Module "4": Assignment 1

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Assignment Details

Purpose

The goal of this assignment is to understand, apply, and interpret detection probability in estimating densities through distance sampling.

Task

Write R code to successfully answer each question below or write text to successfully answer the question.

Criteria for Success

- Code is within the provided code chunks
- Code chunks run without errors
- Code produces the correct result
 - Code attempts will get half points
 - Code that produces the correct answer will receive full points
- Text answers correctly address the question asked

Due Date

May 8 at midnight MST

Assignment Questions

All questions are worth 1 point unless otherwise specified.

Set-Up

We are going to need to load both the tidyverse and the community ecology package, vegan.

library(tidyverse)
library(vegan)

Species Accumulation Curves

For this section of the assignment, we are going to use a survey x species matrix for insects found on a plant.

```
insects <- read_csv("../data_raw/insects.csv")</pre>
```

```
## Rows: 56 Columns: 51
## -- Column specification ------
## Delimiter: ","
## dbl (51): cleta_eckloni, pseudambonea_capeni_schuhistes_lekkersingia, acanth...
##
## 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.
```

1. First, use the specaccum() function to prep the data for the species accumulation curve model.

```
insects_sac <- specaccum(insects)</pre>
```

2. Next, fit a species accumulation model to the data.

```
insects_sac_fit <- fitspecaccum(insects_sac, model = "asymp")</pre>
```

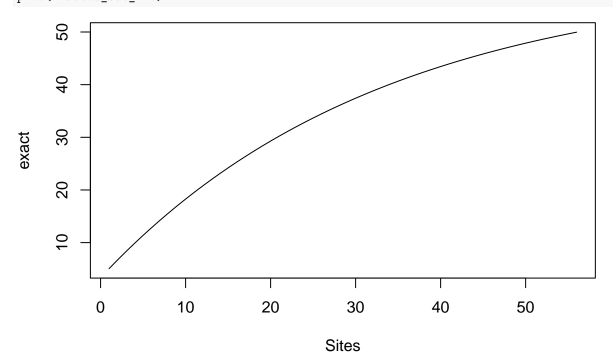
3. What is the predicted asymptote for this data?

```
coef(insects_sac_fit)
```

```
## Asym R0 lrc
## 60.445437 3.357250 -3.497493
```

4. Let's plot species accumulation curve model.

```
plot(insects_sac_fit)
```



5. Based on the predicted asymptote, the number of species we have in the dataset, and the plot you just made, what can we conclude? (2 points)

Answer:

Species Richness and Diversity Indices

For this assignment, you are welcome to use either base R or the tidyverse. If you haven't used the tidyverse much outside of this class, though, base R will probably be the more straightforward way to tackle this part of the assignment.

We are going to keep working with the insect data from above.

Species Richness

6. Use code to calculate the species richness of each community. Which community is more species rich?

```
nrow(comm1)
```

[1] 8

```
nrow(comm2)
```

[1] 5

Shannon Diversity Index We are going to start with comm1.

7. Write a line of code to calculate the number of individuals in the insect community. Use this to create the object N_comm1.

```
N_comm1 <- sum(comm1$individuals)</pre>
```

8. Add a column to the comm1 data frame with each species' proportion of the community.

```
comm1$prop <- comm1$individuals / N_comm1
```

9. Now add a column with the natural logs of the proportions.

```
comm1$ln_prop <- log(comm1$prop)</pre>
```

10. Add a column with the result of the proportion of the community multiplied by the natural log.

```
comm1$prop_lnprop <- comm1$prop * comm1$ln_prop</pre>
```

11. Calculate the Shannon Index for this community.

```
H_comm1 <- sum(comm1$prop_lnprop * -1)</pre>
```

12. Now that you've worked your way through calculating the Shannon Index for community 1, calculate it for community 2.

```
N_comm2 <- sum(comm2$individuals)
comm2$prop <- comm2$individuals / N_comm2
comm2$ln_prop <- log(comm2$prop)
comm2$prop_lnprop <- comm2$prop * comm2$ln_prop
H_comm2 <- sum(comm2$prop_lnprop * -1)</pre>
```

13. Which community is more diverse, according the the Shannon Diversity Index? How do you know? (2 points)

Answer:

Simpson Diversity Index Fortunately, a lot of the work has already been done to calculate the Simpson Diversity Index. We only need to add a little bit more to get there.

14. Add another column to the comm1 data frame that has the squared proportion.

```
comm1$prop_sq <- comm1$prop * comm1$prop</pre>
```

15. Calculate the Simpson Diversity Index for community 1

```
D1 = 1 - sum(comm1$prop_sq)
```

16. Now calculate the Simpson Index for community 2

```
comm2$prop_sq <- comm2$prop * comm2$prop
D2 = 1 - sum(comm2$prop_sq)</pre>
```

17. Do the values you've calculated for the Simpson Index corroborate your answer for which community is more diverse? Why or why not? (2 points)

Answer: