

Statistical_Analysis_Simpson

Daniela Linero

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1. Load data

Load the table that contains the abundance and occurrence measures for frugivores along with the plants dispersed and not dispersed by animals.

```
# Import table
diversityS <- readRDS(file = "./output/cleaned_data/01_Filter_data_frugi_endooPlants_notEndooPlants_rec...
```

2. Correct abundance measures using Sampling effort

```
diversityS <- yarg::CorrectSamplingEffort(diversityS)
```

```
## Correcting 0 missing sampling effort values
## Rescaling sampling effort
## Correcting 114923 values for sensitivity to sampling effort
```

3. Select studies that assessed more than one species

```
# Create a dataset with the SS that assessed more than 1 species
list <- diversityS %>%
  # By grouping by SS we are comparing sites with the same diversity metric
  # type either abundance or occurrence
  dplyr::group_by(SS) %>%
  # Create a new column to calculate the number of species sampled per study
  # n_distinct = length(unique())
  dplyr::mutate(N_species_sampled = dplyr::n_distinct(Taxon_name_entered)) %>%
  # ungroup data frame
  dplyr::ungroup() %>%
  # Filter the studies that sampled more than 1 species
  dplyr::filter(N_species_sampled > 1) %>%
  base::droplevels()
```

```
# Create a character vector of unique SS that assessed more than 1 species
list <- as.character(unique(list$SS))
```

```

diversityS <- diversityS %>%
  # Filter the studies present in the list
  base::subset(SS %in% list) %>%
  droplevels()

```

4. Remove sites that will produce NaN values

```

diversityS <- diversityS %>%
  # Remove the sites that have a maximum abundance of 0
  base::subset(SSBS %nin% c("MJ1_2009__Lehouck 2 Fururu 10",
                            "MJ1_2009__Lehouck 2 Fururu 11",
                            "MJ1_2009__Lehouck 2 Macha 12",
                            "MJ1_2009__Lehouck 2 Macha 13")) %>%
  base::droplevels()

```

5. Split dataset

I will separate the measures of plants not dispersed by animals, in order to avoid that they are merged together with plants dispersed by animals with the MergeSites function

```

# I am going to separate the records that belong to plants not dispersed
# by animals
diversityS_notEndo <- diversityS %>%
  base::subset(Kingdom == "nePlantae") %>%
  base::droplevels()

# Get the table without plants dispersed by animals
diversityS_frugi_endo <- diversityS %>%
  base::subset(Kingdom != "nePlantae") %>%
  base::droplevels()

```

6. Merge Sites

```

diversityS_frugi_endo <- yarg::MergeSites(diversityS_frugi_endo,
                                             silent = TRUE,
                                             merge.extra = "Wilderness_area")

diversityS_notEndo <- yarg::MergeSites(diversityS_notEndo,
                                         silent = TRUE,
                                         merge.extra = "Wilderness_area")

```

7. Rename Predominant Habitat

```

# Rename the column predominant habitat, as the dataset is actually
# referring to land use

```

```


diversityS_frugi_endo <- dplyr::rename(diversityS_frugi_endo,  
                                         Predominant_land_use = Predominant_habitat)



diversityS_notEndo <- dplyr::rename(diversityS_notEndo,  
                                         Predominant_land_use = Predominant_habitat)


```

8. Calculate Diversity Metrics

```


# Calculate diversity metrics for animals and endoozoocoric plants  
diversity1S_frugi_endo <- diversityS_frugi_endo %>%  
  
  # add Diversity_metric_is_valid column  
  dplyr::mutate(Diversity_metric_is_valid = TRUE) %>%  
  
  # The extra.cols parameter is used for columns that we want to  
  # transferred to the final site-level data frame and that the function  
  # does not add automatically  
  args::SiteMetrics(extra.cols = c("SSB", "SSBS", "Predominant_land_use", "Kingdom"))



# Calculate diversity metrics for plants not dispersed by animals  
diversity1S_notendo <- diversityS_notEndo %>%  
  
  # add Diversity_metric_is_valid column  
  dplyr::mutate(Diversity_metric_is_valid = TRUE) %>%  
  
  # The extra.cols parameter is used for columns that we want to  
  # transferred to the final site-level data frame and that the function  
  # does not add automatically  
  args::SiteMetrics(extra.cols = c("SSB", "SSBS", "Predominant_land_use", "Kingdom"))



# Merge the site metrics for all organisms  
diversity_all <- base::rbind.data.frame(diversity1S_frugi_endo,  
                                         diversity1S_notendo)


