

# Rock Lichen data from Sunset Crater

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## Analysis Summary

- Dead trees and non-lichen species were removed from lichen community analyses.
- Lichen communities were adequately sampled, based on species accumulation curves, with moth resistant trees accumulating slightly more lichen species.
- Lichen communities (abundance, richness, diversity, composition) were significantly, generally negatively, affected by moth susceptibility.
- Several tree variables, including light availability, leaf litter abundance and rock abundance, were impacted by moth susceptibility.
- Analysis of causal pathways supported an indirect link between moth susceptibility and impacts on lichen communities via decreasing rock (i.e. habitat) availability through increased leaf abscission and accumulation on rocks under trees.
- These results support a genetically based link between intraspecific variation in susceptibility to an insect herbivore and community dynamics in an arid ecosystem.
- Given the possible impacts of climate change on this system, this study supports the conclusion that community and ecosystem impacts need to be considered in an evolutionary context.

```
# 0. Supporting functions and libraries
## Support functions

dif <- function(x){
  out=x[1]
  for (i in 2:length(x)){
    out=out-x[i]
  }
  return(out)
}

## Libraries
my.libs <- c("vegan", "ecodist")
if (any(!(my.libs %in% installed.packages()[, 1]))){
  sapply(my.libs[!(my.libs %in% installed.packages()[, 1])],
    install.packages)
}else{}
sapply(my.libs, require, character.only = TRUE)
```

## Load Data

The following are variable descriptions (Variable, Type, Range, Definition):

- Moth,categorical,0 or 1,Was the tree susceptible (0) or resistant (1) to moth attack?
- Live/Dead,categorical,0 or 1,Was the tree dead (0) or alive (1)?

- Litter %,continuous,0 to 100,Percent cover inside quadrat
- Rocks > 3cm? %,continuous,0 to 100,Percent cover of rocks > 3cm? inside quadrat
- Rocks < 3cm? %,continuous,0 to 100,Percent cover of rocks < 3cm? inside quadrat
- Shrubs %,continuous,0 to 100,Percent cover of shrubs inside quadrat
- Grass %,continuous,0 to 100,Percent cover of grass inside quadrat
- Branches %,continuous,0 to 100,Percent cover of branches on ground inside quadrat
- Distance,continuous,0 to 100,“Distance from main trunk, converted to percent of crown radius at that azimuth”
- Azimuth,continuous,0 to 360,Compass direction from main trunk
- Slope,continuous,0 to 90,Topographical steepness
- Aspect,continuous,0 to 360,Compass direction of slope
- Light,continuous,,Amount of light available to epiliths

```
## Data are in ../data/scrl
l.dat <- read.csv("../data/scrl/spp_env_combined.csv")

## Summary of data
summary(l.dat)

## remove dead trees
l.dat <- l.dat[l.dat[, "Live.Dead"] != 0, ]

## Lichen species list
spp.l <- c("Acacon", "Acasup", "Acaobp", "Sterile.sp", "Brown.cr",
"Lobalp", "Canros", "Calare", "Phydub", "Rhichr", "Xanlin", "Xanpli",
"Xanele", "GrBr.cr", "Gray.cr")
spp.moss <- c("Synrur", "Cerpur.Bryarg")

## Create a community matrix
com <- l.dat[, colnames(l.dat) %in% spp.l]
com.moss <- l.dat[, colnames(l.dat) %in% spp.moss]

## Add the tree labels to the rownames
rownames(com) <- paste(l.dat[, "Moth"], l.dat[, "Tree.pairs"], sep = "_")
rownames(com.moss) <- paste(l.dat[, "Moth"], l.dat[, "Tree.pairs"], sep = "_")
rownames(l.dat) <- paste(l.dat[, "Moth"], l.dat[, "Tree.pairs"], sep = "_")
```

