## Supplementary Material II

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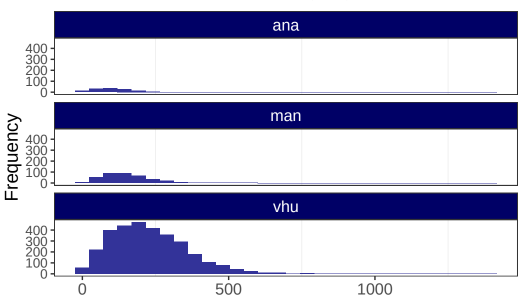
#### **Colony Size**

```
poci_size <- read.csv('coral_size.csv')</pre>
  poci_size.gg <- read_csv("poci_size_main.csv")</pre>
Rows: 3639 Columns: 10
-- Column specification ------
Delimiter: ","
chr (3): level, Site, cover
dbl (7): layer, class, id, area, enn, para, size_cm
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.
  ggplot(poci_size.gg, aes(x = size_cm)) +
    geom_histogram(fill = "#333399") +
    #below here is ylabel, xlabel, and main title
    ylab("Frequency") +
    xlab(NULL) +
    ggtitle(expression("Coral size " (cm**2))) +
    theme bw() +
    facet_wrap(~ Site, ncol = 1) +
    #theme sets sizes, text, etc
    theme(axis.title.x = element_text(size = 14),
          axis.title.y = element_text(size = 14),
          axis.text.y = element_text(size= 10),
          axis.text.x = element_text(size = 12),
```

```
legend.text = element_text(size = 12),
legend.title = element_text(size = 12),
plot.title = element_text(hjust = 0.5, size = 14),
# change plot background, grid lines, etc (just examples so you can see)
panel.background = element_rect(fill = "white"),
panel.grid.minor.y = element_blank(),
panel.grid.major = element_blank(),
plot.background = element_rect(fill = "white"),
legend.background = element_rect(fill = "white"),
strip.text.x = element_text(size = 12, colour = "#FFFFFF"),
strip.background = element_rect(fill = '#000066')
```

`stat\_bin()` using `bins = 30`. Pick better value with `binwidth`.