```

9. Check results

```


# Merge the measures of frugivores, endoplants and not endoplants  
diversityS_combined <- rbind.data.frame(diversityS_frugi_endo,  
                                         diversityS_notEndo)  
  
diversityS_combined %>%  
  
  # subset one site to check  
  base::subset(SSS == "DG1_2013_Zou 1 1") %>% base::droplevels() %>%  
  
  # Calculate each species abundance over total abundance at the site  
  dplyr::mutate(Proportion = Measurement/sum(Measurement)) %>%  
  
  # Sum the proportions and squared them  
  dplyr::mutate(Simpson_D = sum(Proportion^2)) %>%  
  
  # Calculate 1/D  
  dplyr::mutate(one_over_D = 1/Simpson_D) %>%


```

```

# Select only columns we're interested in
dplyr::select(SSS, Best_guess_binomial, Measurement, one_over_D)

##          SSS  Best_guess_binomial Measurement one_over_D
## 1 DG1_2013__Zou 1 1      Megalaima oorti    20.58333  3.698822
## 2 DG1_2013__Zou 1 1     Pycnonotus sinensis   48.75000  3.698822
## 3 DG1_2013__Zou 1 1     Alophoixus pallidus   34.66667  3.698822
## 4 DG1_2013__Zou 1 1     Hemixos castanonotus  35.75000  3.698822

# Compare with the SiteMetrics result
diversity_all %>% base::subset(SSS == "DG1_2013__Zou 1 1") %>%
  dplyr::select(SSS, Simpson_diversity)

##          SSS Simpson_diversity
## 462 DG1_2013__Zou 1 1           3.698822

```

10. Check the abundance measures units

I am going to check that the measures used that weren't number of individuals or ind/km², were reasonably concordant to an abundance measure of each species.

```

# Get the unique diversity metric type and unit for each study
Diversity_metric_unit <- diversityS_combined %>%

# Get only one row for each study
dplyr::distinct(SS, .keep_all = TRUE) %>%

# Select the columns we're interested in
dplyr::select(Source_ID, SS,
             Diversity_metric_unit, Kingdom,
             Study_common_taxon) %>%

# Filter the studies that need to be checked
base::subset(Diversity_metric_unit %in% c("individuals",
                                            "stems/hectare",
                                            "presence/absence",
                                            "effort-corrected individuals",
                                            "individuals/km2",
                                            "groups/colonies per km",
                                            "number of groups",
                                            "individuals/km")) %>%
  droplevels()

kable(Diversity_metric_unit, format="latex", booktabs=TRUE) %>%
  kable_styling(latex_options="scale_down")

```

Source_ID	SS		Diversity_metric_unit	Kingdom	Study_common_taxon
18 DI1_2011_Dawson	DI1_2011_Dawson 1	proportion of plots	Animalia	Aves	
38 HB1_1998_Svenning	HB1_1998_Svenning 1	percentage	Plantae	Arecaceae	
41 HP1_2005_Hietz	HP1_2005_Hietz 1	percentage of sites	Plantae	Tracheophyta	
92 SH1_2002_Sheil	SH1_2002_Sheil 1	number of plots	Plantae	Plantae	
96 SH1_2013_CIFORcameroon	SH1_2013_CIFORcameroon 1	number of plots	Plantae	Plantae	
98 SH1_2013_CIFORphilippines	SH1_2013_CIFORphilippines 1	number of plots	Plantae	Plantae	
99 SH1_2013_CIFORphilippines	SH1_2013_CIFORphilippines 2	number of plots	Plantae	Tracheophyta	
108 YY1_2015_Mandle	YY1_2015_Mandle 1	percentage	Plantae	Tracheophyta	
111 YY1_2016_Mohandass	YY1_2016_Mohandass 1	percentage	Plantae		