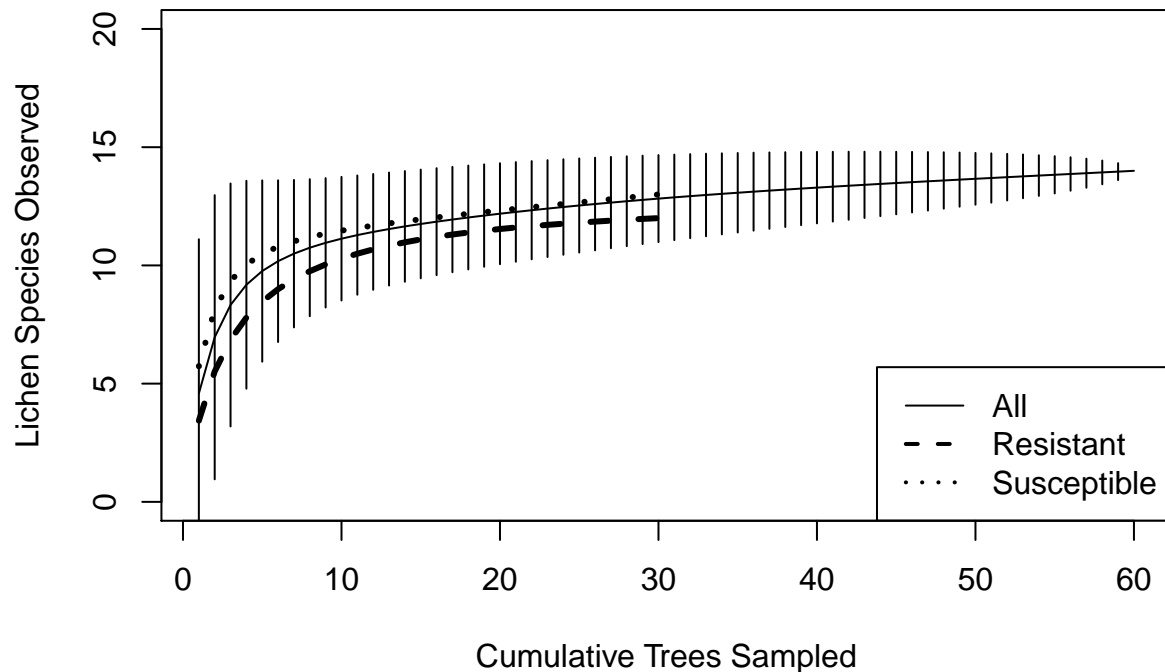
## Species accumulation

Are the communities on each tree type adequately sampled?

```
spa.all <- specaccum(com)
spa.res <- specaccum(com[l.dat[, "Moth"] == 0, ])
spa.sus <- specaccum(com[l.dat[, "Moth"] == 1, ])

plot(spa.all,
     ylim = c(0, 20),
     xlab = "Cumulative Trees Sampled",
     ylab = "Lichen Species Observed")
lines(spa.res$sites, spa.res$richness,
      ylim = c(0, 20), lty = 2, lwd = 3)
lines(spa.sus$sites, spa.sus$richness,
      ylim = c(0, 20), lty = 3, lwd = 3)
```

```
legend("bottomright",
      legend = c("All", "Resistant", "Susceptible"),
      lty = c(1, 2, 3), lwd = c(1, 2, 2))
```



## Moth trees have different microenvironments

- paired t-tests

## Moth trees have different lichen communities (FIGURE ch.plot A, R, H)

less abundant and diverse (paired t-tests, in text)

```
abun <- apply(com, 1, sum)
rich <- apply(com, 1, function(x) sum(sign(x)))
shan <- apply(com, 1, diversity, index = "shannon")
tt.a <- t.test(tapply(abun, l.dat[, "Tree.pairs"], diff))
tt.r <- t.test(tapply(rich, l.dat[, "Tree.pairs"], diff))
tt.h <- t.test(tapply(shan, l.dat[, "Tree.pairs"], diff))
tt.arh <- do.call(rbind,
                  list(a = unlist(tt.a), r = unlist(tt.r), h = unlist(tt.h)))
data.frame(tt.arh)
```

##	statistic.t	parameter.df	p.value	conf.int1
## a	-2.35680534636893	29	0.0253991007560338	-2.89259563276878
## r	-2.83579994251995	29	0.00824742800912123	-3.95880113980294
## h	-2.43278934693583	29	0.0213834528180339	-0.783514237345595
##	conf.int2	estimate.mean.of.x	null.value.mean	stderr
## a	-0.204737700564558	-1.54866666666667	0	0.657104189386137

```
## r -0.641198860197056          -2.3          0 0.811058624239964
## h -0.0678109108683953 -0.425662574106995    0 0.174968940341312
## alternative method data.name
## a two.sided One Sample t-test tapply(abun, l.dat[, "Tree.pairs"], diff)
## r two.sided One Sample t-test tapply(rich, l.dat[, "Tree.pairs"], diff)
## h two.sided One Sample t-test tapply(shan, l.dat[, "Tree.pairs"], diff)
```