# Coral size (cm<sup>2</sup>)

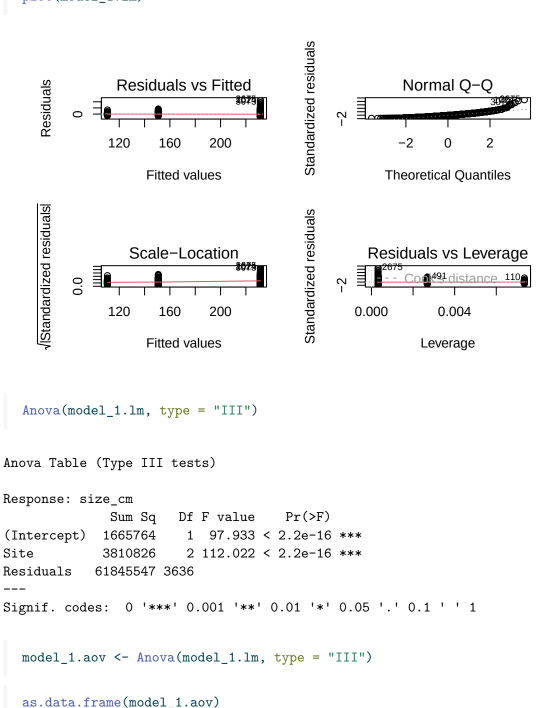


```
poci_size2 <-
poci_size %>%
  as_tibble() %>%
  mutate(size_cm = area*10000) %>%
  group_by(Site) %>%
```

```
dplyr::summarize(mean = mean(size_cm),
                     sd = sd(size_cm),
                     n = n(),
                     se = sd/sqrt(n)
    ) %>%
    mutate(se = sd / sqrt(n),
           lower.ci = mean - qt(1 - (0.05 / 2), n - 1) * se,
           upper.ci = mean + qt(1 - (0.05 / 2), n - 1) * se) %>%
    mutate_at(vars(Site), factor) %>%
    add column(
            location = c('Anakena', 'Manavai', 'Southeast')
            ) %>%
    mutate_at(vars(location), factor)
  poci_size2
# A tibble: 3 x 8
 Site
                             se lower.ci upper.ci location
        mean
                sd
                       n
                                   <dbl>
 <fct> <dbl> <int> <dbl>
                                            <dbl> <fct>
1 ana
        111. 73.8
                      136 6.33
                                   98.2
                                             123. Anakena
2 man
        151.
              82.7
                      372 4.29
                                   142.
                                             159. Manavai
3 vhu
        231. 137.
                    3131 2.44
                                   227.
                                             236. Southeast
  poci_size.gg <-</pre>
    poci_size %>%
    mutate(size_cm = area*10000) %>%
    as tibble() %>%
    mutate_at(vars(Site), factor)
  poci_size.gg
# A tibble: 3,639 x 10
  layer level class
                        id Site cover
                                          area
                                                  enn para size_cm
   <int> <chr> <int> <int> <fct> <chr>
                                         <dbl> <dbl> <dbl>
                                                              <dbl>
 1
       1 patch
                   1
                         2 ana
                                 coral 0.0165 0.111
                                                       42.0
                                                              165.
2
       1 patch
                                 coral 0.0261 0.111
                                                       32.1
                   1
                         3 ana
                                                              261.
                                 coral 0.0162 0.0598 44.3
3
      1 patch
                   1
                        4 ana
                                                              162.
4
      1 patch
                        5 ana coral 0.0149 0.230
                                                       38.6
                                                              149.
                   1
5
       1 patch
                   1
                                 coral 0.00788 0.535
                                                       61.5
                                                               78.8
                         6 ana
6
       1 patch
                        7 ana
                                 coral 0.0148 0.532
                                                       41.1
                                                              148.
```

```
7
      1 patch
                  1 8 ana
                                coral 0.0109 0.0598 50.6
                                                            109.
 8
                       9 ana coral 0.0101 0.346
                                                     52.6
                                                            101.
      1 patch
                  1
 9
      1 patch
                  1
                    10 ana coral 0.0262 0.192
                                                     38.4
                                                            262.
10
      1 patch
                  1
                       11 ana coral 0.00620 0.137 71.6
                                                             62.0
# i 3,629 more rows
  model_1.lm <- lm(size_cm ~ Site, data = poci_size.gg)</pre>
  model_1.lm
Call:
lm(formula = size_cm ~ Site, data = poci_size.gg)
Coefficients:
(Intercept)
                Siteman
                             Sitevhu
     110.67
                  40.18
                              120.81
  summary(model_1.lm)
Call:
lm(formula = size_cm ~ Site, data = poci_size.gg)
Residuals:
   Min
            1Q Median
                            3Q
-231.46 -91.09 -16.44 70.33 1160.47
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept)
             110.67
                         11.18
                                9.896 < 2e-16 ***
Siteman
             40.18
                         13.07
                                3.074 0.00212 **
Sitevhu
             120.81
                         11.42 10.575 < 2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 130.4 on 3636 degrees of freedom
Multiple R-squared: 0.05804, Adjusted R-squared: 0.05752
F-statistic: 112 on 2 and 3636 DF, p-value: < 2.2e-16
```

```
par(mfrow = c(2, 2))
plot(model_1.lm)
```



```
        Sum Sq
        Df
        F value
        Pr(>F)

        (Intercept)
        1665764
        1 97.93297
        8.382392e-23

        Site
        3810826
        2 112.02231
        6.156818e-48

        Residuals
        61845547
        3636
        NA
        NA
```

```
flextable(model_1.aov)
```

