Practical 13: Biodiversity and Productivity APES2039A

Dr. Joseph White

03/06/2022

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1 Background

In this practical we will explore the relationships between different types of biodiversity (species richness, functional diversity, phylogenetic diversity) and productivity (Net Primary Productivity).

We will make use of two different datasets.

- 1) Plants of South Africa (POSA) specimen and habit dataset
- 2) MODIS Terra satellite's derived annual Net Primary Productivity (NPP) at 500m pixel resolution.

The POSA specimen and habit dataset is an extensive collection of plant samples with locations from all of South Africa collected over more than 200 years. These include herbarium records and verified observations amongst others. Additionally, this dataset has information on the type of habit a plant has (e.g. tree, shrub, herb), which we will use as our functional type data.

The MODIS NPP dataset provides annual derived productivity values across the globe. The dataset represents the average NPP values from 2001-2021.

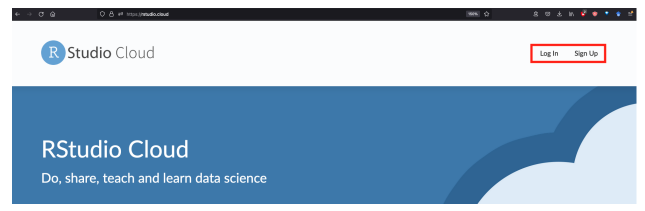
The data for both POSA and NPP is subsetted to only include values for the grassland biome of South Africa. These are linked to a location across the country in the form of a grid system referred to as a Quarter Degree Square (QDS). Each grid cell is roughly 30×30 km in size.

This document provides the code and output (summary values, maps, figures and tables) that you will work from to extract the relevant answers. All of the answers you require for your Practical 13 worksheet submission will be found in this document, your lecture notes from Week 14 Lecture 3 Functional consequences of biodiversity and your own experience and understanding of sampling field data.

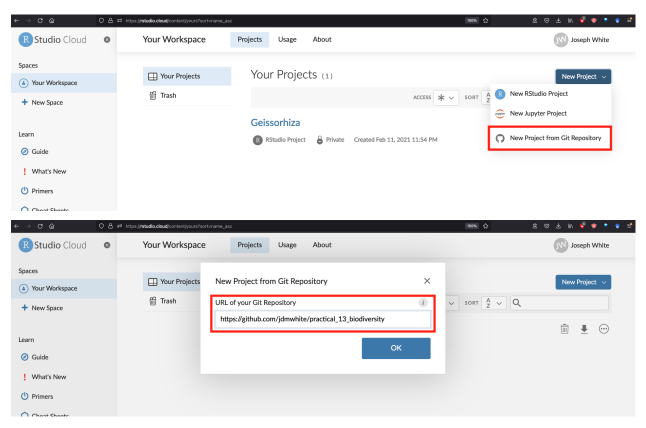
2 If you would like to follow along and run the code yourself

2.1 R Studio Cloud

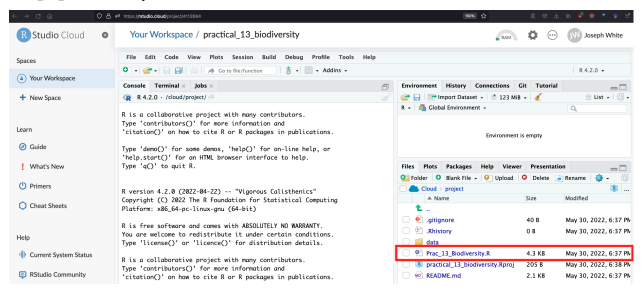
Go to R Studio Cloud. Either **Sign Up** or as you have already run a practical here before, **Log In** to your account.



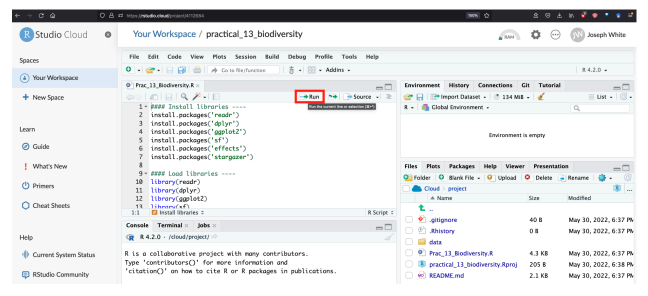
Once you have logged in, click on **New Project** in the top right corner. Select **New Project from Git Repository**. Copy and paste this link into the open space: https://github.com/jdmwhite/practical_13_biodiversity



Wait for the project to deploy (this may take a couple of minutes). Once it has opened, click on the Prac_13_Biodiversity.R file.