composition is different (PERMANOVA, in text and supplement)

```
com.ds <- cbind(com, ds = rep(0.0001, nrow(com)))
set.seed(123)
adonis2(com.ds ~ Moth, data = l.dat,
        strata = l.dat[, "Tree.pairs"],
        by = "margin", nperm = 100000)
```

```
## Permutation test for adonis under NA model
## Marginal effects of terms
## Permutation: free
## Number of permutations: 999
##
## adonis2(formula = com.ds ~ Moth, data = l.dat, by = "margin", strata = l.dat[, "Tree.pairs"], nperm =
##          Df SumOfSqs      R2      F Pr(>F)
## Moth      1   0.8281 0.03894 2.3499 0.023 *
## Residual 58  20.4394 0.96106
## Total    59  21.2676 1.00000
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

three main species were reduced by moths (FDR paired t-tests, in text + supplement)

```
ind.spp <- lapply(com, function(x, p) t.test(tapply(x, p, diff)), p = l.dat[, "Tree.pairs"])
isp <- apply(do.call(rbind, lapply(ind.spp, unlist)), 2, as.numeric)
```

```
## Warning in apply(do.call(rbind, lapply(ind.spp, unlist)), 2, as.numeric): NAs
## introduced by coercion
```

```
## Warning in apply(do.call(rbind, lapply(ind.spp, unlist)), 2, as.numeric): NAs
## introduced by coercion
```

```
## Warning in apply(do.call(rbind, lapply(ind.spp, unlist)), 2, as.numeric): NAs
## introduced by coercion
```

```
rownames(isp) <- names(ind.spp)
isp[, "p.value"] <- p.adjust(isp[, "p.value"], method = "fdr")
isp <- isp[order(isp[, "p.value"]), ]
head(isp[, 1:3])
```

```
##          statistic.t parameter.df      p.value
## Acacon    -3.377629           29 0.01390405
## Acasup    -3.242091           29 0.01390405
## Canros    -3.581884           29 0.01390405
## Lobalp    -2.041361           29 0.17642430
## Phydub    -1.922619           29 0.18031798
## Calare    -1.607607           29 0.22424946
```

## Litter covering rocks was the main driver (FIGURE = ORDINATION)

light not litter predicted lichen composition (PERMANOVA, table 3, Ordination)

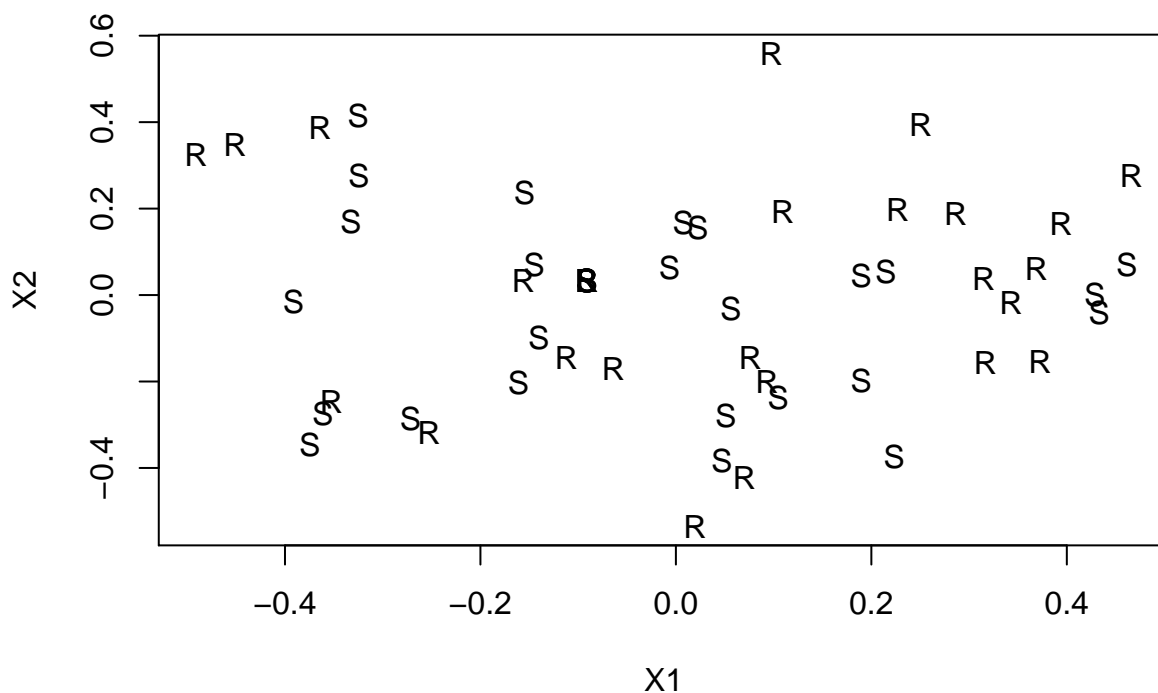
```
set.seed(123)
adonis2(com.ds ~ Light...average + Litter..., data = l.dat,
        strata = l.dat[, "Tree.pairs"],
        by = "margin", nperm = 100000)

