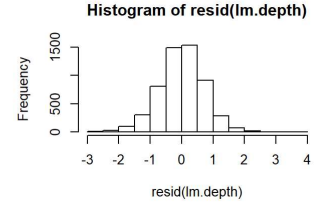


Histogram of tribulusPC\$depth









Summary Results Longest Spine

```
summary(lm.longspine)
```

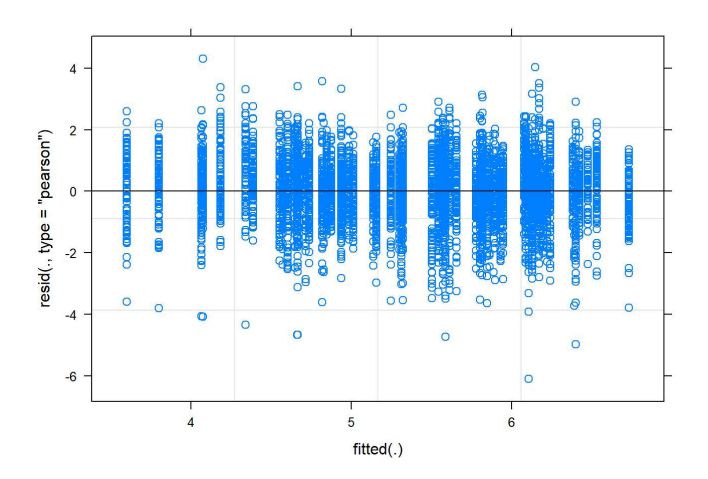
```
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula:
## longest_spine ~ 1 + (1 | year) + (1 | island) + (1 | island:population)
      Data: tribulusPC
##
##
## REML criterion at convergence: 16980.9
##
## Scaled residuals:
##
      Min
               1Q Median
                               3Q
                                      Max
## -5.5275 -0.5493 0.0718 0.6419 3.9052
##
## Random effects:
   Groups
##
                     Name
                                 Variance Std.Dev.
   island:population (Intercept) 0.30468 0.5520
                     (Intercept) 0.34562 0.5879
                     (Intercept) 0.08652 0.2941
##
   vear
##
   Residual
                                 1.22096 1.1050
## Number of obs: 5552, groups: island:population, 25; island, 4; year, 3
##
## Fixed effects:
##
               Estimate Std. Error
                                      df t value Pr(>|t|)
                           0.3584 4.5178
                                          14.99 5.03e-05 ***
## (Intercept) 5.3717
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Testing significance of each random effects:

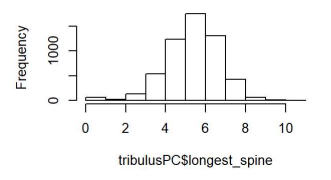
```
ranova(lm.longspine)
```

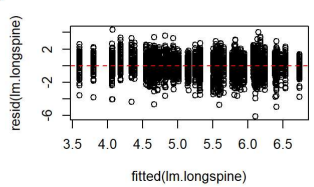
```
## ANOVA-like table for random-effects: Single term deletions
##
## Model:
## longest_spine ~ (1 | year) + (1 | island) + (1 | island:population)
##
                          npar logLik
                                       AIC
                                               LRT Df Pr(>Chisq)
## <none>
                             5 -8490.5 16991
## (1 | year)
                             4 -8604.5 17217 228.03 1 < 2.2e-16 ***
## (1 | island)
                             4 -8495.1 16998
                                              9.29 1
                                                      0.002309 **
## (1 | island:population)
                             4 -8843.1 17694 705.24 1 < 2.2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

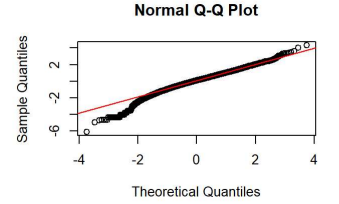
Checking assumptions longest spine:

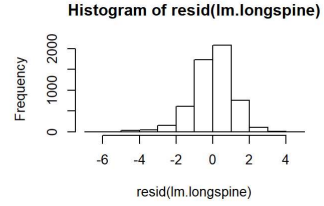


Histogram of tribulusPC\$longest_spine









Summary Results Spine Tip Distance

Spine tip distance model removed NAs only uses data from 2017-2018

```
summary(lm.tipdist)
```