You can now run the code, either using (Cntrl + ENTER) or click the **Run** button in the top middle of your screen.



Make sure to **LOG OUT** of your R Studio Cloud session when you are finished using the platform. You are granted 25 free hours per month. But these hours will be depleted quickly if you don't log out!

2.2 R Studio run locally on your computer

- Step 1: Download R: https://cran.r-project.org/ (select the correct operating system for your computer)
- Step 2: Download R Studio: https://www.rstudio.com/products/rstudio/download/#download (select the correct operating system for your computer)

Once you have download and installed both, all you need is to open R Studio. The next step is to go to https://github.com/jdmwhite/practical_13_biodiversity, click "Code" in the top right corner and the click "Download ZIP". Unzip this file, by double-clicking on it, and then open the practical_13_biodiversity.Rproj file. Make sure to open the file with the .Rproj file extension. This will automatically open R Studio and begin your session. Once it has opened, click on the Prac_13_Biodiversity.R file. You can now run the code, either using (Cntrl + ENTER) or click the Run button in the top middle of your screen (as shown above in the R Studio Cloud example).

3 Let's code

3.1 Install and load libraries

The first step in most coding languages is to install and load the libraries (normally called packages). These packages have special functions, which users created for specific tasks. The huge variety of R packages are one of the major reasons R is so popular with the ecology community.

```
#### Load libraries ----
library(readr)
library(dplyr)
library(ggplot2)
library(sf)
library(effects)
library(stargazer)
```

Now we will load in the data. The POSA data is stored in a csv file. This file type is similar to an excel file, but is more efficient in its use of space.

We will also load in the QDS and covariate dataset. This is a **shapefile**, which is a *spatial* file format. It is a table, with each row representing one grid cell (QDS). The columns include variables associated with each QDS. Then there is a special column called **geometry**. This represents the *spatial* component of the table, and tells R where each polygon or point is found in space.

We will also load the South African border **shapefile** for our maps.

```
#### Load in data ----
# POSA richness
posa_rich <- read_csv('data/posa_richness.csv')

# QDS and covariates
qds_cov <- read_sf('data/grass_qds_cov.shp')

# Map of SA
sa <- read_sf('data/SA_border/SA_border.shp')</pre>
```

3.2 Explore the data

3.2.1 POSA data

Now that the data is loaded into our working environment, let's take a look at it:

```
#### Explore the data ----
#### Plants of SA data ----

# Take a look at the data
posa_rich
```

```
## # A tibble: 86,859 x 7
##
      QDS
             Taxon
                          Family func_type spp_rich func_diversity phylog_diversity
##
      <chr> <chr>
                           <chr> <chr>
                                               <dbl>
                                                               <dbl>
                                                                                 <dbl>
##
   1 3030CA Melica race~ Poace~ graminoid
                                                 731
                                                                  24
                                                                                   139
  2 2528CC Themeda tri~ Poace~ graminoid
                                                 972
                                                                  24
                                                                                   139
##
   3 3226DB Indigofera ~ Fabac~ herb
                                                1110
                                                                  23
                                                                                   165
##
  4 2731CB Pavetta ede~ Rubia~ shrub
                                                                  18
                                                                                   100
                                                 449
##
   5 2831CD Miscanthus ~ Poace~ graminoid
                                                 990
                                                                  24
                                                                                   182
  6 2930CB Asterella b~ Ayton~ bryophyte
                                                                  26
                                                                                   219
                                                1578
  7 3029CB Eragrostis ~ Poace~ graminoid
                                                 574
                                                                  20
                                                                                   98
## 8 2926AA Paspalum di~ Poace~ graminoid
                                                 605
                                                                  18
                                                                                   96
## 9 3128AA Gladiolus l~ Irida~ geophyte
                                                 631
                                                                  19
                                                                                   101
## 10 2930CB Schoenoplec~ Cyper~ cyperoid
                                                1578
                                                                  26
                                                                                   219
## # ... with 86,849 more rows
```

As we can see it is a big dataset! 86859 rows and 7 columns. By default R only shows us the first 10 rows. We have information about the QDS (the grid cell the species is found in), the taxon name, the plant family, the functional type, and then summary information for each QDS on the species richness, functional diversity and phylogenetic diversity.