Warning in raw\_block(x, "latex", ...): raw\_block() requires Pandoc >= 2.0.0

Sum Sq	Df	F value	Pr(>F)
1,665,764	1	97.932970	.000000000
3,810,826	2	112.022310	.000000000
61,845,547	3,636		

```
mutate_at(vars(cld), factor)
  poci_size3
# A tibble: 3 x 9
 Site
        mean
                          se lower.ci upper.ci location cld
               sd
                      n
 <fct> <dbl> <int> <dbl>
                                <dbl>
                                        <dbl> <fct>
                                                       <fct>
1 ana
        111. 73.8 136 6.33
                                 98.2
                                         123. Anakena
                                                       a
2 man
        151. 82.7
                    372 4.29
                                142.
                                         159. Manavai
        231. 137.
                                         236. Southeast c
3 vhu
                   3131 2.44
                                227.
```

#### Percent Cover

From Damgaard & Irvine (2019) Using the beta distribution to analyse plant cover data Journal of Ecology. 107:2747-2759

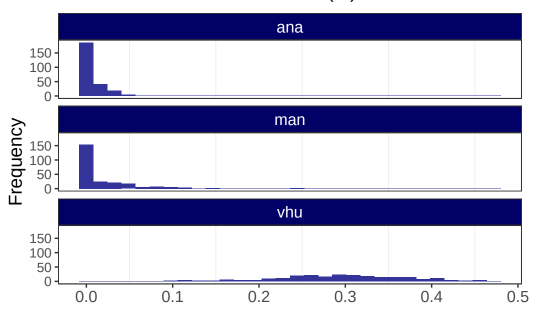
```
p_cover = read_csv('rpn_percent_cover.csv') %>%
    group_by(Site) %>%
    mutate(
     pland_decimal = pland*0.01
    ) %>%
    mutate_at(vars(Site, cover, plot_id), factor)
Rows: 750 Columns: 12
-- Column specification ------
Delimiter: ","
chr (3): Site, cover, level
dbl (7): plot_id, class, layer, percentage_inside, clumpy, pd, pland
lgl (2): id, NA
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.
  p_cover
# A tibble: 750 x 13
# Groups:
           Site [3]
  plot_id class Site cover level layer id percentage_inside clumpy
                                                                      pd
```

```
<dbl> <fct> <fct> <chr> <dbl> <lgl>
                                                            <dbl> <dbl> <dbl>
                                                                   0.992 2.00
1 1
               1 ana
                       coral class
                                        1 NA
                                                            100.
2 1
                       coral class
                                      NA NA
                                                                  NA
                                                                          NA
               0 man
                                                             NA
3 1
                                        1 NA
                                                            100.
                                                                   0.987 19.0
               0 vhu
                       coral class
4 2
                                                                   0.990 8.01
               1 ana
                       coral class
                                        1 NA
                                                             99.9
5 2
               0 man
                       coral class
                                      NA NA
                                                             NA
                                                                  NA
                                                                          NA
6 2
               0 vhu
                       coral class
                                       1 NA
                                                             99.9
                                                                   0.987 20.0
7 3
               1 ana
                       coral class
                                       1 NA
                                                            100.
                                                                   0.995 4.00
8 3
               0 man
                       coral class
                                     NA NA
                                                             NA
                                                                  NA
                                                                         NA
9 3
               0 vhu
                       coral class
                                       1 NA
                                                            100.
                                                                   0.989 16.0
10 4
                                                                   0.995 3.00
                       coral class
                                       1 NA
                                                            100.
               1 ana
# i 740 more rows
# i 3 more variables: pland <dbl>, `NA` <lgl>, pland_decimal <dbl>
  head(p_cover)
# A tibble: 6 x 13
# Groups:
            Site [3]
 plot_id class Site cover level layer id
                                               percentage_inside clumpy
          <dbl> <fct> <fct> <chr> <dbl> <lgl>
                                                           <dbl>
                                                                  <dbl> <dbl>
1 1
              1 ana
                      coral class
                                      1 NA
                                                           100.
                                                                  0.992 2.00
2 1
              0 man
                      coral class
                                     NA NA
                                                            NA
                                                                 NA
                                                                        NA
3 1
                      coral class
              0 vhu
                                                                  0.987 19.0
                                      1 NA
                                                           100.
4 2
              1 ana
                      coral class
                                      1 NA
                                                            99.9
                                                                  0.990 8.01
5 2
                      coral class
                                     NA NA
                                                                 NA
              0 man
                                                            NA
                                                                        NA
              0 vhu
                      coral class
                                      1 NA
                                                            99.9 0.987 20.0
# i 3 more variables: pland <dbl>, `NA` <lgl>, pland_decimal <dbl>
  ggplot(p_cover, aes(x = pland_decimal)) +
    geom_histogram(fill = "#333399") +
    #below here is ylabel, xlabel, and main title
    ylab("Frequency") +
    xlab(NULL) +
    ggtitle(expression("Coral Cover (%)")) +
    theme_bw() +
    facet_wrap(~ Site, ncol = 1) +
    #theme sets sizes, text, etc
    theme(axis.title.x = element_text(size = 14),
          axis.title.y = element_text(size = 14),
          axis.text.y = element_text(size= 10),
```

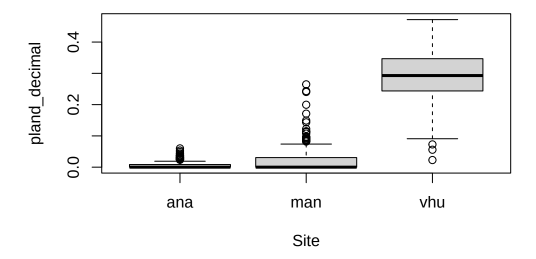
```
axis.text.x = element_text(size = 12),
legend.text = element_text(size = 12),
legend.title = element_text(size = 12),
plot.title = element_text(hjust = 0.5, size = 14),
# change plot background, grid lines, etc (just examples so you can see)
panel.background = element_rect(fill = "white"),
panel.grid.minor.y = element_blank(),
panel.grid.major = element_blank(),
plot.background = element_rect(fill = "white"),
legend.background = element_rect(fill = "white"),
strip.text.x = element_text(size = 12, colour = "#FFFFFF"),
strip.background = element_rect(fill = '#000066')
```

`stat\_bin()` using `bins = 30`. Pick better value with `binwidth`.

### Coral Cover (%)



```
plot(pland_decimal ~ Site, data = p_cover)
```



```
p_cover.mean <-</pre>
    p_cover %>%
    group_by(Site) %>%
    dplyr::summarise(mean = mean(pland_decimal))
  p_cover.mean
# A tibble: 3 x 2
 Site
           mean
          <dbl>
  <fct>
        0.00602
1 ana
2 man
        0.0224
3 vhu
        0.290
```

### Methods for analyzing percent cover of pocilloporid coral

One option for dealing with the 0 and 1 values is to transform them to be slightly less than one or more than zero. This approach assumes that the data are consistent with a common beta distribution. We fit five models to the data: three variations on the beta model and two linear model approaches.

A beta regression assuming a common spatial aggregation  $\delta$  or precision parameter  $(\phi)$  (object named: mod.beta1). Notice that  $\delta = \frac{1}{1+\phi}$  and  $\phi = \frac{(1-\delta)}{\delta}$ . A beta regression assuming each reef location had a different  $\phi$  parameter.

Other options based on assuming that the residuals are normally distributed is to use a linear model with a logit-transformed response or a linear model with response untransformed proportions.

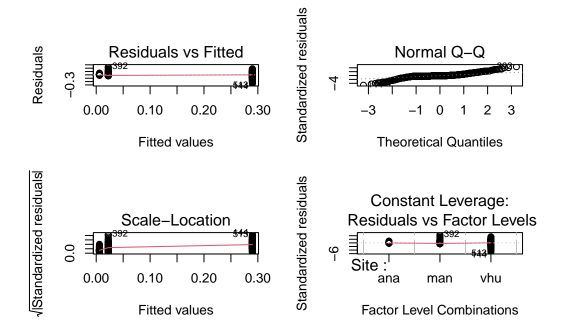
For comparison this applies a logit-transformation to the empirical proportions and then uses a standard linear regression model.

