Richness & Diversity

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Comparison of richness and diversity between collection methods

We are interested to compare Malaise trap and Pollard walk surveys to see if there is a difference in the observed species richness, R, and diversity. For the latter, we will use Shannon's H as a measure of diversity (Hill 1973).

Methods

Using the BioSCAN data, we first calculate R (richness) and H (diversity) for each site, including only those sites for which data from both survey types is available. For the latter, we use the **vegan** package in R (Oksanen et al. 2018). We then compare the two methods through paired t-tests, to determine if the difference in means between collection methods is significant. We use the **ggplot2** package (Wickham 2009) for visualization. Additional data-wrangling provided by **tidyr** package.

Starting by loading dependencies and reading in the data

```
library("tidyr")  # converting data to long and wide format
library("ggplot2")  # plotting
library("vegan")  # calculating Shannon's H

## Loading required package: permute

## Loading required package: lattice

## This is vegan 2.5-2

source(file = "bioscan-functions.R")
bioscan <- CompleteBioscan()

# Identify those columns with species data
species.cols <- c(5:33)</pre>
```

Richness

Diversity

t-test

Richness

```
# Create data frame for richness t-test. One column for Pollard, one for Malaise
richness.df <- bioscan[, c("Site.Number", "Collection.Method", "Richness")]</pre>
richness.df <- richness.df %>%
  spread(Collection.Method, Richness)
\# Run the t-test
richness.t.test <- t.test(x = richness.df$Malaise,
                           y = richness.df$ Pollard Walk,
                          paired = TRUE)
# Store the value of t
richness.t <- round(richness.t.test$statistic, 3)
# Rounding may turn p-values into zero, so account for that possibility here
richness.p <- "< 0.001"
if (richness.t.test$p.value > 0.001) {
  richness.p <- round(richness.t.test$p.value, 3)</pre>
# Store the means
richness.means <- bioscan %>%
  group_by(Collection.Method) %>%
  summarise(means = round(mean(Richness), 3))
```

Diversity

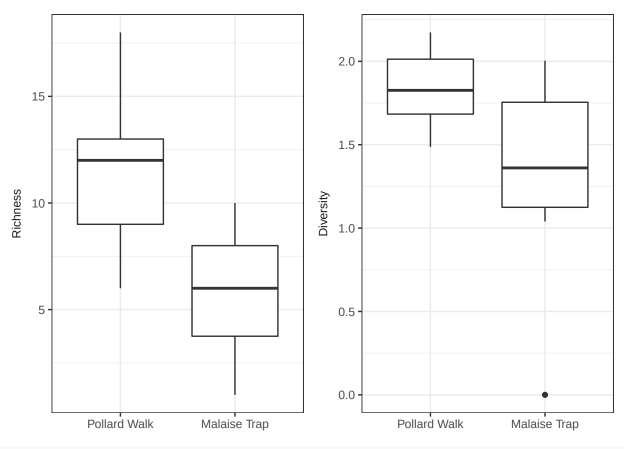
```
diversity.p <- round(diversity.t.test$p.value, 3)
}

# Store the means
diversity.means <- bioscan %>%
    group_by(Collection.Method) %>%
    summarise(means = round(mean(Diversity), 3))
```

Results

Observed istribution of richness and diversity:

```
# We need to create long-format data for use with ggplot
bioscan.long <- bioscan[, c("Collection.Method", "Richness", "Diversity")] %>%
  gather(key = "statistic",
         value = "value",
         -Collection.Method)
# For consistency, replace "Malaise" with "Malaise trap"
bioscan.long$Collection.Method <- gsub(pattern = "Malaise",</pre>
                                        replacement = "Malaise Trap",
                                        x = bioscan.long$Collection.Method)
# Re-level the two factors so they appear in desired order
bioscan.long Collection.Method <- factor(bioscan.long Collection.Method,
                                            levels = c("Pollard Walk", "Malaise Trap"))
bioscan.long$statistic <- factor(bioscan.long$statistic,</pre>
                                    levels = c("Richness", "Diversity"))
# Boxplot, with a separate plot for each statistic
statistics.plot <- ggplot(data = bioscan.long,</pre>
                          mapping = aes(x = Collection.Method, y = value)) +
  geom_boxplot() +
 facet_wrap(~ statistic, scales = "free_y", strip.position = "left") +
  theme_bw() +
 ylab(NULL) +
  xlab(NULL) +
  theme(strip.background = element_blank(),
        strip.placement = "outside")
print(statistics.plot)
```



ggsave(filename = "output/richness-diversity-boxplot.png", plot = statistics.plot)

Saving 6.5 x 4.5 in image

Means table:

Collection Method	R	Н
Malaise trap	5.938	1.393
Pollard walk	11.312	1.846

t-table:

Statistic	t	p
\overline{R}	-4.657	< 0.001
H	-3.307	0.005