HW8

Chapters 11 and 12

TA Solutions

The code below just loads some packages and makes it so that enough digits are printed that you won't get confused by rounding errors.

```
library(dplyr) # functions like summarize
library(ggplot2) # for making plots
library(gridExtra)
library(GGally)
library(readr)
library(car)
options("pillar.sigfig" = 10) # print 10 significant digits in summarize output
```

Problem 1: Galapagos (Adapted from Sleuth3 12.20)

Quote from book:

The data [read in below] come from a 1973 study. (Data from M. P. Johnson and P. H. Raven, "Species Number and Endemism: The Galapagos Archipelago Revisited," *Science* 179 (1973): 893-5.) The number of species on an island is known to be related to the island's area. Of interest is what other variables are also related to the number of species, after island area is accounted for. The data set includes the following variables:

- Island: a character vector indicating the island
- Total: total number of observed species
- Native: number of native species
- Area: area (km²)
- Elev: elevation (m)
- DistNear: distance from nearest island (km)
- DistSc: distance from Santa Cruz (km)
- AreaNear: area of nearest island (km²)

In this analysis, our response variable is Native, the number of native species. You will use Area, Elev, DistNear, DistSc, and AreaNear as possible explanatory variables.

```
galapagos <- read_csv("http://www.evanlray.com/data/sleuth3/ex1220_galapagos.csv")</pre>
```

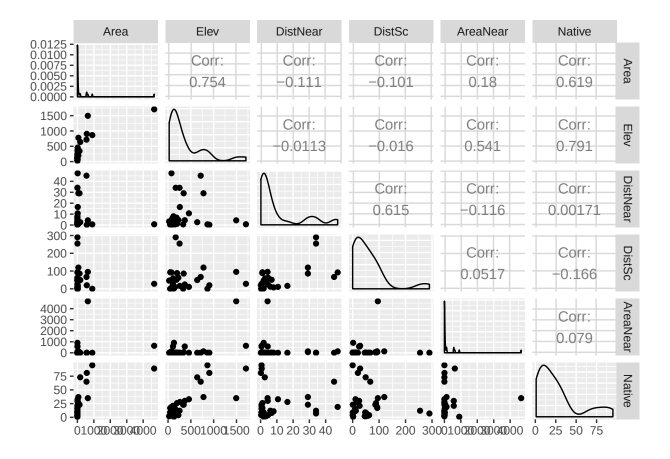
```
## Parsed with column specification:
## cols(
##
     Island = col_character(),
##
     Total = col_double(),
     Native = col_double(),
##
     Area = col double(),
##
    Elev = col_double(),
##
##
    DistNear = col_double(),
    DistSc = col_double(),
##
     AreaNear = col_double()
##
## )
```

head(galapagos)

```
## # A tibble: 6 x 8
##
     Island
                   Total Native Area Elev DistNear DistSc AreaNear
                   <dbl>
                                                 <dbl>
                                                        <dbl>
##
     <chr>>
                          <dbl> <dbl> <dbl>
                                                                  <dbl>
                                                           0.6
                              23 25.09
                                                   0.6
## 1 Baltra
                      58
                                         332
                                                                   1.84
## 2 Bartolome
                      31
                              21
                                  1.24
                                         109
                                                   0.6
                                                          26.3
                                                                 572.33
                       3
                                                                   0.78
## 3 Caldwell
                               3
                                  0.21
                                         114
                                                   2.8
                                                          58.7
## 4 Champion
                      25
                               9
                                  0.1
                                          46
                                                   1.9
                                                          47.4
                                                                   0.18
                                                                 903.82
## 5 Coamano
                       2
                                  1.05
                                         130
                                                   1.9
                               1
                                                           1.9
## 6 Daphne Major
                      18
                              11
                                  0.34
                                         119
                                                   8
                                                           8
                                                                   1.84
```

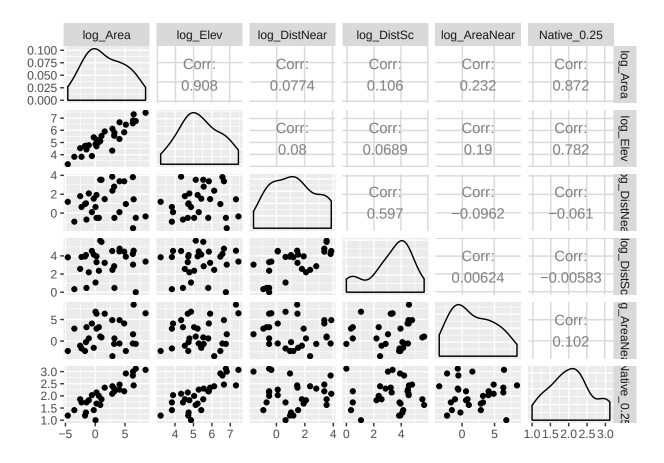
(a) Make a pairs plot of the data including only the variables you will use in your analysis (with the response variable last).

ggpairs(galapagos %>% select(Area, Elev, DistNear, DistSc, AreaNear, Native))

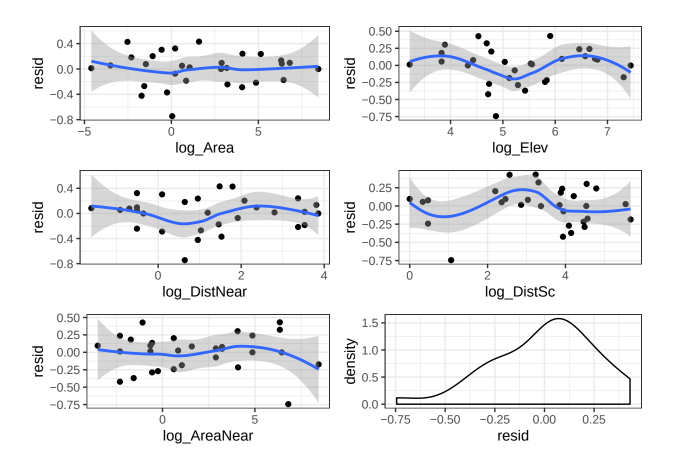


(b) Identify a set of transformations for all variables in the model so that the regression conditions appear to be fairly well satisfied. In doing this, consider pairs plots of the transformed data and plots of residuals vs explanatory variables in a regression model including all transformed explanatory variables. (You should have both of these types of plots.) After this step, you should feel fairly confident that any models you fit will either have approximately linear relationships among transformed variables, or know how you will handle non-linearity by adding polynomial terms in the model. You should also feel confident that the variance of residuals is fairly constant across values of explanatory variables. Note that DistSc includes some 0 values. A common trick in cases like this is to add 1 to the observed values of that variable before doing transformations (this means that things like a log transformation are an option).

```
galapagos_transformed <- galapagos %>%
transmute(
  log_Area = log(Area),
  log_Elev = log(Elev),
  log_DistNear = log(DistNear),
  log_DistSc = log(DistSc + 1),
  log_AreaNear = log(AreaNear),
  Native_0.25 = Native^0.25
)
ggpairs(galapagos_transformed)
```