Next, let's count what the most common species across all of the QDSs is. We will use the count() function and arrange() function. count() tells us how many times each species name is found. arrange() then sorts this for us. We use a negative sign to tell R we would like the result in *descending* order.

```
# What is the species found in most Quarter Degree Square (QDS)?
posa_rich %>% count(Taxon) %>% arrange(-n)
```

```
## # A tibble: 7,278 x 2
##
     Taxon
##
      <chr>>
                             <int>
## 1 Eragrostis curvula
                               281
## 2 Helichrysum nudifolium
                               155
## 3 Diospyros lycioides
                               147
## 4 Searsia pyroides
                               147
## 5 Themeda triandra
                               147
## 6 Commelina africana
                               134
## 7 Setaria sphacelata
                               134
## 8 Scabiosa columbaria
                               132
## 9 Digitaria eriantha
                               131
## 10 Helichrysum rugulosum
                               124
## # ... with 7,268 more rows
```

We will now continue to look at different summary metrics for our POSA dataset.

How many different plant families are there?

```
# How many different plant families are there?
n_distinct(posa_rich$Family)
```

```
## [1] 319
```

What are the different functional types? To make R print out more rows, we specify n=29 within the print function.

```
# What different functional types are there?
posa_rich %>% distinct(func_type) -> functional_types
print(functional_types, n = 29)
```

```
## # A tibble: 29 x 1
##
     func_type
##
      <chr>
##
  1 graminoid
##
  2 herb
##
  3 shrub
## 4 bryophyte
## 5 geophyte
  6 cyperoid
##
##
  7 tree
## 8 helophyte
## 9 scrambler
## 10 succulent
## 11 dwarf shrub
## 12 emergent hydrophyte
## 13 climber
## 14 mesophyte
## 15 suffrutex
```

```
## 16 hydrophyte
## 17 parasite
## 18 epiphyte
## 19 lithophyte
## 20 carnivore
## 21 tenagophyte
## 22 epihydate
## 23 pleustophyte
## 24 hyperhydate
## 25 restioid
## 26 sudd hydrophyte
## 27 creeper
## 28 haptophyte
## 29 lichen
```

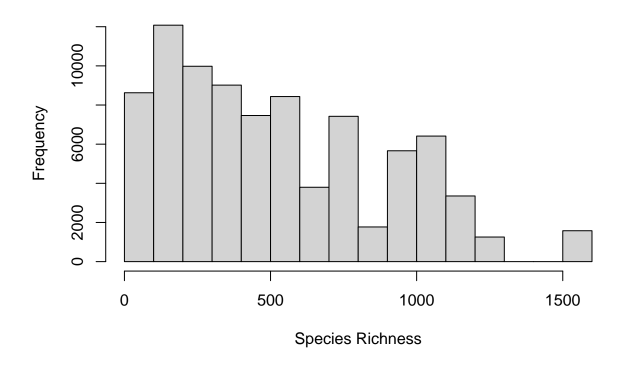
What is the most common functional type? Again, we use the count() and arrange() functions.

```
# What is the most common functional type?
posa_rich %>% count(func_type) %>% arrange(-n)
```

```
## # A tibble: 29 x 2
##
     func_type
                     n
##
      <chr>
                  <int>
   1 herb
##
                 29910
                 10638
   2 shrub
  3 graminoid
                  9766
##
## 4 geophyte
                  7609
## 5 succulent
                  7006
   6 dwarf shrub 4369
##
   7 bryophyte
                  2768
## 8 tree
                   2611
                  2572
## 9 climber
## 10 lithophyte
                  2171
## # ... with 19 more rows
```

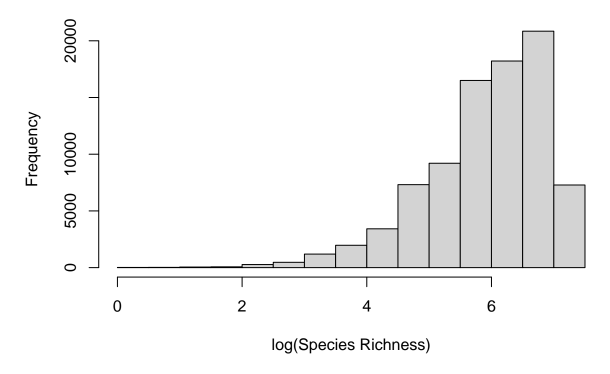
Before we do any analysis of our data, it is important to consider whether our data is **normally distributed** or not. This is due to the *assumptions* that some models, such as linear regression make. Your input data does not necessarily need to be normally distributed, but rather your prediction error (or residuals). If you want to learn more on this, look HERE. We will log-transform our species richness data to use later in our regression analysis.