#### Raw data - no transformation

```
p_cover_mod.aov1 <- aov(pland_decimal ~ Site, data = p_cover)</pre>
  summary(p_cover_mod.aov1)
            Df Sum Sq Mean Sq F value Pr(>F)
             2 12.699
                        6.350
                                2203 <2e-16 ***
Site
           747 2.153
                        0.003
Residuals
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
  p_cover_mod.lm1 <- lm(pland_decimal ~ Site, data = p_cover)</pre>
  summary(p_cover_mod.lm1)
Call:
lm(formula = pland_decimal ~ Site, data = p_cover)
Residuals:
     Min
                1Q
                      Median
                                   3Q
                                            Max
-0.267087 -0.022444 -0.006021 0.015304 0.242376
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.006021
                      0.003395
                                 1.773 0.07656 .
Siteman
           0.016423
                      0.004802
                                3.420 0.00066 ***
           Sitevhu
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.05368 on 747 degrees of freedom
```

Multiple R-squared: 0.8551, Adjusted R-squared: 0.8547 F-statistic: 2203 on 2 and 747 DF, p-value: < 2.2e-16

```
par(mfrow = c(2, 2))
plot(p_cover_mod.lm1)
```



```
Anova(p_cover_mod.lm1, type = "III")
```

```
Anova Table (Type III tests)
```

```
Response: pland_decimal
```

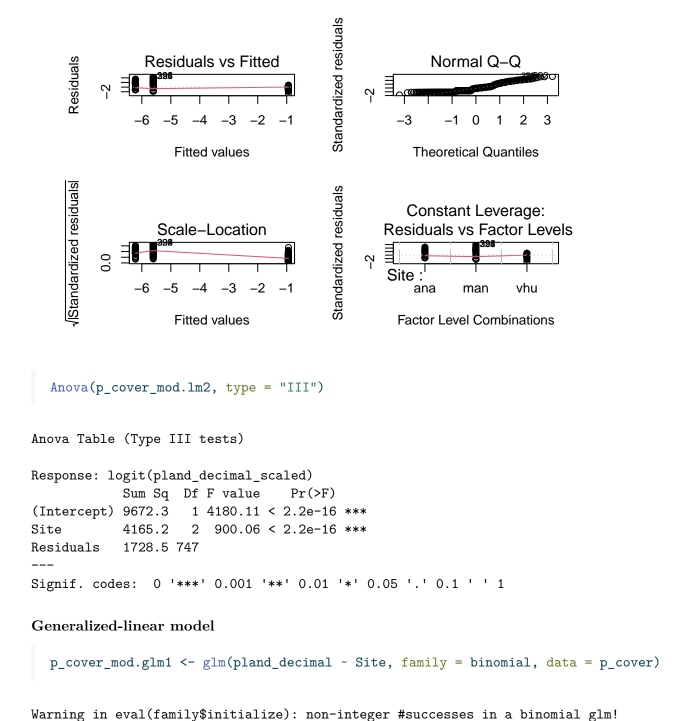
Sum Sq Df F value Pr(>F)
(Intercept) 0.0091 1 3.1451 0.07656 .
Site 12.6991 2 2203.2548 < 2e-16 \*\*\*
Residuals 2.1528 747

---

Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

One option to deal with 0 and 100 percent cover is to add and subtract a small amount to those values

```
transform01 <- function(x) {</pre>
    (x * (length(x) - 1) + 0.5) / (length(x))
  p_cover$pland_decimal_scaled <- transform01(p_cover$pland_decimal)</pre>
Logit-transformation
  p_cover_mod.lm2 <- lm(logit(pland_decimal_scaled) ~ Site, data = p_cover)</pre>
  summary(p_cover_mod.lm2)
Call:
lm(formula = logit(pland_decimal_scaled) ~ Site, data = p_cover)
Residuals:
            1Q Median
                           3Q
                                  Max
-2.7875 -1.0925 -0.1617 0.7543 4.5854
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
0.61521
                     0.13606 4.522 7.13e-06 ***
Siteman
Sitevhu
            5.27828    0.13606    38.795    < 2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Residual standard error: 1.521 on 747 degrees of freedom
Multiple R-squared: 0.7067,
                             Adjusted R-squared: 0.7059
F-statistic: 900.1 on 2 and 747 DF, p-value: < 2.2e-16
  par(mfrow = c(2, 2))
  plot(p_cover_mod.lm2)
```