11. Models

Count number of sites for the SiteMetrics table in order to know the sample size for each land-use type and intensity.

```
# Remove sites that don't have a simpson diversity value or land-use type
diversity_simpson <- drop_na(diversity_all,
                                Simpson_diversity,
                                Predominant_land_use) %>%
  droplevels()

# Number of sites
table(diversity_simpson$Predominant_land_use,
      diversity_simpson$Use_intensity,
      diversity_simpson$Kingdom)
```

	Minimal use	Light use
## , , = Animalia		
##		
##		
##		
## Primary forest	325	113
## Primary non-forest	0	0
## Young secondary vegetation	37	8
## Intermediate secondary vegetation	81	50
## Mature secondary vegetation	11	4
## Secondary vegetation (indeterminate age)	27	6
## Plantation forest	27	207
## Pasture	0	4
## Cropland	59	20
## Urban	0	29
##		
##		
##		
## Intense use		
## Primary forest	15	9
## Primary non-forest	0	0
## Young secondary vegetation	0	22
## Intermediate secondary vegetation	2	26
## Mature secondary vegetation	0	21
## Secondary vegetation (indeterminate age)	0	10
## Plantation forest	18	0
## Pasture	1	24
## Cropland	40	18
## Urban	1	0
##		
## , , = Plantae		

```

##                                     Minimal use Light use
## Primary forest                      247      55
## Primary non-forest                   2        0
## Young secondary vegetation          55      15
## Intermediate secondary vegetation   36      72
## Mature secondary vegetation         34       8
## Secondary vegetation (indeterminate age) 7      15
## Plantation forest                  71     166
## Pasture                           19      17
## Cropland                          18       0
## Urban                            0        0
##
##                                     Intense use Cannot decide
## Primary forest                      70      59
## Primary non-forest                   0        0
## Young secondary vegetation          0        1
## Intermediate secondary vegetation   2      11
## Mature secondary vegetation         6       0
## Secondary vegetation (indeterminate age) 0     112
## Plantation forest                  18       3
## Pasture                           2        0
## Cropland                          36       8
## Urban                            0        0
##
## , , = nePlantae
##
##                                     Minimal use Light use
## Primary forest                      196      41
## Primary non-forest                   3        1
## Young secondary vegetation          61      11
## Intermediate secondary vegetation   38      31
## Mature secondary vegetation         10       7
## Secondary vegetation (indeterminate age) 8      18
## Plantation forest                  68      65
## Pasture                           19      24
## Cropland                          19       0
## Urban                            0        0
##
##                                     Intense use Cannot decide
## Primary forest                      50      51
## Primary non-forest                   0        0
## Young secondary vegetation          0        3
## Intermediate secondary vegetation   2      18
## Mature secondary vegetation         3       0
## Secondary vegetation (indeterminate age) 0     118
## Plantation forest                  4        6
## Pasture                           4        0
## Cropland                          6        8
## Urban                            0        0

```

According to the number of sites, I am going to try this initial combination:

- Primary can be divided into the two level intensities in all cases
- Cropland has to be merged
- ISV can be divided in minimal use and light/intense use
- MSV has to be merged
- Pasture has to be merged
- Plantation forest can be divided in minimal use and light/intense
- YSV has to be merged

```
# Call the function that merges lan-uses and intensities
source("./R/02_Statistical_Analysis_merge_LandUses_Intensities.R")

# Create the vectors that hold the land-uses that we want to
# keep with different use intensities
land_uses_separate_1 <- c("Primary", "ISV", "Plantation forest")

# Create a vector with the land-uses where we want to merge the
# light and intense use intensities
land_uses_light_intense_1 <- c("Primary", "ISV", "Plantation forest")

# Merge landuse intensities
diversity_simpson <- Merge_landUses_and_intensities(dataset = diversity_simpson,
                                                       index = 1,
                                                       land_uses_separate_intensities = land_uses_separate_1,
                                                       land_uses_merge_light_intense = land_uses_light_intense_1,
                                                       "Primary Minimal use")

# Check number of sites
addmargins(table(diversity_simpson$LandUse.1, diversity_simpson$Kingdom), 2)
```

```
##                                     Animalia Plantae nePlantae Sum
## Primary Minimal use                  325    249     199 773
## Cropland All                         137     62      33 232
## ISV Light-intense use                 52     74      33 159
## ISV Minimal use                      81     36      38 155
## MSV All                             36     48      20 104
## Pasture All                          29     38      47 114
## Plantation forest Light-intense use  225    184      69 478
## Plantation forest Minimal use         27     71      68 166
## Primary Light-intense use            128    125      92 345
## YSV All                            67     71      75 213
```