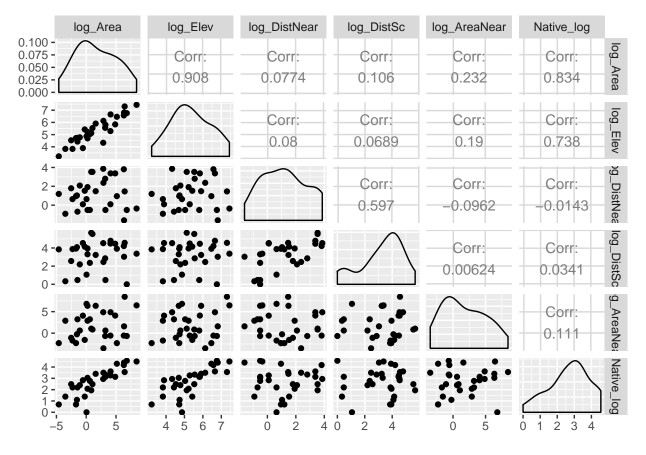


```
lm_full <- lm(Native_0.25 ~ log_Area + log_Elev + log_DistNear + log_DistSc + log_AreaNear,</pre>
  data = galapagos_transformed)
galapagos_transformed <- galapagos_transformed %>%
   resid = residuals(lm_full)
p1 <- ggplot(data = galapagos_transformed, mapping = aes(x = log_Area, y = resid)) +
  geom point() +
  geom smooth() +
  theme bw()
p2 <- ggplot(data = galapagos_transformed, mapping = aes(x = log_Elev, y = resid)) +
  geom_point() +
  geom smooth() +
 theme_bw()
p3 <- ggplot(data = galapagos_transformed, mapping = aes(x = log_DistNear, y = resid)) +
  geom_point() +
  geom_smooth() +
  theme_bw()
p4 <- ggplot(data = galapagos_transformed, mapping = aes(x = log_DistSc, y = resid)) +
  geom_point() +
  geom_smooth() +
 theme_bw()
p5 <- ggplot(data = galapagos_transformed, mapping = aes(x = log_AreaNear, y = resid)) +
  geom_point() +
  geom smooth() +
 theme bw()
p6 <- ggplot(data = galapagos_transformed, mapping = aes(x = resid)) +</pre>
  geom_density() +
  theme_bw()
grid.arrange(p1, p2, p3, p4, p5, p6)
## `geom_smooth()` using method = 'loess' and formula 'y ~ x'
## geom_smooth() using method = 'loess' and formula 'y ~ x'
## `geom_smooth()` using method = 'loess' and formula 'y ~ x'
## `geom_smooth()` using method = 'loess' and formula 'y ~ x'
## `geom_smooth()` using method = 'loess' and formula 'y ~ x'
```



Grader note: a log transformation for Native is also OK. The fourth root transformation looks slightly better, but log also works, and has interpretation advantages. The output with log(Native) as the response is included as the second output for each part of the homework for reference.

```
galapagos_transformed2 <- galapagos %>%
    transmute(
    log_Area = log(Area),
    log_Elev = log(Elev),
    log_DistNear = log(DistNear),
    log_DistSc = log(DistSc + 1),
    log_AreaNear = log(AreaNear),
    Native_log = log(Native)
    )
ggpairs(galapagos_transformed2)
```



```
lm_full2 <- lm(Native_log ~ log_Area + log_Elev + log_DistNear + log_DistSc + log_AreaNear,</pre>
  data = galapagos_transformed2)
galapagos_transformed2 <- galapagos_transformed2 %>%
  mutate(
    resid = residuals(lm_full2)
p1 <- ggplot(data = galapagos_transformed2, mapping = aes(x = log_Area, y = resid)) +
  geom_point() +
  geom_smooth() +
  theme_bw()
p2 <- ggplot(data = galapagos_transformed2, mapping = aes(x = log_Elev, y = resid)) +
  geom_point() +
  geom_smooth() +
  theme_bw()
p3 <- ggplot(data = galapagos_transformed2, mapping = aes(x = log_DistNear, y = resid)) +
  geom_point() +
  geom_smooth() +
  theme bw()
p4 <- ggplot(data = galapagos_transformed2, mapping = aes(x = log_DistSc, y = resid)) +
  geom_point() +
  geom_smooth() +
  theme bw()
p5 <- ggplot(data = galapagos transformed2, mapping = aes(x = log AreaNear, y = resid)) +
  geom_point() +
  geom_smooth() +
theme_bw()
```

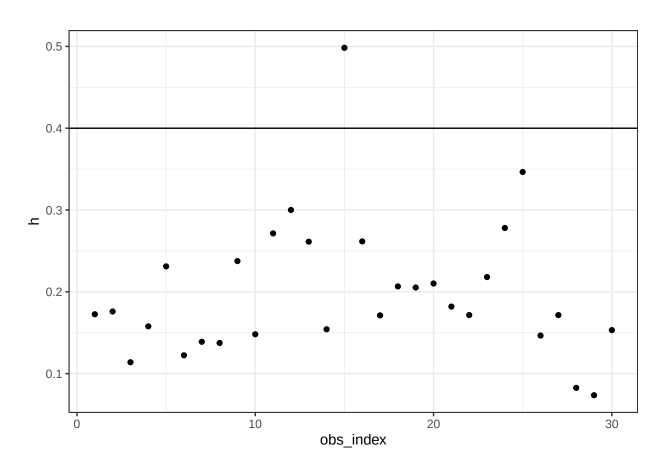
```
p6 <- ggplot(data = galapagos_transformed2, mapping = aes(x = resid)) +</pre>
  geom_density() +
  theme_bw()
grid.arrange(p1, p2, p3, p4, p5, p6)
## `geom_smooth()` using method = 'loess' and formula 'y ~ x'
## `geom_smooth()` using method = 'loess' and formula 'y ~ x'
## geom_smooth() using method = 'loess' and formula 'y ~ x'
## geom_smooth() using method = 'loess' and formula 'y ~ x'
## `geom_smooth()` using method = 'loess' and formula 'y ~ x'
                                                 resid
resid
       -5
                                                                       log_Elev
                      log_Area
resid
                                                 resid
                                                                     2
                    log_DistNear
                                                                      log_DistSc
                                                    0.6
                                                 density
0.4
0.2
 resid
                                                    0.0
                    log_AreaNear
                                                                         resid
```

(c) Check for influential observations, outliers or high leverage observations.

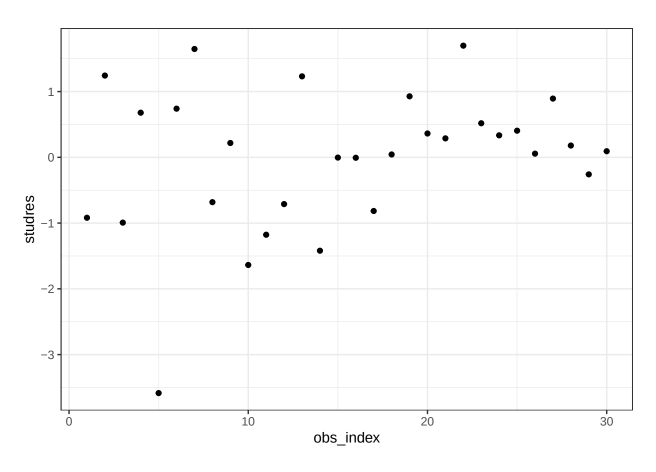
With Native $^{0.25}$ transformation:

```
galapagos_transformed <- galapagos_transformed %>%
mutate(
   obs_index = row_number(),
   h = hatvalues(lm_full),
   studres = rstudent(lm_full),
   D = cooks.distance(lm_full)
)
ggplot(data = galapagos_transformed, mapping = aes(x = obs_index, y = h)) +
   geom_hline(yintercept = 2 * 6 / nrow(galapagos_transformed))+
```

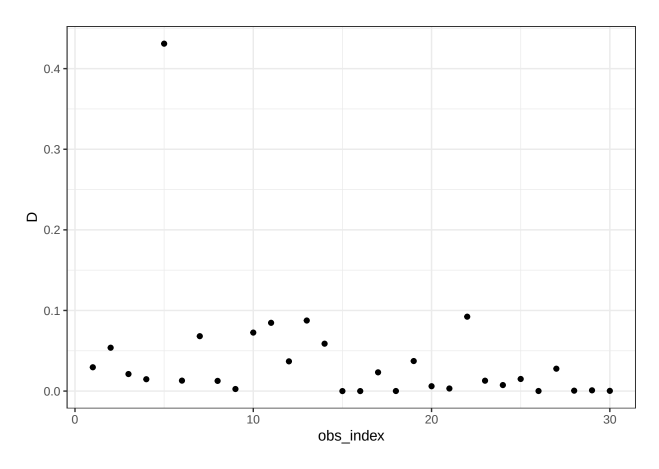
```
geom_point() +
theme_bw()
```



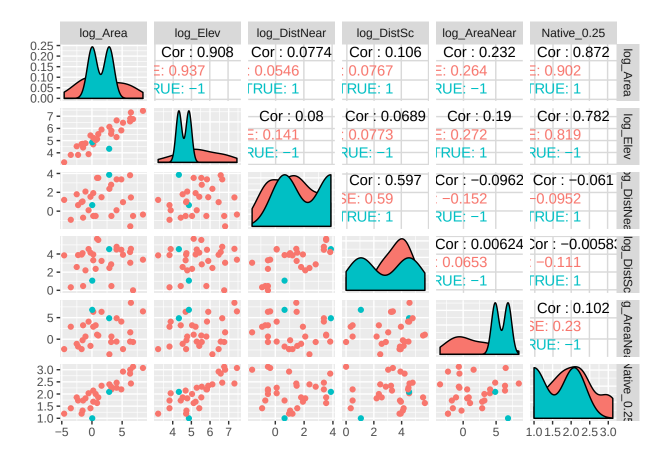
```
ggplot(data = galapagos_transformed, mapping = aes(x = obs_index, y = studres)) +
  geom_point() +
  theme_bw()
```



```
ggplot(data = galapagos_transformed, mapping = aes(x = obs_index, y = D)) +
geom_point() +
theme_bw()
```

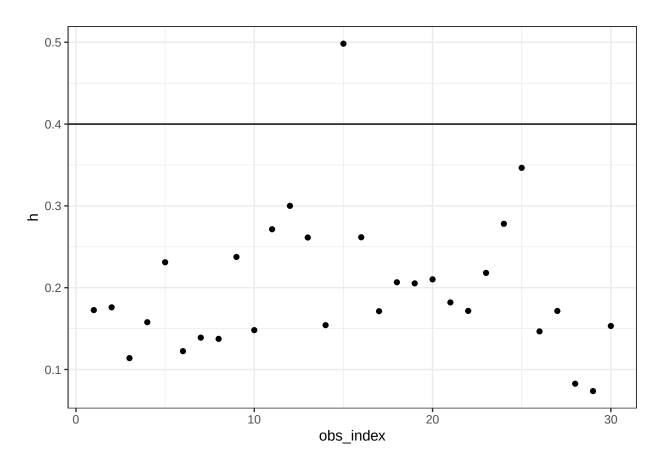


```
galapagos_transformed <- galapagos_transformed %>%
  mutate(suspicious = obs_index %in% c(5, 15))
ggpairs(galapagos_transformed, mapping = aes(color = suspicious), columns = 1:6)
```

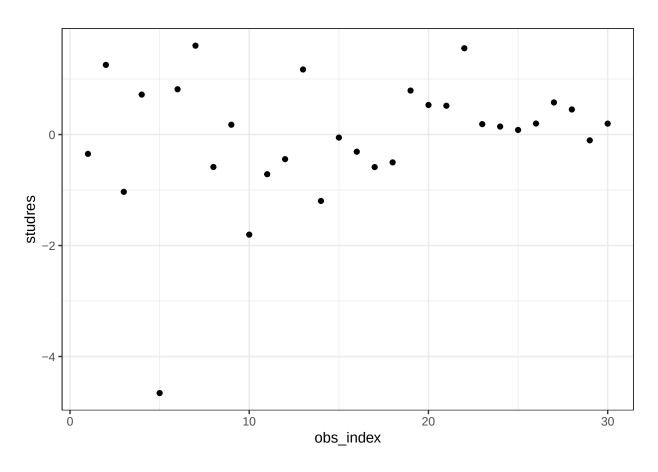


With log(Native) transformation:

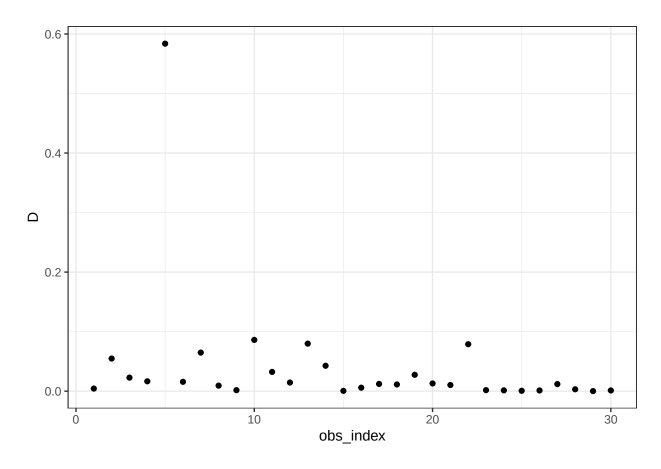
```
galapagos_transformed2 <- galapagos_transformed2 %>%
  mutate(
    obs_index = row_number(),
    h = hatvalues(lm_full2),
    studres = rstudent(lm_full2),
    D = cooks.distance(lm_full2)
)
ggplot(data = galapagos_transformed2, mapping = aes(x = obs_index, y = h)) +
    geom_hline(yintercept = 2 * 6 / nrow(galapagos_transformed2))+
    geom_point() +
    theme_bw()
```



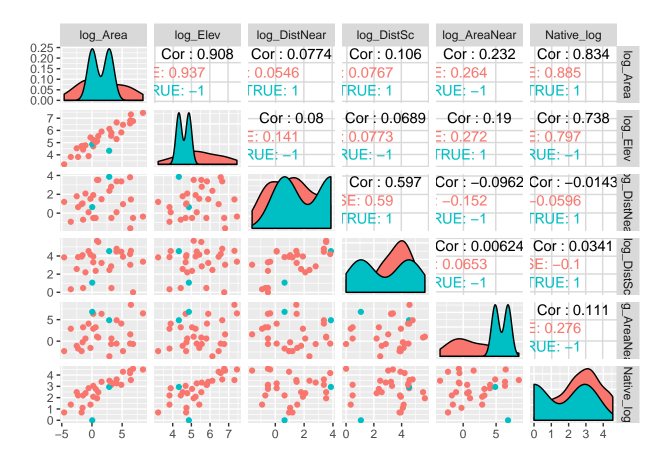
```
ggplot(data = galapagos_transformed2, mapping = aes(x = obs_index, y = studres)) +
  geom_point() +
  theme_bw()
```



```
ggplot(data = galapagos_transformed2, mapping = aes(x = obs_index, y = D)) +
geom_point() +
theme_bw()
```



```
galapagos_transformed2 <- galapagos_transformed2 %>%
  mutate(suspicious = obs_index %in% c(5, 15))
ggpairs(galapagos_transformed2, mapping = aes(color = suspicious), columns = 1:6)
```



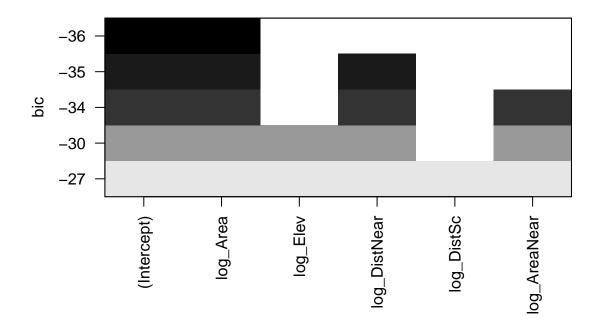
(d) Use all subsets regression to identify a set of models with similar ability to model these data well, based on your transformed variables. If necessary, perform this step both with and without the outliers or influential observations included.

With Native $^{0.25}$ transformation:

```
library(leaps)
```

```
## Warning: package 'leaps' was built under R version 3.6.3
```

```
candidate_models <- regsubsets(Native_0.25 ~ log_Area + log_Elev + log_DistNear +
    log_DistSc + log_AreaNear, data = galapagos_transformed)
plot(candidate_models)</pre>
```

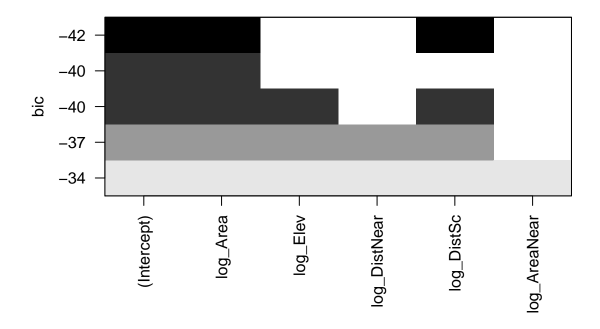


summary(candidate_models)

```
## Subset selection object
## Call: regsubsets.formula(Native_0.25 ~ log_Area + log_Elev + log_DistNear +
##
       log_DistSc + log_AreaNear, data = galapagos_transformed)
## 5 Variables (and intercept)
##
                Forced in Forced out
                    FALSE
                                FALSE
## log_Area
                                FALSE
## log_Elev
                    FALSE
## log_DistNear
                    FALSE
                                FALSE
## log_DistSc
                    FALSE
                                FALSE
## log_AreaNear
                    FALSE
                                FALSE
## 1 subsets of each size up to 5
## Selection Algorithm: exhaustive
##
            log_Area log_Elev log_DistNear log_DistSc log_AreaNear
## 1
      (1) "*"
                     11 11
                                            11 11
     (1)"*"
                      11 11
                               "*"
                                            11 11
## 2
     (1)"*"
                      11 11
                               "*"
                                            11 11
                                                        "*"
     (1)"*"
                      "*"
                               "*"
                                                        "*"
## 4
                      "*"
                                            "*"
     (1)"*"
                               "*"
                                                        "*"
```

summary(candidate_models)\$bic

```
## [1] -36.15035 -34.91195 -33.51806 -30.22219 -26.89620
```



summary(candidate_models2)

```
## Subset selection object
## Call: regsubsets.formula(Native_0.25 ~ log_Area + log_Elev + log_DistNear +
       log_DistSc + log_AreaNear, data = galapagos_transformed %>%
##
       filter(!suspicious))
## 5 Variables (and intercept)
##
                Forced in Forced out
## log_Area
                    FALSE
                                FALSE
## log_Elev
                    FALSE
                                FALSE
## log_DistNear
                    FALSE
                                FALSE
## log_DistSc
                    FALSE
                                FALSE
## log_AreaNear
                    FALSE
                                FALSE
## 1 subsets of each size up to 5
## Selection Algorithm: exhaustive
##
            log_Area log_Elev log_DistNear log_DistSc log_AreaNear
## 1 ( 1 ) "*"
                     11 11
                               11 11
     (1)"*"
                                                        11 11
## 2
                                            "*"
                     "*"
                               11 11
                                            "*"
## 3 (1) "*"
## 4 ( 1 ) "*"
                     "*"
                               "*"
                                            "*"
                                                        11 11
## 5 (1) "*"
                     "*"
                               "*"
                                            "*"
                                                        "*"
```

summary(candidate_models2)\$bic

```
## [1] -40.33967 -42.38225 -40.04481 -36.84316 -33.51199
```