## Permutation test for adonis under reduced model
## Marginal effects of terms
## Permutation: free
## Number of permutations: 999
##
## adonis2(formula = com.ds ~ Light...average + Litter..., data = l.dat, by = "margin", strata = l.dat[,
##          Df SumOfSqs      R2      F Pr(>F)
## Light...average  1   0.4115 0.01935 1.2257  0.240
## Litter...       1   1.0096 0.04747 3.0072  0.007 **
## Residual       57  19.1362 0.89979
## Total          59  21.2676 1.00000
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

nm.ds.out <- nm.ds(vegdist(com.ds), 5, 5)
ord <- nm.ds.min(nm.ds.out, dims = 5)

## Minimum stress for given dimensionality: 0.08730395
## r^2 for minimum stress configuration: 0.9159319

ord.pch <- c("R", "S")[(l.dat[, "Moth"] + 1)]
plot(X2~ X1, data = ord, pch = ord.pch)
```



litter not light was correlated with large rocks (dist cor, in text)

```
cor.test(tapply(l.dat[, "Big.rocks.."], l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Litter.."], l.dat[, "Tree.pairs"], diff))
```

```
##
## Pearson's product-moment correlation
##
## data:  tapply(l.dat[, "Big.rocks.."], l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Litter.."], l.dat[, "Tree.pairs"], diff)
## t = -11.106, df = 28, p-value = 9.054e-12
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
##  -0.9530598 -0.8039735
## sample estimates:
##          cor
## -0.9027609
```

```
cor.test(tapply(l.dat[, "Big.rocks.."], l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Light...average"], l.dat[, "Tree.pairs"], diff))
```

```
##
## Pearson's product-moment correlation
##
## data:  tapply(l.dat[, "Big.rocks.."], l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Light...average"], l.dat[, "Tree.pairs"], diff)
## t = 0.71624, df = 28, p-value = 0.4798
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
##  -0.2376184  0.4716125
## sample estimates:
##          cor
##  0.1341335
```

## Community Analyses

The following is an analysis of the univariate community metrics, abundance, richness and diversity (Shannon's index). These three metrics were strongly intercorrelated and all responded to the environmental effects of moth infection. Litter cover, light and rock abundance were all also responsive to moth infection. Other variables, such as branches, grasses and mosses were not impacted by moth. Differences are analyzed with univariate t-tests in order to account for the paired structure of the data. Also presented here are Shapiro-Wilks tests for homogeneity of variance, which as assumption of the t-test. These effects should be shown using dotplots with mean differences as dots and confidence intervals as horizontal lines.

```
## Calculate metrics
abun <- apply(com, 1, sum)
rich <- apply(com, 1, function(x) sum(sign(x)))
shan <- apply(com, 1, diversity, index = "shannon")
```

```
## Examine intercorrelations
cor.test(abun, rich, method = "k")
```

```
##
## Kendall's rank correlation tau
##
## data:  abun and rich
## z = 8.3541, p-value < 2.2e-16
```

```

## alternative hypothesis: true tau is not equal to 0
## sample estimates:
##      tau
## 0.7845321
cor.test(abun, shan, method = "k")

##
## Kendall's rank correlation tau
##
## data:  abun and shan
## z = 5.9531, p-value = 2.63e-09
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
##      tau
## 0.5497907
cor.test(rich, shan, method = "k")

##
## Kendall's rank correlation tau
##
## data:  rich and shan
## z = 7.8403, p-value = 4.496e-15
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
##      tau
## 0.7471965

## Test for moth effects via a one-way test on the
## differences between susceptible and resistance trees.
## diff = Susceptible - Resistant
## If diff is less than 0, susceptibility had a negative effect
## If diff is greater than 0, susceptibility had a positive effect