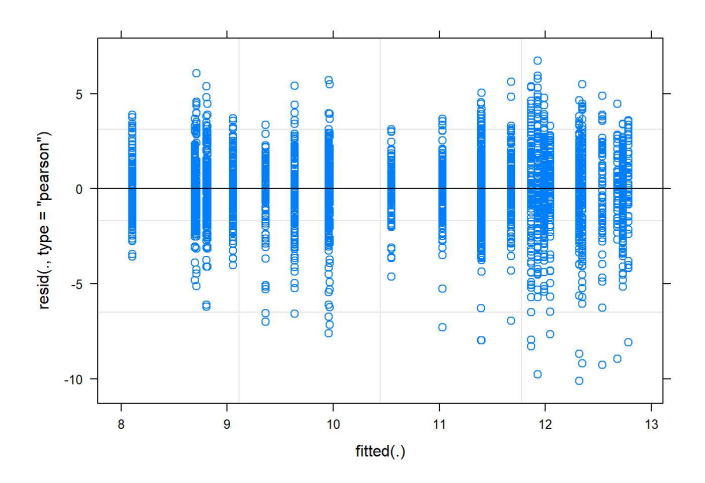
```
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula:
## spine_tip_distance ~ 1 + (1 | year) + (1 | island) + (1 | island:population)
      Data: tribulusPC
##
##
## REML criterion at convergence: 13918
##
## Scaled residuals:
##
      Min
                1Q Median
                                3Q
                                       Max
##
  -5.2632 -0.5685 0.0254 0.6195 3.5055
##
## Random effects:
##
   Groups
                      Name
                                  Variance Std.Dev.
   island:population (Intercept) 1.077
                                           1.038
##
##
   island
                      (Intercept) 1.690
                                           1.300
                      (Intercept) 0.000
                                           0.000
##
   year
   Residual
                                  3.697
                                           1.923
## Number of obs: 3334, groups: island:population, 25; island, 4; year, 3
##
## Fixed effects:
##
               Estimate Std. Error
                                        df t value Pr(>|t|)
## (Intercept) 10.6863 0.6854 3.0548
                                             15.59 0.000518 ***
## ---
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
## convergence code: 0
## boundary (singular) fit: see ?isSingular
```

Testing significance of each random effects:

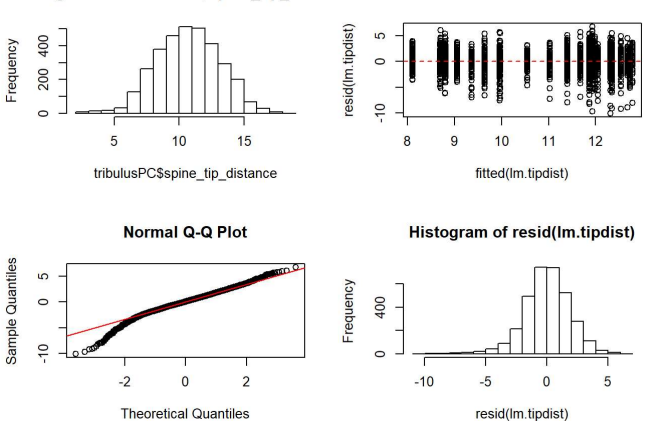
```
ranova(lm.tipdist)
```

```
## boundary (singular) fit: see ?isSingular
```

Checking assumptions spine tip distance:



Histogram of tribulusPC\$spine_tip_distan



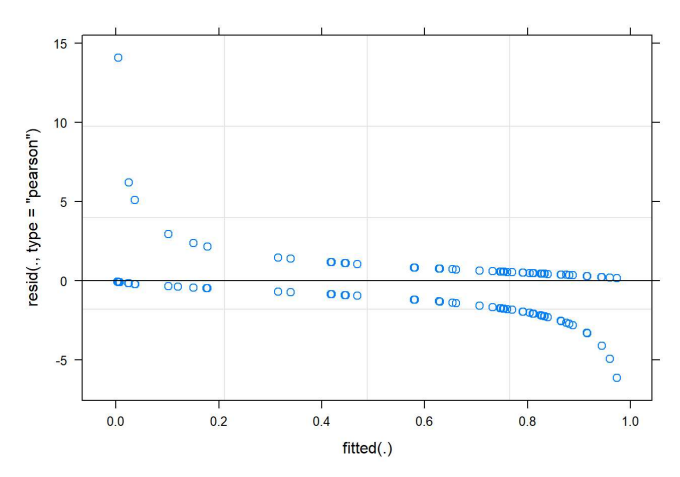
Summary Results Lower Spine

```
summary(lm.lowspines)
```

```
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
##
##
   Family: binomial ( logit )
## Formula: lower_spine \sim 1 + (1 \mid year) + (1 \mid island) + (1 \mid island:population)
      Data: tribulusPC
##
##
##
        AIC
                       logLik deviance df.resid
##
     4798.8
              4825.2 -2395.4
                                4790.8
                                            5548
##
## Scaled residuals:
                                3Q
##
       Min
                1Q Median
                                       Max
##
  -6.1378 -0.3373 0.3023 0.5127 14.1088
##
## Random effects:
   Groups
##
                      Name
                                  Variance Std.Dev.
   island:population (Intercept) 4.64714 2.1557
   island
                      (Intercept) 2.35354 1.5341
##
##
   year
                      (Intercept) 0.07092 0.2663
## Number of obs: 5552, groups: island:population, 25; island, 4; year, 3
##
## Fixed effects:
##
               Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.1810
                            0.8992 -0.201
```