When all observations are included, the following three models have roughly similar performance:

Model 1: log Area, log DistNear, and log AreaNear as explanatory variables

Model 2: log Area and log DistNear as explanatory variables

Model 3: log Area as the only explanatory variable

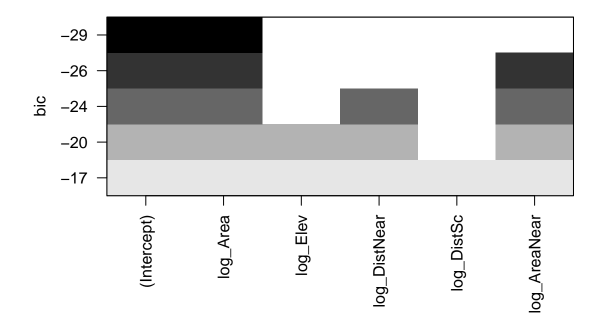
When one outlier and one high leverage observation are omitted, the following three models have roughly similar performance:

Model 1: log Area, log DistSc, and log Elev as explanatory variables

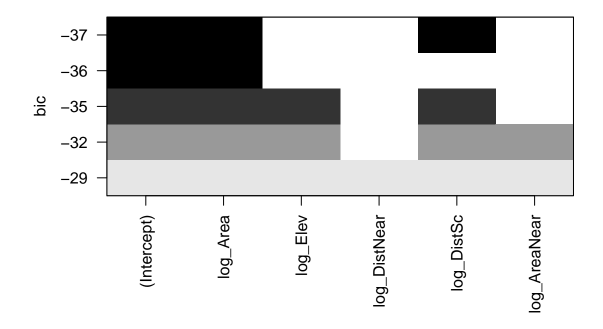
Model 2: log Area and log DistSc as explanatory variables

Model 3: log Area as the only explanatory variable

With log(Native) transformation:



```
summary(candidate_models2_1)
## Subset selection object
## Call: regsubsets.formula(Native_log ~ log_Area + log_Elev + log_DistNear +
       log_DistSc + log_AreaNear, data = galapagos_transformed2)
## 5 Variables (and intercept)
##
                Forced in Forced out
## log_Area
                    FALSE
                              FALSE
## log_Elev
                    FALSE
                               FALSE
## log_DistNear
                               FALSE
                   FALSE
## log_DistSc
                    FALSE
                               FALSE
## log_AreaNear
                               FALSE
                   FALSE
## 1 subsets of each size up to 5
## Selection Algorithm: exhaustive
           log_Area log_Elev log_DistNear log_DistSc log_AreaNear
##
                    11 11
                              11 11
                                    11 11
                                                     11 11
## 1 ( 1 ) "*"
                     11 11
                              11 11
                                           11 11
                                                      "*"
## 2 (1)"*"
                     11 11
                              "*"
                                          11 11
                                                      "*"
## 3 (1) "*"
                                           11 11
## 4 (1) "*"
                     "*"
                              "*"
                                                      "*"
## 5 (1) "*"
                     "*"
                              "*"
                                           "*"
                                                      "*"
summary(candidate_models2_1)$bic
## [1] -28.79992 -26.11391 -23.53182 -20.38876 -16.99462
candidate_models2_2 <- regsubsets(Native_log ~ log_Area + log_Elev + log_DistNear +</pre>
    log_DistSc + log_AreaNear, data = galapagos_transformed2 %% filter(!suspicious))
plot(candidate_models2_2)
```



summary(candidate_models2_2)

```
## Subset selection object
## Call: regsubsets.formula(Native_log ~ log_Area + log_Elev + log_DistNear +
##
       log_DistSc + log_AreaNear, data = galapagos_transformed2 %>%
       filter(!suspicious))
## 5 Variables (and intercept)
##
                Forced in Forced out
                    FALSE
                               FALSE
## log_Area
## log_Elev
                    FALSE
                               FALSE
## log_DistNear
                    FALSE
                               FALSE
## log_DistSc
                    FALSE
                               FALSE
## log_AreaNear
                    FALSE
                               FALSE
## 1 subsets of each size up to 5
## Selection Algorithm: exhaustive
##
            log_Area log_Elev log_DistNear log_DistSc log_AreaNear
## 1 ( 1 ) "*"
                     11 11
                               11 11
## 2 (1) "*"
                                            "*"
                                                       11 11
                     "*"
                               11 11
                                            "*"
     (1)"*"
## 4 ( 1 ) "*"
                     "*"
                               11 11
                                            "*"
                     "*"
                               "*"
                                            "*"
## 5 (1)"*"
                                                       "*"
```

summary(candidate_models2_2)\$bic

```
## [1] -36.14739 -36.74069 -34.61666 -31.81287 -28.67545
```

When all observations are included, the following two models have roughly similar performance:

```
Model 1: log Area as the only explanatory variables (BIC=-28.8)
```

```
Model 2: log Area and log AreaNear as explanatory variables (BIC=-26.1)
```

When one outlier and one high leverage observation are omitted, the following three models have roughly similar performance:

```
Model 1: log Area, log DistSc, and log Elev as explanatory variables (BIC=-34.6)
```

Model 2: log Area and log DistSc as explanatory variables (BIC=-36.7)

```
Model 3: log Area as the only explanatory variable (BIC=-36.1)
```

Note: Model 2 and Model 3 are the same models identified for the fourth root transformation for Native, both with and without the outlier and high leverage point. If someone makes a log transformation, you don't need to take off points, but make a note that they should investigate a fourth root transformation - it is more stabilizing in this case for outliers and leverage points (removing them from the analysis doesn't have the same impact that it does for the log transformation) They may not really see that until part (d).

(e) Obtain the model fits for all models you identified in part (d) as explaining the data about as well as each other, and print the model summaries.