Test for collinearity

```
# Get the function
source("https://highstat.com/Books/Book2/HighstatLibV10.R")

# Calculate the VIF
corvif(diversity_simpson[ , c("LandUse.1", "Kingdom")])
```

```
##                                     Animalia Plantae nePlantae Sum
## Primary Minimal use                  325    249     199 773
## Cropland All                         137     62      33 232
## ISV Light-intense use                 52     74      33 159
## ISV Minimal use                      81     36      38 155
## MSV All                             36     48      20 104
## Pasture All                          29     38      47 114
## Plantation forest Light-intense use  225    184      69 478
## Plantation forest Minimal use         27     71      68 166
## Primary Light-intense use            128    125      92 345
## YSV All                            67     71      75 213
```

```

##                                     GVIF Df GVIF^(1/2Df)
## LandUse.1 1.065772  9     1.003545
## Kingdom    1.065772  2     1.016052

```

Choose random effects structure

```

# Simplest random effects structure
m1 <- lmer(Simpson_diversity ~ LandUse.1 + Kingdom + LandUse.1:Kingdom +
             (1|SS) + (1|SSB), data = diversity_simpson)

```

```

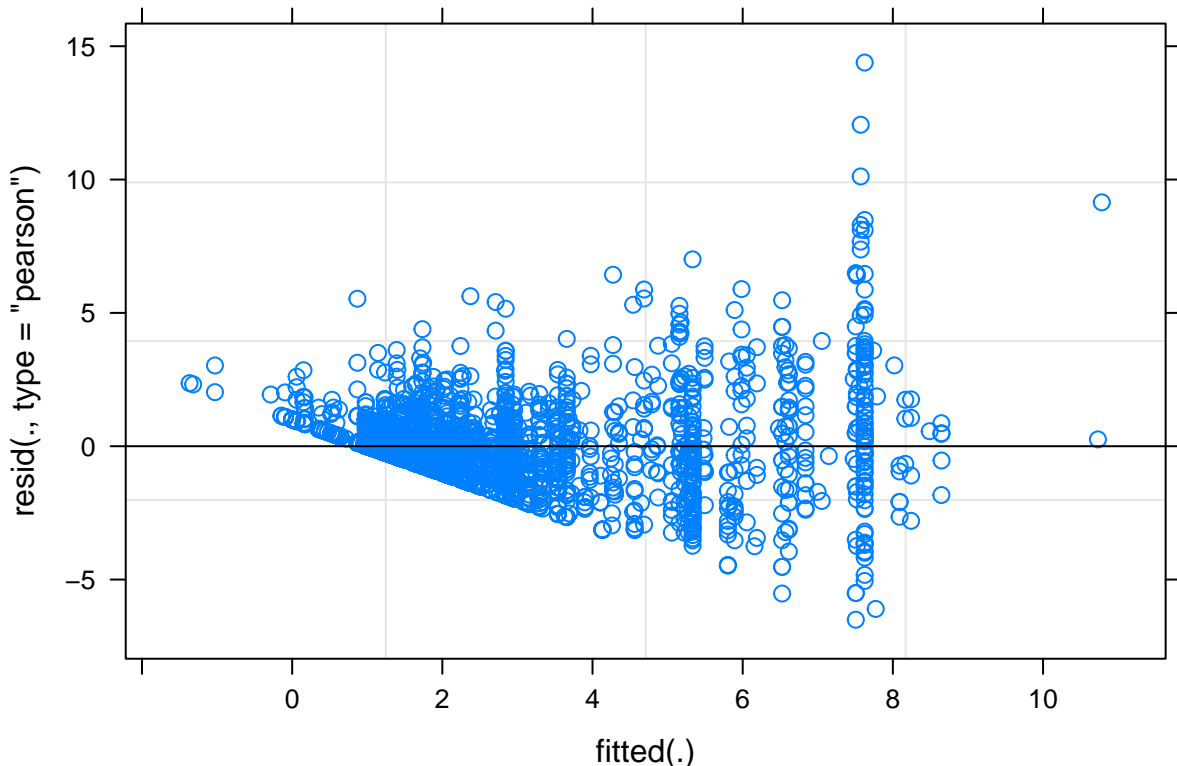
## boundary (singular) fit: see ?isSingular