Lower Spine plot

```
plot(lm.lowspines)
```



Testing significance of each random effects:

This was done manually given that lower spine is a binomial trait. Similar to Sofia's approach I created a model with all the random effects: year, island, island:population and then compared with other models removing each effect and test for significance.

Island:Population removed

summary(mred)

```
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
##
   Family: binomial (logit)
## Formula: lower_spine ~ 1 + (1 | year) + (1 | island)
      Data: tribulusPC
##
## Control: glmerControl(optimizer = "bobyqa", optCtrl = list(maxfun = 2e+05))
##
##
        AIC
                 BIC
                      logLik deviance df.resid
##
     6060.1
             6079.9 -3027.0
                               6054.1
                                           5549
##
## Scaled residuals:
##
      Min
                1Q Median
                                3Q
                                      Max
## -2.0760 -0.4049 0.4969 0.6871 2.8873
##
## Random effects:
   Groups Name
                      Variance Std.Dev.
##
   island (Intercept) 1.54519 1.2431
##
          (Intercept) 0.02984 0.1727
## Number of obs: 5552, groups: island, 4; year, 3
##
## Fixed effects:
               Estimate Std. Error z value Pr(>|z|)
## (Intercept)
               0.1356
                            0.6290
                                    0.216
                                              0.829
```

```
anova(mfull, mred)
```

Island removed

```
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
##
##
   Family: binomial (logit)
## Formula: lower_spine ~ 1 + (1 | year) + (1 | island:population)
      Data: tribulusPC
##
##
##
        AIC
                 BIC
                      logLik deviance df.resid
##
     4801.9
              4821.8 -2397.9 4795.9
                                           5549
##
## Scaled residuals:
##
       Min
                10 Median
                                3Q
                                       Max
  -6.1372 -0.3396 0.3029 0.5140 13.9500
##
## Random effects:
##
   Groups
                      Name
                                 Variance Std.Dev.
   island:population (Intercept) 7.48875 2.7366
##
   year
                      (Intercept) 0.06921 0.2631
##
## Number of obs: 5552, groups: island:population, 25; year, 3
##
## Fixed effects:
##
               Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.1649
                            0.5757 -0.286
```

```
## Data: tribulusPC
## Models:
## mred2: lower spine ~ 1 + (1 | year) + (1 | island:population)
## mfull: lower_spine \sim 1 + (1 | year) + (1 | island) + (1 | island:population)
                       BIC logLik deviance Chisq Df Pr(>Chisq)
        npar
                AIC
##
           3 4801.9 4821.8 -2397.9
## mred2
                                     4795.9
           4 4798.8 4825.2 -2395.4 4790.8 5.1273 1
## mfull
                                                        0.02355 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Population removed

```
summary(mred3)
```

```
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
##
##
   Family: binomial (logit)
## Formula: lower_spine ~ 1 + (1 | island) + (1 | island:population)
      Data: tribulusPC
##
##
##
        AIC
                BIC
                      logLik deviance df.resid
##
     4824.6
             4844.4 -2409.3 4818.6
                                          5549
##
## Scaled residuals:
##
      Min
               10 Median
                               3Q
                                      Max
## -5.4260 -0.3808 0.3277 0.5179 11.7428
##
## Random effects:
## Groups
                     Name
                                 Variance Std.Dev.
  island:population (Intercept) 4.625
                                          2.151
##
   island
                     (Intercept) 2.311
                                          1.520
##
## Number of obs: 5552, groups: island:population, 25; island, 4
##
## Fixed effects:
##
               Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.1790 0.8844 -0.202
```

```
anova(mfull, mred3)
```