### Test for violation of normality of differences
shapiro.test(tapply(abun , l.dat[, "Tree.pairs"], diff))

##
## Shapiro-Wilk normality test
##
## data:  tapply(abun, l.dat[, "Tree.pairs"], diff)
## W = 0.95143, p-value = 0.1847
shapiro.test(tapply(rich , l.dat[, "Tree.pairs"], diff))

##
## Shapiro-Wilk normality test
##
## data:  tapply(rich, l.dat[, "Tree.pairs"], diff)
## W = 0.9541, p-value = 0.2175
shapiro.test(tapply(shan , l.dat[, "Tree.pairs"], diff))

##
## Shapiro-Wilk normality test
##
## data:  tapply(shan, l.dat[, "Tree.pairs"], diff)

```

```

## W = 0.95291, p-value = 0.2021
shapiro.test(tapply(l.dat[, "Litter.."] , l.dat[, "Tree.pairs"], diff))

##
## Shapiro-Wilk normality test
##
## data:  tapply(l.dat[, "Litter.."], l.dat[, "Tree.pairs"], diff)
## W = 0.94289, p-value = 0.1088
shapiro.test(tapply(l.dat[, "Light...average"] , l.dat[, "Tree.pairs"], diff))

##
## Shapiro-Wilk normality test
##
## data:  tapply(l.dat[, "Light...average"], l.dat[, "Tree.pairs"], diff)
## W = 0.96945, p-value = 0.5242
shapiro.test(tapply(l.dat[, "Light...N"] , l.dat[, "Tree.pairs"], diff))

##
## Shapiro-Wilk normality test
##
## data:  tapply(l.dat[, "Light...N"], l.dat[, "Tree.pairs"], diff)
## W = 0.98758, p-value = 0.9727
shapiro.test(tapply(l.dat[, "Light...S"] , l.dat[, "Tree.pairs"], diff))

##
## Shapiro-Wilk normality test
##
## data:  tapply(l.dat[, "Light...S"], l.dat[, "Tree.pairs"], diff)
## W = 0.98356, p-value = 0.9102
shapiro.test(tapply(l.dat[, "Big.rocks.." ] , l.dat[, "Tree.pairs"], diff))

##
## Shapiro-Wilk normality test
##
## data:  tapply(l.dat[, "Big.rocks.."], l.dat[, "Tree.pairs"], diff)
## W = 0.94775, p-value = 0.1471
### Too small for habitat
shapiro.test(tapply(l.dat[, "Small.rocks.." ] , l.dat[, "Tree.pairs"], diff))

##
## Shapiro-Wilk normality test
##
## data:  tapply(l.dat[, "Small.rocks.."], l.dat[, "Tree.pairs"], diff)
## W = 0.79098, p-value = 4.509e-05
### Too few observations
shapiro.test(tapply(l.dat[, "Synrur"] , l.dat[, "Tree.pairs"], diff))

##
## Shapiro-Wilk normality test
##
## data:  tapply(l.dat[, "Synrur"], l.dat[, "Tree.pairs"], diff)
## W = 0.3766, p-value = 3.307e-10

```



```

shapiro.test(tapply(l.dat[, "Branches.."] , l.dat[, "Tree.pairs"], diff))

##
## Shapiro-Wilk normality test
##
## data:  tapply(l.dat[, "Branches.."], l.dat[, "Tree.pairs"], diff)
## W = 0.17962, p-value = 7.766e-12

shapiro.test(tapply(l.dat[, "Grass.."] , l.dat[, "Tree.pairs"], diff))

##
## Shapiro-Wilk normality test
##
## data:  tapply(l.dat[, "Grass.."], l.dat[, "Tree.pairs"], diff)
## W = 0.17962, p-value = 7.766e-12

shapiro.test(tapply(l.dat[, "Cerpur.Bryarg"] , l.dat[, "Tree.pairs"], diff))

##
## Shapiro-Wilk normality test
##
## data:  tapply(l.dat[, "Cerpur.Bryarg"], l.dat[, "Tree.pairs"], diff)
## W = 0.2428, p-value = 2.398e-11

#### Test for effect of moth
t.test(tapply(abun, l.dat[, "Tree.pairs"], diff))

##
## One Sample t-test
##
## data:  tapply(abun, l.dat[, "Tree.pairs"], diff)
## t = -2.3568, df = 29, p-value = 0.0254
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  -2.8925956 -0.2047377
## sample estimates:
## mean of x
## -1.548667

t.test(tapply(rich, l.dat[, "Tree.pairs"], diff))