Histogram of Species Richness



```
hist(log(posa_rich$spp_rich), xlab = 'log(Species Richness)',
    main = 'Histogram of log-transformed Species Richness')
```

Histogram of log-transformed Species Richness



3.2.2 NPP data

Now we have an idea of what's inside our POSA data. Let's look at our Net Primary Productivity data, which is stored in the QDS shapefile:

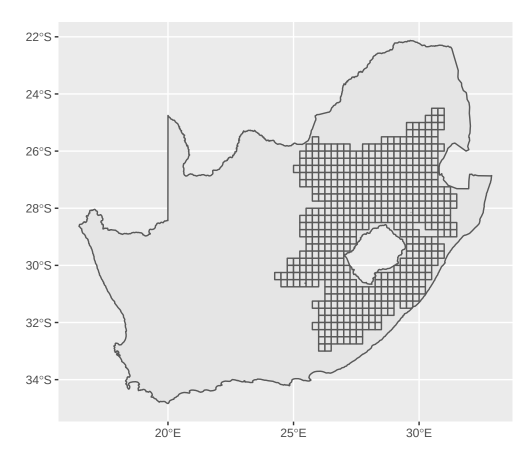
```
#### Spatial data (QDS) ----
qds_cov
## Simple feature collection with 510 features and 5 fields
## Geometry type: POLYGON
## Dimension:
                  XY
## Bounding box:
                  xmin: 24.25 ymin: -33 xmax: 31.5 ymax: -24.5
## Geodetic CRS:
                  WGS 84
## # A tibble: 510 x 6
##
      QDS
               MAP Elev_STD
                              NPP NPP_var
                                                                             geometry
                                                                        <POLYGON [°]>
##
      <chr>
             <dbl>
                      <dbl> <dbl>
                                     <dbl>
   1 2430CD
             816.
                      212.
                            0.878
                                   0.0974 ((30.5 -25, 30.41667 -25, 30.33333 -25, ~
##
                                           ((30.75 -24.75, 30.66667 -24.75, 30.6347~
##
   2 2430DA
              933.
                      179.
                            0.806
                                   0.106
                                           ((31 -24.75, 30.91667 -24.75, 30.83333 -~
##
   3 2430DB
              928.
                      315.
                            1.07
                                   0.117
   4 2430DC
              967.
                      210.
                            0.983
                                   0.0926 ((30.75 -25, 30.66667 -25, 30.58333 -25,~
                                           ((31 -25, 30.91667 -25, 30.83333 -25, 30~
##
   5 2430DD 1035.
                      268.
                            1.49
                                   0.113
##
   6 2525DB
              568.
                       31.3 0.402
                                   0.0762 ((26 -25.75, 25.91667 -25.75, 25.83333 -~
              542.
                       35.8 0.330 0.0749 ((25.75 -26, 25.66667 -26, 25.58333 -26,~
##
   7 2525DC
   8 2525DD
                       36.4 0.412 0.0741 ((26 -26, 25.91667 -26, 25.83333 -26, 25~
              579.
                       28.2 0.435 0.0636 ((26.25 -26, 26.16667 -26, 26.08333 -26,~
   9 2526CC
##
              618.
```

```
## 10 2526CD 647. 44.2 0.445 0.0667 ((26.5 -26, 26.41667 -26, 26.33333 -26, ~ ## # ... with 500 more rows
```

There are 510 different grid cells, each with its own associated covariate data (i.e. other columns with environmental values). You should also notice the geometry column, which includes the coordinates of each grid cell.

Using the flexible ggplot2 package, we will now make some figures First, plot the QDSs on top of the South Africa border to see where our data is found within the country. We simply specify geom_sf (a geometry feature, plotted in ggplot2 using the sf package extension). We tell each geom_sf layer which data to plot (sa and then qds_cov). The order is important. Each layer plots on top of the next. So in this case geom_sf(data = qds_cov) will plot on top of the geom_sf(data = sa) layer.

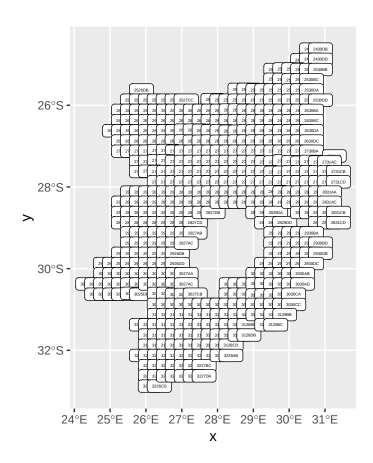
```
# Where are our grids inside SA?
ggplot() +
  geom_sf(data = sa) +
  geom_sf(data = qds_cov)
```