summary(p\_cover\_mod.glm1)

```
Call:
```

glm(formula = pland\_decimal ~ Site, family = binomial, data = p\_cover)

#### Deviance Residuals:

Min 1Q Median 3Q Max -0.71274 -0.15856 -0.10518 0.07292 0.94245

#### Coefficients:

---

Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

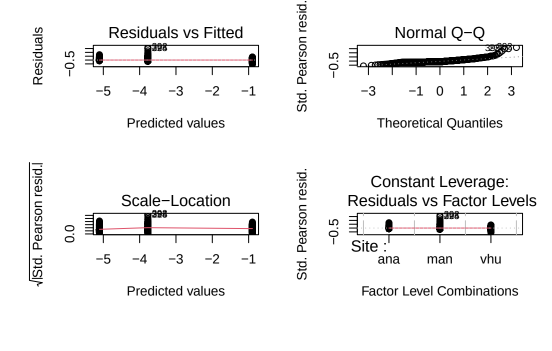
(Dispersion parameter for binomial family taken to be 1)

Null deviance: 161.289 on 749 degrees of freedom Residual deviance: 26.922 on 747 degrees of freedom

AIC: 191.54

Number of Fisher Scoring iterations: 8

```
par(mfrow = c(2, 2))
plot(p_cover_mod.glm1)
```



```
Anova(p_cover_mod.glm1, type = "III")
```

```
Warning in eval(family$initialize): non-integer #successes in a binomial glm!
```

Analysis of Deviance Table (Type III tests)

```
Response: pland_decimal
     LR Chisq Df Pr(>Chisq)
Site    134.37   2   < 2.2e-16 ***
---
Signif. codes:   0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1</pre>
```

#### Beta Regression I: $\phi$ does not vary

```
p_cover_mod.beta1 <- betareg(pland_decimal_scaled ~ Site, data = p_cover, link = c("logit"
summary(p_cover_mod.beta1)</pre>
```

#### Call:

```
betareg(formula = pland_decimal_scaled ~ Site, data = p_cover, link = c("logit"),
    link.phi = NULL, type = c("ML"))
Standardized weighted residuals 2:
             1Q Median
                             3Q
                                    Max
-6.1915 -0.6623 -0.3219 0.6315 1.8412
Coefficients (mean model with logit link):
            Estimate Std. Error z value Pr(>|z|)
                      0.07391 -57.953 <2e-16 ***
(Intercept) -4.28350
Siteman
             0.20537
                        0.08442
                                  2.433
                                           0.015 *
             3.38355
                        0.07733 43.754
Sitevhu
                                           <2e-16 ***
Phi coefficients (precision model with identity link):
      Estimate Std. Error z value Pr(>|z|)
(phi)
        25.321
                    1.561
                            16.22
                                    <2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Type of estimator: ML (maximum likelihood)
Log-likelihood: 2094 on 4 Df
Pseudo R-squared: 0.7048
Number of iterations: 19 (BFGS) + 2 (Fisher scoring)
Beta Regression II: \phi does vary (by Site)
  p_cover_mod.beta2 <- betareg(pland_decimal_scaled ~ Site | Site, data = p_cover, link = c(</pre>
Extract AIC from beta regression models
  p_cover_mod.beta1_aic <- AIC(p_cover_mod.beta1)</pre>
  p_cover_mod.beta2_aic <- AIC(p_cover_mod.beta2)</pre>
  p_cover_mod.beta1_aic
[1] -4179.365
  p_cover_mod.beta2_aic
[1] -4259.399
```