##
## One Sample t-test
##
## data:  tapply(rich, l.dat[, "Tree.pairs"], diff)
## t = -2.8358, df = 29, p-value = 0.008247
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  -3.9588011 -0.6411989
## sample estimates:
## mean of x
##      -2.3

t.test(tapply(shan, l.dat[, "Tree.pairs"], diff))

##
## One Sample t-test

```

```

##
## data:  tapply(shan, l.dat[, "Tree.pairs"], diff)
## t = -2.4328, df = 29, p-value = 0.02138
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  -0.78351424 -0.06781091
## sample estimates:
##  mean of x
## -0.4256626
t.test(tapply(l.dat[, "Litter.."] , l.dat[, "Tree.pairs"], diff))

##
## One Sample t-test
##
## data:  tapply(l.dat[, "Litter.."], l.dat[, "Tree.pairs"], diff)
## t = 2.8665, df = 29, p-value = 0.00765
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  4.317792 25.822208
## sample estimates:
## mean of x
## 15.07
t.test(tapply(l.dat[, "Light...average"] , l.dat[, "Tree.pairs"], diff))

##
## One Sample t-test
##
## data:  tapply(l.dat[, "Light...average"], l.dat[, "Tree.pairs"], diff)
## t = -9.2728, df = 29, p-value = 3.557e-10
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  -18.47119 -11.79547
## sample estimates:
## mean of x
## -15.13333
t.test(tapply(l.dat[, "Light...N"] , l.dat[, "Tree.pairs"], diff))

##
## One Sample t-test
##
## data:  tapply(l.dat[, "Light...N"], l.dat[, "Tree.pairs"], diff)
## t = -8.0191, df = 29, p-value = 7.634e-09
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  -20.05142 -11.90191
## sample estimates:
## mean of x
## -15.97667
t.test(tapply(l.dat[, "Light...S"] , l.dat[, "Tree.pairs"], diff))

##
## One Sample t-test
##

```

```

## data:  tapply(l.dat[, "Light...S"], l.dat[, "Tree.pairs"], diff)
## t = -7.5187, df = 29, p-value = 2.748e-08
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  -18.17717 -10.40283
## sample estimates:
## mean of x
##    -14.29

t.test(tapply(l.dat[, "Big.rocks.."] , l.dat[, "Tree.pairs"], diff))

##
## One Sample t-test
##
## data:  tapply(l.dat[, "Big.rocks.."], l.dat[, "Tree.pairs"], diff)
## t = -2.4617, df = 29, p-value = 0.02001
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  -17.728936 -1.638397
## sample estimates:
## mean of x
## -9.683667

                                # Ignore these results
t.test(tapply(l.dat[, "Small.rocks.."] , l.dat[, "Tree.pairs"], diff))

##
## One Sample t-test
##
## data:  tapply(l.dat[, "Small.rocks.."], l.dat[, "Tree.pairs"], diff)
## t = -2.0792, df = 29, p-value = 0.04655
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  -9.86878788 -0.08121212
## sample estimates:
## mean of x
##    -4.975

t.test(tapply(l.dat[, "Branches.."] , l.dat[, "Tree.pairs"], diff))

##
## One Sample t-test
##
## data:  tapply(l.dat[, "Branches.."], l.dat[, "Tree.pairs"], diff)
## t = 1, df = 29, p-value = 0.3256
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  -0.1484226 0.4324226
## sample estimates:
## mean of x
##    0.142

t.test(tapply(l.dat[, "Grass.."] , l.dat[, "Tree.pairs"], diff))

##
## One Sample t-test
##

```

```
## data:  tapply(l.dat[, "Grass.."], l.dat[, "Tree.pairs"], diff)
## t = -1, df = 29, p-value = 0.3256
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  -0.15023133  0.05156466
## sample estimates:
##  mean of x
## -0.04933333
```

```
t.test(tapply(l.dat[, "Synrur"] , l.dat[, "Tree.pairs"], diff))
```

```
##
##  One Sample t-test
##
## data:  tapply(l.dat[, "Synrur"], l.dat[, "Tree.pairs"], diff)
## t = 0.36285, df = 29, p-value = 0.7194
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  -0.1020054  0.1460054
## sample estimates:
## mean of x
##  0.022
```

```
t.test(tapply(l.dat[, "Cerpur.Bryarg"] , l.dat[, "Tree.pairs"], diff))
```

```
##
##  One Sample t-test
##
## data:  tapply(l.dat[, "Cerpur.Bryarg"], l.dat[, "Tree.pairs"], diff)
## t = -1.2357, df = 29, p-value = 0.2265
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  -0.04602247  0.01135580
## sample estimates:
##  mean of x
## -0.01733333
```

Lichen communities as a whole responded to the effects of moth infection. Using PerMANOVA, a permutational analysis of variance, with the tree pairs as blocks in the model, we observed a significant effect of moth susceptibility on lichen community composition. This should be shown using an NMDS ordination plot.