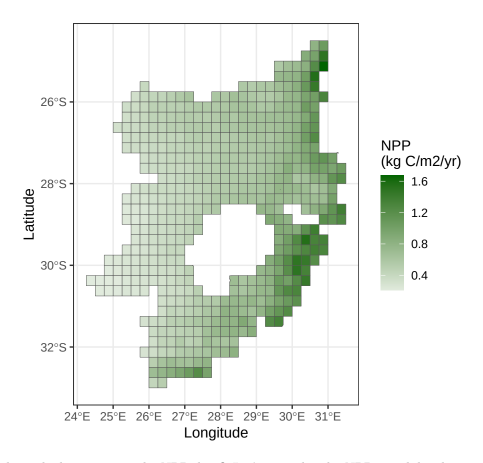
Each QDS is associated with a **unique identifying code**. The code represents the latitude and longitude of the grid and then which block this falls in to. There are 16 blocks within each combination of one degree of latitude and longitude, respectively. These are represented by letters. The top left block within each one degree is A. The smaller block within is AA. The bottom right block is DD and so on. Let's plot this to try and visualise it by labeling each QDS:

```
# What are the labels of our grids?
ggplot() +
```

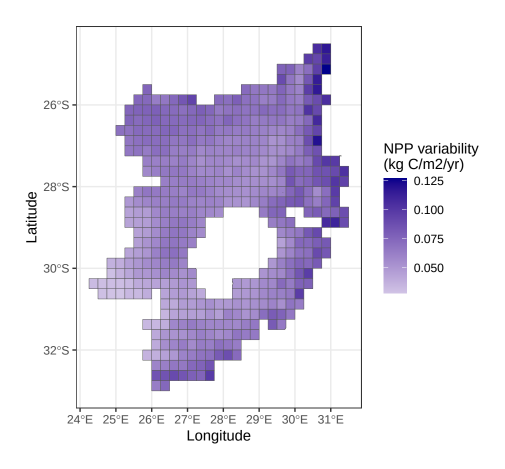
```
geom_sf(data = qds_cov) +
geom_sf_label(data = qds_cov, aes(label = QDS), size = 1)
```



We can now visualise our Net Primary Productivity (NPP) layer. We have the core of our code. But now we will add in a mapping aesthetic (or aes()). In this case we want each cell to be filled by the corresponding value of NPP. Then we can chose a colour gradient to fill this in with (white to darkgreen) and give our legend a good label (NPP (kg C/m2/yr)). We can also change the x and y labels using labs() and lastly edit the overall theme of the plot/map using a present function called $theme_bw()$.



What spatial trends do you see in the NPP data? Let's now plot the NPP variability layer. This uses the same copy and pasted code from above. We have just changed the fill variable in aes(), the colour gradient (white to darkblue) and the legend label. What spatial trends do you see in the NPP variability data?



3.3 Clean the data

The next step is to clean the data a bit further. In the POSA data, we only want one row per QDS. To do this we run the distinct() function. Next we select() only the columns we want. Lastly, take a look at each dataset.

```
#### Clean the data and join it together ----
# only select one row for each QDS
posa_rich %>% distinct(QDS, .keep_all = TRUE) -> posa_rich

# select the columns we want
posa_rich %>% select(QDS, spp_rich, func_diversity, phylog_diversity) -> posa_rich

# join the two datasets together
# reminder of each dataset
head(posa_rich)
```

```
## # A tibble: 6 x 4
##
             spp_rich func_diversity phylog_diversity
     QDS
##
     <chr>>
                <dbl>
                                <dbl>
                                                   <dbl>
                  731
## 1 3030CA
                                   24
                                                     139
## 2 2528CC
                  972
                                    24
                                                     139
                                   23
                                                     165
## 3 3226DB
                 1110
## 4 2731CB
                  449
                                   18
                                                     100
                                    24
## 5 2831CD
                  990
                                                     182
```

6 2930CB 1578 26 219

head(qds_cov)

```
## Simple feature collection with 6 features and 5 fields
## Geometry type: POLYGON
## Dimension:
                 XY
                 xmin: 25.75 ymin: -25.75 xmax: 31 ymax: -24.5
## Bounding box:
                 WGS 84
## Geodetic CRS:
## # A tibble: 6 x 6
##
     QDS
             MAP Elev_STD
                            NPP NPP_var
                                                                           geometry
     <chr> <dbl>
                     <dbl> <dbl>
                                  <dbl>
                                                                      <POLYGON [°]>
##
## 1 2430CD 816.
                     212. 0.878 0.0974 ((30.5 -25, 30.41667 -25, 30.33333 -25, 3~
## 2 2430DA 933.
                     179. 0.806 0.106 ((30.75 -24.75, 30.66667 -24.75, 30.63474~
                                  0.117 ((31 -24.75, 30.91667 -24.75, 30.83333 -2~
## 3 2430DB 928.
                     315. 1.07
                     210. 0.983 0.0926 ((30.75 -25, 30.66667 -25, 30.58333 -25, ~
## 4 2430DC 967.
                                 0.113 ((31 -25, 30.91667 -25, 30.83333 -25, 30.~
## 5 2430DD 1035.
                     268. 1.49
## 6 2525DB 568.
                     31.3 0.402 0.0762 ((26 -25.75, 25.91667 -25.75, 25.83333 -2~
```