With Native^{0.25} transformation

```
fit1 <- lm(Native_0.25 ~ log_Area + log_DistNear + log_AreaNear, data = galapagos_transformed)
summary(fit1)</pre>
```

```
##
## Call:
## lm(formula = Native_0.25 ~ log_Area + log_DistNear + log_AreaNear,
##
       data = galapagos_transformed)
##
## Residuals:
##
       Min
                  1Q
                      Median
                                    3Q
                                            Max
## -0.72395 -0.19768 0.05207 0.17589
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                            0.07354
                 1.89972
                                    25.834 < 2e-16 ***
## log_Area
                 0.15514
                            0.01572
                                      9.872 2.77e-10 ***
## log_DistNear -0.05232
                            0.03292
                                    -1.589
                                               0.124
## log AreaNear -0.02200
                            0.01640
                                    -1.341
                                               0.191
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.279 on 26 degrees of freedom
## Multiple R-squared: 0.7921, Adjusted R-squared: 0.7681
## F-statistic: 33.02 on 3 and 26 DF, p-value: 5.091e-09
fit2 <- lm(Native_0.25 ~ log_Area + log_DistNear, data = galapagos_transformed)
summary(fit2)
```

```
##
## Call:
## lm(formula = Native_0.25 ~ log_Area + log_DistNear, data = galapagos_transformed)
## Residuals:
##
       \mathtt{Min}
                 1Q Median
                                   3Q
## -0.84087 -0.19373 0.04474 0.21246 0.46082
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                1.86380
                           0.06949 26.820 < 2e-16 ***
                           0.01547
                                    9.696 2.74e-10 ***
## log_Area
                0.15005
## log_DistNear -0.04713
                           0.03317 - 1.421
                                              0.167
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.2831 on 27 degrees of freedom
## Multiple R-squared: 0.7777, Adjusted R-squared: 0.7613
## F-statistic: 47.24 on 2 and 27 DF, p-value: 1.524e-09
fit3 <- lm(Native_0.25 ~ log_Area, data = galapagos_transformed)
summary(fit3)
##
## Call:
## lm(formula = Native_0.25 ~ log_Area, data = galapagos_transformed)
## Residuals:
       Min
                 1Q
                     Median
## -0.81591 -0.15749 0.05753 0.23017 0.42722
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
##
                          0.05869 30.818 < 2e-16 ***
## (Intercept) 1.80867
## log Area
               0.14834
                          0.01571
                                   9.445 3.34e-10 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.2882 on 28 degrees of freedom
## Multiple R-squared: 0.7611, Adjusted R-squared: 0.7526
## F-statistic: 89.21 on 1 and 28 DF, p-value: 3.342e-10
fit1a <- lm(Native_0.25 ~ log_Area + log_DistSc + log_AreaNear,
            data = galapagos_transformed %>% filter(!suspicious))
summary(fit1a)
##
## Call:
## lm(formula = Native_0.25 ~ log_Area + log_DistSc + log_AreaNear,
       data = galapagos_transformed %>% filter(!suspicious))
##
## Residuals:
##
       Min
                 1Q
                                   3Q
                     Median
                                           Max
```

```
## -0.36159 -0.15993 0.00683 0.14532 0.36227
##
## Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                2.068e+00 1.095e-01 18.879 6.61e-16 ***
               1.471e-01 1.337e-02 11.004 7.36e-11 ***
## log Area
## log_DistSc -6.729e-02 2.989e-02 -2.251 0.0338 *
## log_AreaNear -1.513e-05 1.477e-02 -0.001
                                              0.9992
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.2347 on 24 degrees of freedom
## Multiple R-squared: 0.846, Adjusted R-squared: 0.8267
## F-statistic: 43.94 on 3 and 24 DF, p-value: 6.651e-10
fit2a <- lm(Native_0.25 ~ log_Area + log_DistSc, data = galapagos_transformed %>% filter(!suspicious))
summary(fit2a)
##
## lm(formula = Native_0.25 ~ log_Area + log_DistSc, data = galapagos_transformed %>%
      filter(!suspicious))
##
##
## Residuals:
##
       Min
                 1Q
                    Median
                                   3Q
## -0.36155 -0.15995 0.00682 0.14536 0.36234
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 2.06787
                          0.10681
                                  19.36 < 2e-16 ***
## log_Area
               0.14711
                          0.01265
                                   11.63 1.39e-11 ***
## log_DistSc -0.06729
                          0.02926
                                   -2.30 0.0301 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.23 on 25 degrees of freedom
## Multiple R-squared: 0.846, Adjusted R-squared: 0.8337
## F-statistic: 68.66 on 2 and 25 DF, p-value: 6.994e-11
fit3a <- lm(Native_0.25 ~ log_Area, data = galapagos_transformed %>% filter(!suspicious))
summary(fit3a)
##
## lm(formula = Native_0.25 ~ log_Area, data = galapagos_transformed %>%
##
      filter(!suspicious))
##
## Residuals:
##
       Min
                 1Q Median
                                   3Q
## -0.41110 -0.18423 0.04419 0.22051 0.37839
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
```

```
## (Intercept) 1.84874
                         0.05213 35.47 < 2e-16 ***
             0.14488
                          0.01361 10.64 5.67e-11 ***
## log_Area
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.2482 on 26 degrees of freedom
## Multiple R-squared: 0.8134, Adjusted R-squared: 0.8062
## F-statistic: 113.3 on 1 and 26 DF, p-value: 5.674e-11
With log(Native transformation):
fit2_1 <- lm(Native_log ~ log_Area, data = galapagos_transformed2)</pre>
summary(fit2_1)
##
## Call:
## lm(formula = Native_log ~ log_Area, data = galapagos_transformed2)
## Residuals:
               1Q Median
                               3Q
## -2.2471 -0.2140 0.1364 0.4145 0.9884
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 2.23288 0.13646 16.363 7.27e-16 ***
## log_Area
               0.29155
                          0.03652 7.984 1.08e-08 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.67 on 28 degrees of freedom
## Multiple R-squared: 0.6948, Adjusted R-squared: 0.6839
## F-statistic: 63.74 on 1 and 28 DF, p-value: 1.076e-08
fit2_2 <- lm(Native_log ~ log_Area + log_AreaNear, data = galapagos_transformed2)
summary(fit2_2)
##
## Call:
## lm(formula = Native_log ~ log_Area + log_AreaNear, data = galapagos_transformed2)
##
## Residuals:
                 1Q
                     Median
                                   3Q
## -2.07427 -0.27298 0.04037 0.41418 0.92891
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
                          0.14733 15.448 6.30e-15 ***
## (Intercept)
                2.27597
## log_Area
                0.29863
                           0.03778 7.904 1.69e-08 ***
```