```

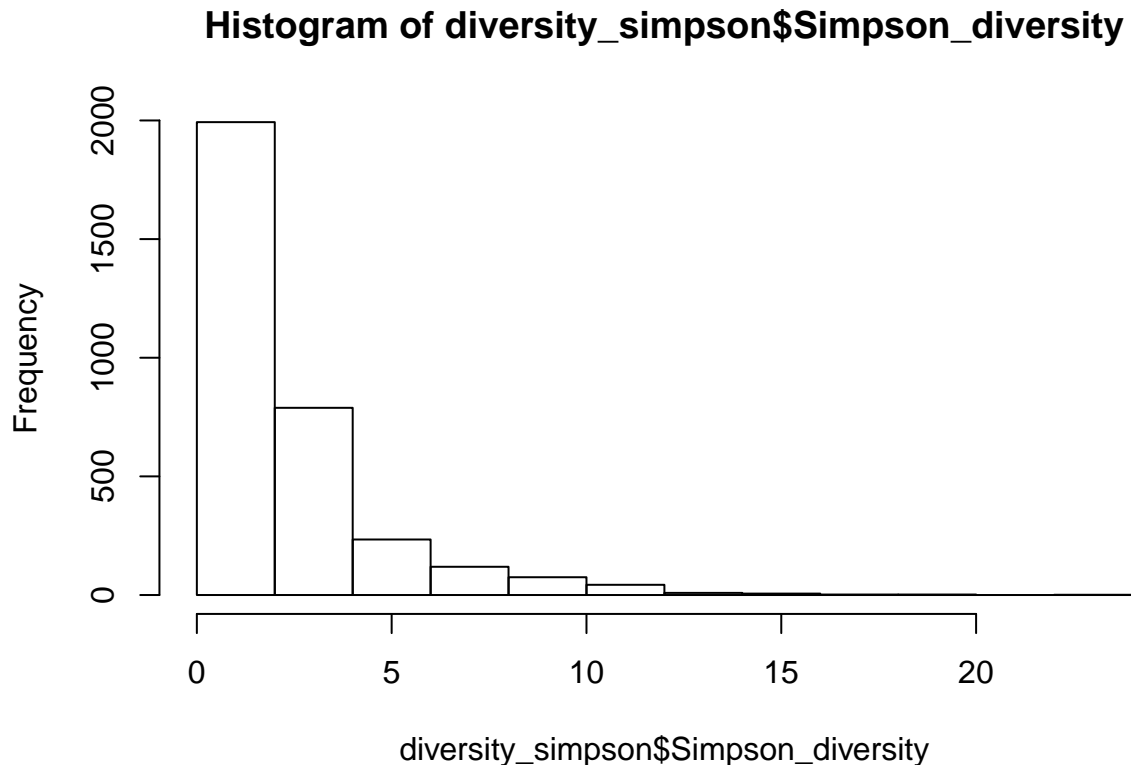
```

plot(m1)

```



```
hist(diversity_simson$Simpson_diversity)
```



```
# The simplest model does not converge
```