3.3.1 Join the datasets

Both datasets have a column with the unique QDS identifier. We will use this to column to join them together with the left_join() function. Lastly, select() the columns we want to use and filter() out any columns that have no species richness data.

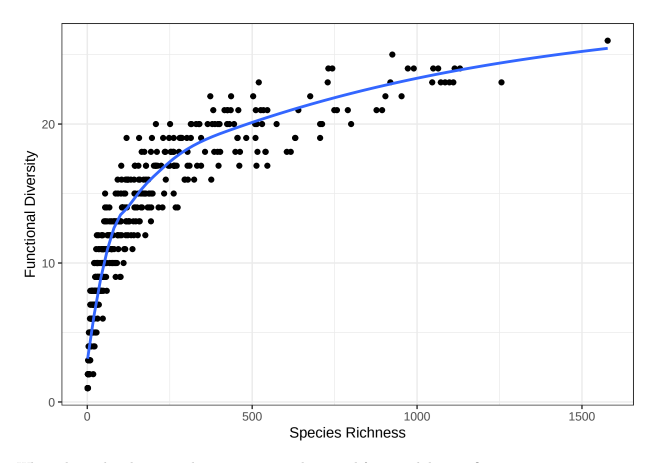
```
# join the two datasets based on the shared column 'QDS'
qds_cov %>% left_join(posa_rich, by = 'QDS') %>%
select(QDS:NPP_var, spp_rich:phylog_diversity, geometry) %>%
filter(!is.na(spp_rich)) -> joined_data
```

3.4 Analyse the data

After loading, exploring and cleaning, we can now begin to analyse the trends in the data. Let's make some figures to explore the relationships between the variables.

3.4.1 Species richness vs. functional diversity

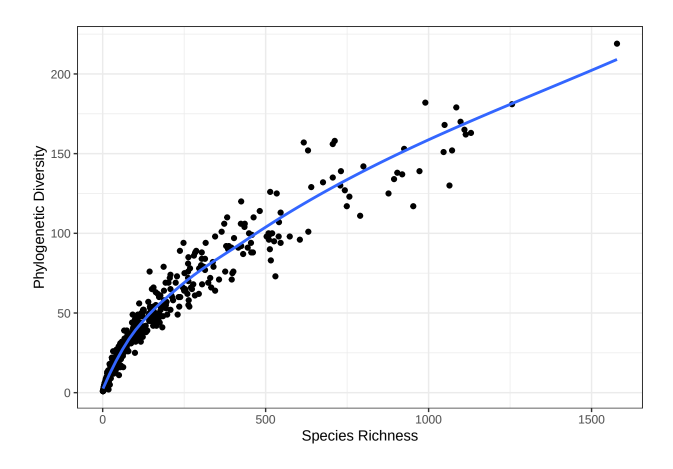
```
#### Analyse the data ----
# 1) Species richness vs. functional diversity
ggplot(data = joined_data, aes(x = spp_rich, y = func_diversity)) +
    geom_point() +
    geom_smooth(se = F) +
    labs(x = 'Species Richness', y = 'Functional Diversity') +
    theme_bw()
```



What relationship do you see between species richness and functional diversity?

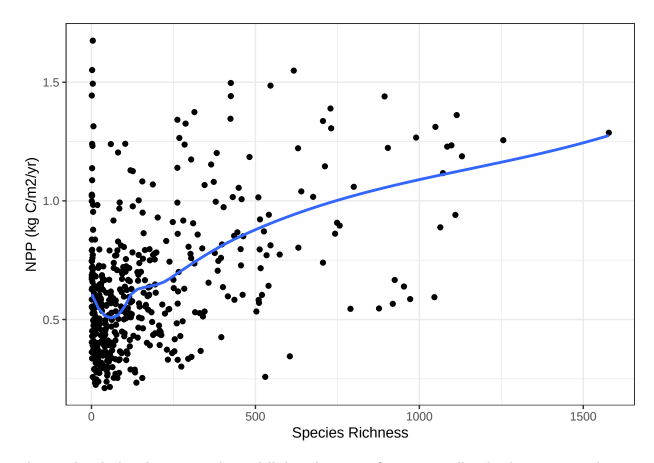
3.4.2 Species richness vs. phylogenetic diversity

```
# 2) Species richness vs. Phylogenetic Diversity
ggplot(data = joined_data, aes(x = spp_rich, y = phylog_diversity)) +
  geom_point() +
  geom_smooth(se = F) +
  labs(x = 'Species Richness', y = 'Phylogenetic Diversity') +
  theme_bw()
```