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

```
##
## Residual standard error: 0.6743 on 27 degrees of freedom
## Multiple R-squared: 0.702, Adjusted R-squared: 0.6799
## F-statistic: 31.8 on 2 and 27 DF, p-value: 7.987e-08
fit2_1a <- lm(Native_log ~ log_Area + log_DistSc + log_Elev,</pre>
           data = galapagos_transformed2 %>% filter(!suspicious))
summary(fit2_1a)
##
## Call:
## lm(formula = Native_log ~ log_Area + log_DistSc + log_Elev, data = galapagos_transformed2 %>%
      filter(!suspicious))
##
## Residuals:
##
       Min
                 1Q
                      Median
                                   3Q
                                           Max
## -0.99171 -0.33379 -0.05635 0.35918 0.87608
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 4.01055
                          1.25788
                                   3.188 0.003950 **
## log Area
               0.36111
                          0.07885
                                   4.580 0.000121 ***
## log_DistSc -0.12303
                          0.06389 -1.926 0.066087 .
## log Elev
              -0.25903
                          0.25180 -1.029 0.313873
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.5022 on 24 degrees of freedom
## Multiple R-squared: 0.8196, Adjusted R-squared: 0.797
## F-statistic: 36.34 on 3 and 24 DF, p-value: 4.383e-09
fit2_2a <- lm(Native_log ~ log_Area + log_DistSc, data = galapagos_transformed2 %>% filter(!suspicious)
summary(fit2_2a)
##
## Call:
## lm(formula = Native_log ~ log_Area + log_DistSc, data = galapagos_transformed2 %>%
      filter(!suspicious))
##
## Residuals:
##
       Min
                 1Q Median
                                   3Q
## -1.06862 -0.31592 -0.01162 0.25476 0.81829
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 2.73900
                          0.23350
                                    11.73 1.17e-11 ***
               0.28514
                          0.02765
                                    10.31 1.71e-10 ***
## log_Area
                          0.06396
                                    -1.94 0.0638 .
## log_DistSc -0.12406
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.5027 on 25 degrees of freedom
## Multiple R-squared: 0.8116, Adjusted R-squared: 0.7965
## F-statistic: 53.85 on 2 and 25 DF, p-value: 8.68e-10
```

```
fit2_3a <- lm(Native_log ~ log_Area, data = galapagos_transformed2 %>% filter(!suspicious))
summary(fit2_3a)
```

```
##
## Call:
  lm(formula = Native_log ~ log_Area, data = galapagos_transformed2 %>%
       filter(!suspicious))
##
##
##
  Residuals:
##
       Min
                  1Q
                       Median
                                     3Q
                                             Max
                      0.08423
##
   -1.15996 -0.32394
                               0.35402
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
                                    21.028 < 2e-16 ***
                2.33501
                           0.11104
## (Intercept)
## log Area
                0.28102
                           0.02899
                                      9.693 4.04e-10 ***
## ---
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.5288 on 26 degrees of freedom
## Multiple R-squared: 0.7832, Adjusted R-squared: 0.7749
## F-statistic: 93.95 on 1 and 26 DF, p-value: 4.043e-10
```

(f) Summarize what your analysis has to say about the association of each of the explanatory variables in the data set with the response, after accounting for the explanatory variables in your models. Indicate which of your findings are consistent across the various models considered and which depend on the details of your analysis.

With Native ^0.25 transformation:

All of the models with low BIC showed very strong evidence of a positive association between an island's area and the number of native species found on the island, among islands similar to those in this study. This result held whether or not one high leverage observation and one outlier were included. If those two observations were removed, there was some weak evidence of a negative association between distance from Santa Cruz and the number of native species in the population of islands similar to those in this study, after accounting for the area of the island; this finding is not reliable since it depends on the removal of those two observations. After accounting for the size of the island, there was not evidence of an association between any of the other explanatory variables and the number of native species on the island.

With log(Native) transformation:

All of the models with low BIC showed very strong evidence of a positive association between an island's area and the number of native species found on the island, among islands similar to those in this study. This result held whether or not one high leverage observation and one outlier were included. This is the most reliable finding, since this transformation was more sensitive to the removal of observations (outliers/leverage points). Regardless, after accounting for the size of the island, there was not evidence of an association between any of the other explanatory variables and the number of native species on the island. This is consistent with the conclusions using the first response variable transformation, which is encouraging.

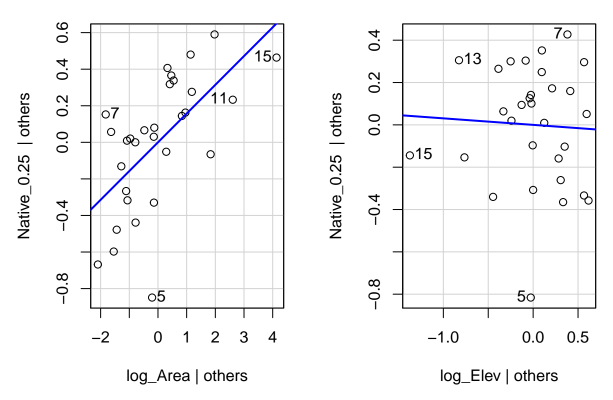
- (g) In this part we'll think through what's going on in a model that includes only your transformed Elev variable and your transformed Area variable.
- i. Fit a model that has your (potentially transformed) Native as the response and your (potentially transformed) Elev and Area variables as explanatory variables. Print the model summary and also use the avPlots function to create added variables plots for these variables.

With Native $^{0.25}$ transformation:

```
fit_both <- lm(Native_0.25 ~ log_Area + log_Elev, data = galapagos_transformed)
summary(fit_both)
##
## Call:
## lm(formula = Native_0.25 ~ log_Area + log_Elev, data = galapagos_transformed)
##
## Residuals:
##
                               ЗQ
      Min
               1Q Median
                                      Max
## -0.8167 -0.1842 0.0616 0.2340 0.4390
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 1.95900
                          0.58985
                                    3.321 0.002579 **
                          0.03821
                                    4.115 0.000327 ***
## log_Area
               0.15724
## log_Elev
              -0.03081
                          0.12025 -0.256 0.799759
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.2931 on 27 degrees of freedom
## Multiple R-squared: 0.7617, Adjusted R-squared: 0.744
## F-statistic: 43.15 on 2 and 27 DF, p-value: 3.903e-09
```

avPlots(fit_both)

Added-Variable Plots

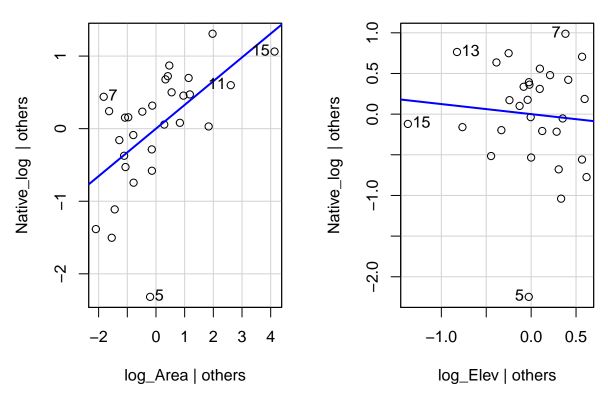


With log(Native) transformation:

```
fit_both2 <- lm(Native_log ~ log_Area + log_Elev, data = galapagos_transformed2)
summary(fit_both2)</pre>
```

```
##
## Call:
## lm(formula = Native_log ~ log_Area + log_Elev, data = galapagos_transformed2)
##
## Residuals:
##
      Min
               1Q Median
                               3Q
                                      Max
## -2.2504 -0.2812 0.1135 0.4521
                                  1.0356
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
##
                          1.36815
                                    2.073 0.047839 *
## (Intercept) 2.83622
## log_Area
               0.32724
                          0.08864
                                    3.692 0.000994 ***
## log_Elev
              -0.12364
                          0.27893 -0.443 0.661107
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.6799 on 27 degrees of freedom
## Multiple R-squared: 0.697, Adjusted R-squared: 0.6745
## F-statistic: 31.05 on 2 and 27 DF, p-value: 9.992e-08
```

Added-Variable Plots



ii. Fit a model that has (potentially transformed) Elev as the response and (potentially transformed) Area as the only explanatory variable. Add the residuals from this model to your data set with transformed variables.