```
#### Composition analysis
com.ds <- cbind(com, ds = rep(0.01, nrow(com)))
adonis2(com.ds ~ Moth, data = l.dat, strata = factor(l.dat$Tree.pairs), perm = 9999)
```

```
## Permutation test for adonis under reduced model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 9999
##
## adonis2(formula = com.ds ~ Moth, data = l.dat, permutations = 9999, strata = factor(l.dat$Tree.pairs))
##           Df SumOfSqs      R2      F Pr(>F)
## Moth       1   0.8849 0.04412 2.6772 0.0224 *
## Residual  58  19.1700 0.95588
## Total     59  20.0548 1.00000
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```

set.seed(159)
adonis2(com.ds ~ Litter.. + Light...average , data = l.dat,
        strata = factor(l.dat$Tree.pairs), perm = 10000, by = "margin")

## Permutation test for adonis under reduced model
## Marginal effects of terms
## Permutation: free
## Number of permutations: 10000
##
## adonis2(formula = com.ds ~ Litter.. + Light...average, data = l.dat, permutations = 10000, by = "margin")
##              Df SumOfSqs      R2      F Pr(>F)
## Litter..      1   1.2076 0.06021 3.9089 0.0025 **
## Light...average 1   0.4195 0.02092 1.3579 0.2055
## Residual     57  17.6095 0.87807
## Total        59  20.0548 1.00000
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

set.seed(159)
adonis2(com.ds ~ Light...average + Litter.. , data = l.dat,
        strata = factor(l.dat$Tree.pairs), perm = 10000, by = "margin")

## Permutation test for adonis under reduced model
## Marginal effects of terms
## Permutation: free
## Number of permutations: 10000
##
## adonis2(formula = com.ds ~ Light...average + Litter.., data = l.dat, permutations = 10000, by = "margin")
##              Df SumOfSqs      R2      F Pr(>F)
## Light...average 1   0.4195 0.02092 1.3579 0.2055
## Litter..        1   1.2076 0.06021 3.9089 0.0025 **
## Residual       57  17.6095 0.87807
## Total          59  20.0548 1.00000
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

cor.test(tapply(l.dat[, "Litter.."] , l.dat[, "Tree.pairs"], diff),
        tapply(l.dat[, "Big.rocks.." ] , l.dat[, "Tree.pairs"], diff))

##
## Pearson's product-moment correlation
##
## data:  tapply(l.dat[, "Litter.."], l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Big.rocks.."], l.dat[, "Tree.pairs"], diff)
## t = -11.106, df = 28, p-value = 9.054e-12
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
##  -0.9530598 -0.8039735
## sample estimates:
##          cor
## -0.9027609

cor.test(tapply(l.dat[, "Light...average"] , l.dat[, "Tree.pairs"], diff),
        tapply(l.dat[, "Big.rocks.." ] , l.dat[, "Tree.pairs"], diff))

##
## Pearson's product-moment correlation

```

```
##
## data:  tapply(l.dat[, "Light...average"], l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Big.rock",
## t = 0.71624, df = 28, p-value = 0.4798
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
##  -0.2376184  0.4716125
## sample estimates:
##      cor
## 0.1341335

cor.test(tapply(l.dat[, "Light...average"] , l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Small.rock", l.dat[, "Tree.pairs"], diff))

##
## Pearson's product-moment correlation
##
## data:  tapply(l.dat[, "Light...average"], l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Small.rock",
## t = 1.1133, df = 28, p-value = 0.275
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
##  -0.1667471  0.5270644
## sample estimates:
##      cor
## 0.2058908
```

## Indicator species

Although susceptibility to moth infection generally had a negative impact on lichen, several species showed significant responses. Here, because of the paired design, we analyze the response of the differences using a one-way t-test on the differences of each species. I don't think we need to plot this, but perhaps the results could be shown as a table.