3.4.3 Species richness vs. NPP

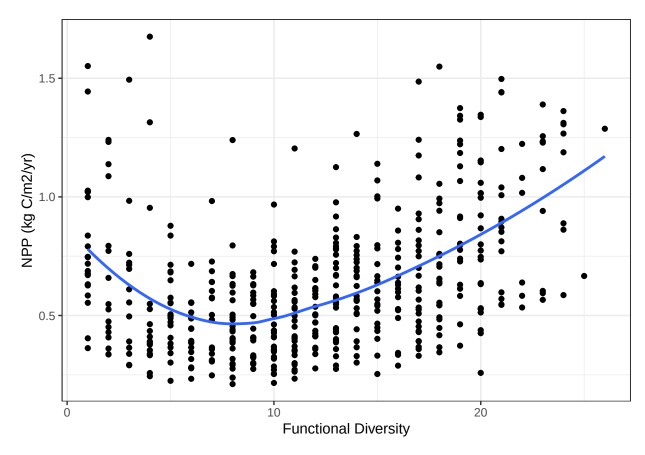
```
# 3) Species richness vs. NPP
ggplot(data = joined_data, aes(x = spp_rich, y = NPP)) +
  geom_point() +
  geom_smooth(se = F) +
  labs(x = 'Species Richness', y = 'NPP (kg C/m2/yr)') +
  theme_bw()
```



The raw data looks a bit messy. The modelled predictions in Section 2.5 will make these patterns clearer.

3.4.4 Functional diversity vs. NPP

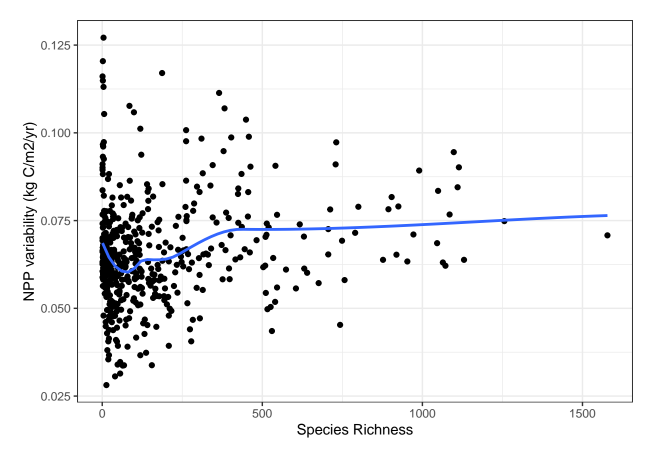
```
# 4) Functional Diversity vs. NPP
ggplot(data = joined_data, aes(x = func_diversity, y = NPP)) +
  geom_point() +
  geom_smooth(se = F) +
  labs(x = 'Functional Diversity', y = 'NPP (kg C/m2/yr)') +
  theme_bw()
```



Again, the raw data looks a bit messy. The modelled predictions in Section 2.5 will make these patterns clearer.

3.4.5 Species richness vs. NPP variability

```
# 5) Species richness vs. NPP variability
ggplot(data = joined_data, aes(x = spp_rich, y = NPP_var)) +
  geom_point() +
  geom_smooth(se = F) +
  labs(x = 'Species Richness', y = 'NPP variability (kg C/m2/yr)') +
  theme_bw()
```



There does not seem to be a clear trend in the species richness versus NPP variability plot. So we will stick to only analysing the relationships between NPP and 1) species richness and 2) functional diversity, for our linear models.

3.5 Run a linear model

To statistically analyse the relationships between our variables we will use linear models/regressions. These models will explain the relationships between our dependent (response) variables and our independent (predictor) variables. Think of these as a straight line equation:

$$y = mx + c \label{eq:y}$$
 In this equation,
$$y \label{eq:y}$$
 is your response variable and
$$x \label{eq:x}$$
 is predictor variable.
$$m$$

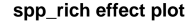
represents your slope or gradient. In R, it will be stored under the **coefficient** output using the summary function. c

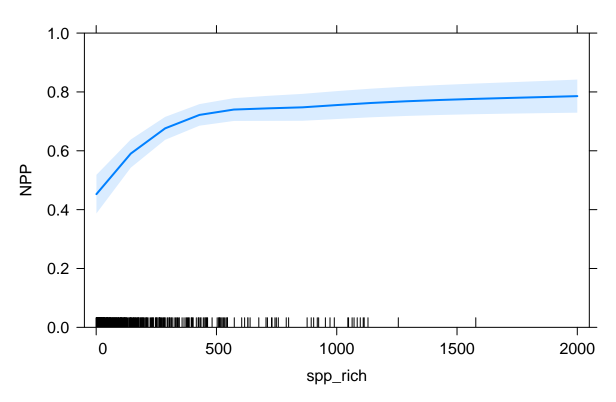
represents your intercept.

To run a linear model in R, we simply call the lm() function. Inside it, we need to specify our formula, which is NPP ~ log(spp_rich), and the dataset we are using: joined_data.

3.5.1 NPP versus species richness

```
#### run a linear model ----
# 1) What is the statistical relationship between NPP and species richness?
model1 <- lm(NPP ~ log(spp_rich), data = joined_data)</pre>
# look at regression coefficients
summary(model1)
##
## Call:
## lm(formula = NPP ~ log(spp_rich), data = joined_data)
##
## Residuals:
##
       Min
                1Q Median
                                 3Q
                                         Max
## -0.46929 -0.19989 -0.06139 0.12888 1.16177
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
               ## log(spp_rich) 0.043815 0.007428 5.899 6.93e-09 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.2795 on 480 degrees of freedom
## Multiple R-squared: 0.06759, Adjusted R-squared: 0.06564
## F-statistic: 34.79 on 1 and 480 DF, p-value: 6.929e-09
\# Plot the predicted relationship between NPP and species richness
plot(allEffects(model1), ylim = c(0,1))
```





Take a careful look at the summary() output and identify the important values, such as the **coefficient** estimates and the R^2 (R-squared).

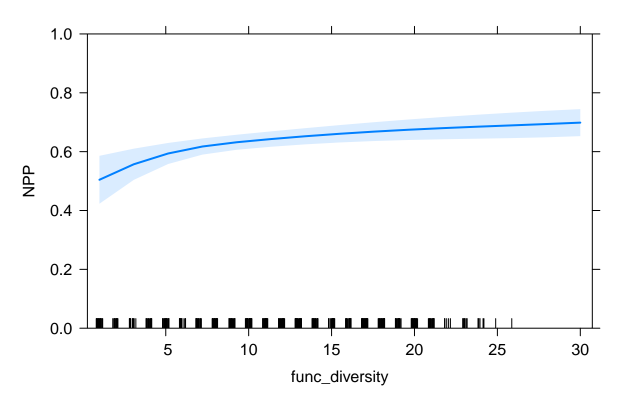
Lastly, we use the plot(allEffects()) nested functions to produce the predicted slope from our model. What is the **predicted** relationship between NPP and species richness?

3.5.2 NPP versus functional diversity

Now do the same thing for NPP and functional diversity.

```
# 2) What is the statistical relationship between NPP and functional diversity?
model2 <- lm(NPP ~ log(func_diversity), data = joined_data)
# look at regression coefficients
summary(model2)</pre>
```

func_diversity effect plot



What is the **predicted** relationship between NPP and functional diversity?

3.6 Linear model summaries

The outputs from summary() can look a bit messy. To tidy this up and provide a neater output, we will use the stargazer() function.

```
#### Summaries of linear models ----
# This function provides a neat summary of the regressions for both models
stargazer(model1, model2, type = 'text')
```

##

##		=========	==========	
##		Dependent variable:		
## ##		NPP		
##		(1)	(2)	
## ##	log(spp_rich)	0.044***		
##		(0.007)		
## ##	<pre>log(func_diversity)</pre>		0.057***	
##			(0.017)	
## ##	Constant	0.453***	0.504***	
##		(0.033)	(0.041)	
## ##				
##	Observations	482	482	
	R2 Adjusted R2	0.068 0.066	0.022 0.020	
##	Residual Std. Error (df = 480)		0.286	
	F Statistic (df = 1; 480)	34.793***	10.942***	
	Note:		.05; ***p<0.01	

These are the same values from our earlier outputs using summary(), but they are organised in a much better way for reading.

3.7 Submission

Your submission for this practical needs to be:

- Answer the 9 questions in the Practical 13 worksheet.
- Submit in word document (.docx) format.
- Submit by Monday 6 June 23h59.