This doesn't depend on the y transformation

```
fit_elev <- lm(log_Elev ~ log_Area, data = galapagos_transformed)
galapagos_transformed <- galapagos_transformed %>%
  mutate(
    resid_elev = residuals(fit_elev)
)

galapagos_transformed2 <- galapagos_transformed2 %>%
  mutate(
    resid_elev = residuals(fit_elev)
)
```

iii. Fit a model that has (potentially transformed) Native as the response and (potentially transformed) Area as the only explanatory variable. Add the residuals from this model to your data set with transformed variables.

With Native^{0.25} transformation:

```
fit_native <- lm(Native_0.25 ~ log_Area, data = galapagos_transformed)
galapagos_transformed <- galapagos_transformed %>%
  mutate(
    resid_native = residuals(fit_native)
)
```

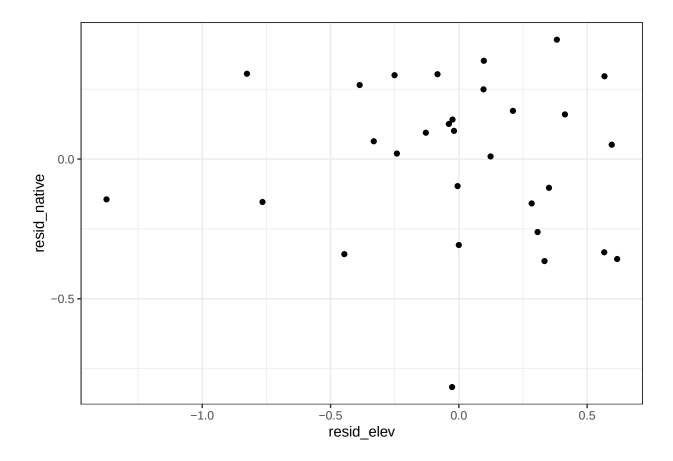
With log(Native) transformation:

```
fit_native2 <- lm(Native_log ~ log_Area, data = galapagos_transformed2)
galapagos_transformed2 <- galapagos_transformed2 %>%
  mutate(
    resid_native2 = residuals(fit_native2)
)
```

iv. Make a plot that has the residuals from part ii on the horizontal axis and the residuals from part iii on the vertical axis. Compare this plot to the added variable plot for Elev from part i.

With Native⁰.25 transformation:

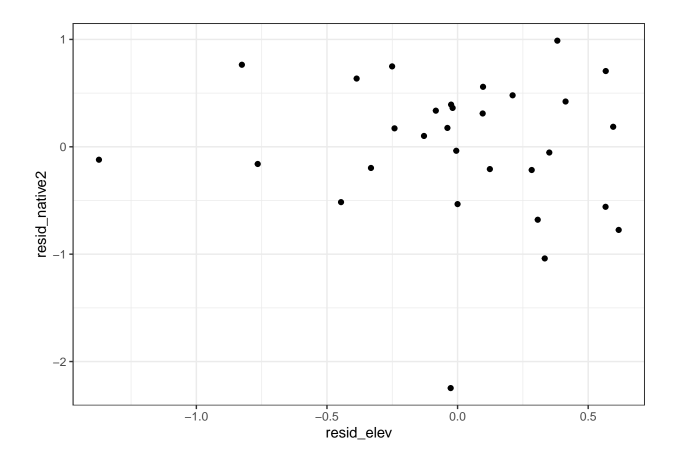
```
ggplot(data = galapagos_transformed, mapping = aes(x = resid_elev, y = resid_native)) +
   geom_point() +
   theme_bw()
```



This plot matches the added variable plot above.

With log(Native) transformation:

```
ggplot(data = galapagos_transformed2, mapping = aes(x = resid_elev, y = resid_native2)) +
  geom_point() +
  theme_bw()
```



This plot matches the added variable plot above.

v. Fit a linear model that has the residuals from part iii as the response and the residuals from part ii as the explanatory variable. Print out the model summary. Compare the coefficient estimate for the slope to the coefficient estimate for Elev from your model in part i.

With Native $^{0.25}$ transformation:

##

```
av_fit <- lm(resid_native ~ resid_elev, data = galapagos_transformed)</pre>
summary(av_fit)
##
## Call:
## lm(formula = resid_native ~ resid_elev, data = galapagos_transformed)
##
## Residuals:
##
                1Q Median
                                ЗQ
                                       Max
  -0.8167 -0.1842 0.0616 0.2340 0.4390
##
##
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
## (Intercept) -1.528e-17 5.255e-02
                                                 1.000
                                       0.000
## resid_elev -3.081e-02 1.181e-01 -0.261
                                                 0.796
```

```
## Residual standard error: 0.2878 on 28 degrees of freedom
## Multiple R-squared: 0.002425, Adjusted R-squared: -0.0332
## F-statistic: 0.06806 on 1 and 28 DF, p-value: 0.7961
```

The coefficient estimate from this fit is the same as the coefficient estimate for elevation in the fit from part i.

With log(Native) transformation:

```
av_fit2 <- lm(resid_native2 ~ resid_elev, data = galapagos_transformed2)
summary(av_fit2)</pre>
```

```
##
## Call:
## lm(formula = resid_native2 ~ resid_elev, data = galapagos_transformed2)
##
## Residuals:
##
      Min
               1Q Median
                                      Max
## -2.2504 -0.2812 0.1135 0.4521 1.0356
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) 4.278e-17 1.219e-01 0.000
                                               1.000
## resid_elev -1.236e-01 2.739e-01 -0.451
##
## Residual standard error: 0.6676 on 28 degrees of freedom
## Multiple R-squared: 0.007225,
                                   Adjusted R-squared:
## F-statistic: 0.2038 on 1 and 28 DF, \, p-value: 0.6552
```

The coefficient estimate from this fit is the same as the coefficient estimate for elevation in the fit from part i