12. Second attempt

```
# I am going to merge all categories

# Create the vectors that hold the land-uses that we want to keep
# with different use intensities
land_uses_separate_2 <- "NA"
# Create a vector with the land-uses where we want to merge the
# light and intense use intensities
land_uses_light_intense_2 <- "NA"

diversity_simson <- Merge_landUses_and_intensities(dataset = diversity_simson,
                                                    index = 2,
                                                    land_uses_separate_intensities = land_uses_separate_2,
                                                    land_uses_merge_light_intense = land_uses_light_intense_2,
                                                    "Primary All")

addmargins(table(diversity_simson$LandUse.2, diversity_simson$Kingdom), 2)

##
```

```

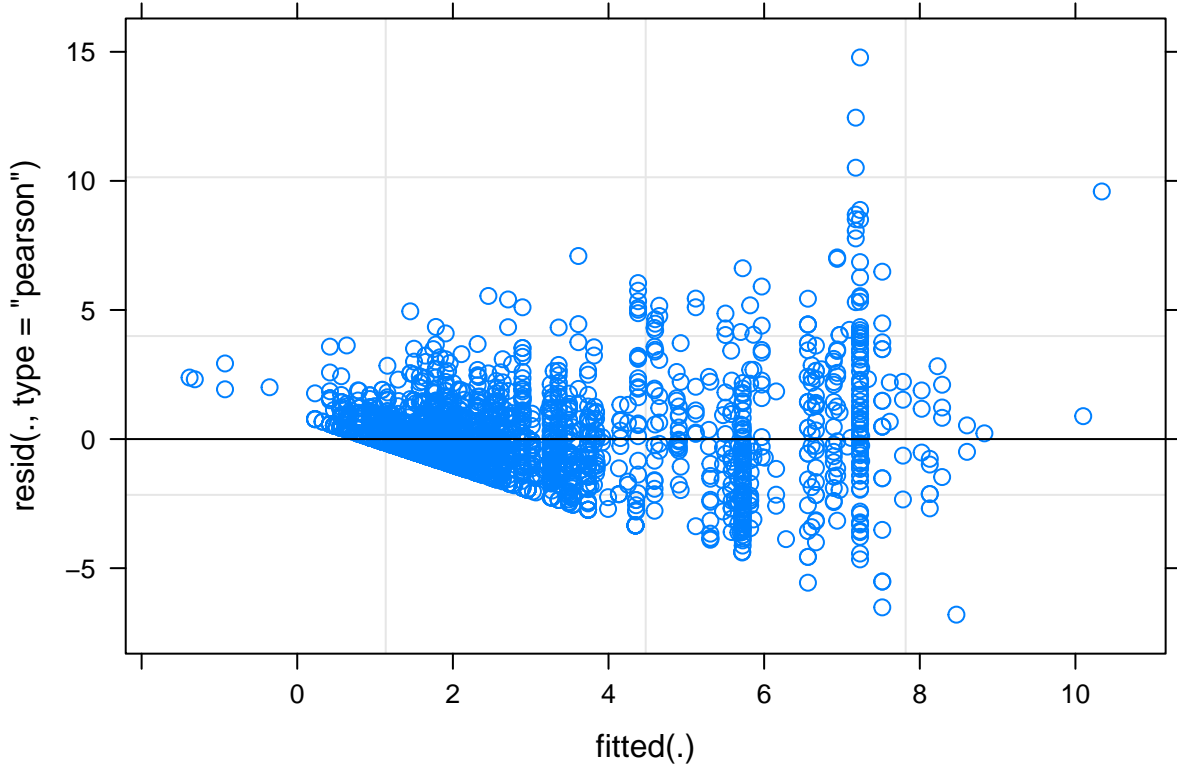
##                                     Animalia Plantae nePlantae Sum
## Primary All                      462     433      342 1237
## Cropland All                     137      62       33  232
## ISV All                          159     121      89  369
## MSV All                          36      48       20 104
## Pasture All                      29      38       47 114
## Plantation forest All            252     258      143 653
## YSV All                          67      71       75 213

m2 <- lmer(Simpson_diversity ~ LandUse.2 + Kingdom + LandUse.2:Kingdom +
           (1|SS) + (1|SSB), data = diversity_simpson)

## boundary (singular) fit: see ?isSingular

plot(m2)

```



I am going to try to remove the MSV, since it only has 20 sites in total for nePlants

```

# Remove MSV records
diversity_simpson1 <- diversity_simpson %>%
  subset(LandUse.2 != "MSV All") %>% droplevels()

# Run the model
m3 <- lmer(Simpson_diversity ~ LandUse.2 + Kingdom + LandUse.2:Kingdom +
           (1|SS) + (1|SSB), data = diversity_simpson1) #Is Singular

```

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

```
plot(m3)
```