```
ind.spp <- list()
ind.spp$p.value[1] <- t.test(tapply(com[, "Acacon"], l.dat[, "Tree.pairs"], diff))$p.value
ind.spp$p.value[2] <- t.test(tapply(com[, "Acasup"], l.dat[, "Tree.pairs"], diff))$p.value
ind.spp$p.value[3] <- t.test(tapply(com[, "Canros"], l.dat[, "Tree.pairs"], diff))$p.value
ind.spp$p.value[4] <- t.test(tapply(com[, "Lobalp"], l.dat[, "Tree.pairs"], diff))$p.value
ind.spp$p.value[5] <- t.test(tapply(com[, "Phydub"], l.dat[, "Tree.pairs"], diff))$p.value
ind.spp$p.value[6] <- t.test(tapply(com[, "Acaobp"], l.dat[, "Tree.pairs"], diff))$p.value
ind.spp$p.value[7] <- t.test(tapply(com[, "Sterile.sp"], l.dat[, "Tree.pairs"], diff))$p.value
ind.spp$p.value[8] <- t.test(tapply(com[, "Calare"], l.dat[, "Tree.pairs"], diff))$p.value
ind.spp$p.value[9] <- t.test(tapply(com[, "Rhichr"], l.dat[, "Tree.pairs"], diff))$p.value
ind.spp$p.value[10] <- t.test(tapply(com[, "Xanlin"], l.dat[, "Tree.pairs"], diff))$p.value
ind.spp$p.value[11] <- t.test(tapply(com[, "Xanpli"], l.dat[, "Tree.pairs"], diff))$p.value
ind.spp$p.value[12] <- t.test(tapply(com[, "Xanele"], l.dat[, "Tree.pairs"], diff))$p.value
ind.spp$p.value[13] <- t.test(tapply(com[, "GrBr.cr"], l.dat[, "Tree.pairs"], diff))$p.value
ind.spp$t[1] <- t.test(tapply(com[, "Acacon"], l.dat[, "Tree.pairs"], diff))$statistic
ind.spp$t[2] <- t.test(tapply(com[, "Acasup"], l.dat[, "Tree.pairs"], diff))$statistic
ind.spp$t[3] <- t.test(tapply(com[, "Canros"], l.dat[, "Tree.pairs"], diff))$statistic
ind.spp$t[4] <- t.test(tapply(com[, "Lobalp"], l.dat[, "Tree.pairs"], diff))$statistic
ind.spp$t[5] <- t.test(tapply(com[, "Phydub"], l.dat[, "Tree.pairs"], diff))$statistic
ind.spp$t[6] <- t.test(tapply(com[, "Acaobp"], l.dat[, "Tree.pairs"], diff))$statistic
ind.spp$t[7] <- t.test(tapply(com[, "Sterile.sp"], l.dat[, "Tree.pairs"], diff))$statistic
ind.spp$t[8] <- t.test(tapply(com[, "Calare"], l.dat[, "Tree.pairs"], diff))$statistic
```

```
ind.spp$t[9] <- t.test(tapply(com[, "Rhichr"], l.dat[, "Tree.pairs"], diff))$statistic
ind.spp$t[10] <- t.test(tapply(com[, "Xanlin"], l.dat[, "Tree.pairs"], diff))$statistic
ind.spp$t[11] <- t.test(tapply(com[, "Xanpli"], l.dat[, "Tree.pairs"], diff))$statistic
ind.spp$t[12] <- t.test(tapply(com[, "Xanele"], l.dat[, "Tree.pairs"], diff))$statistic
ind.spp$t[13] <- t.test(tapply(com[, "GrBr.cr"], l.dat[, "Tree.pairs"], diff))$statistic
## This species had zero abundance, perhaps it was only observed under dead trees
t.test(tapply(com[, "Brown.cr"], l.dat[, "Tree.pairs"], diff))
```

```
##
## One Sample t-test
##
## data: tapply(com[, "Brown.cr"], l.dat[, "Tree.pairs"], diff)
## t = NaN, df = 29, p-value = NA
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
## NaN NaN
## sample estimates:
## mean of x
## 0
## Assign names to vectors
names(ind.spp$p.value) <- c("Acacon", "Acasup", "Canros", "Lobalp", "Phydub",
"Acabop", "Sterile.sp", "Calare", "Rhichr", "Xanlin", "Xanpli",
"Xanele", "GrBr.cr")
names(ind.spp$t) <- c("Acacon", "Acasup", "Canros", "Lobalp", "Phydub",
"Acabop", "Sterile.sp", "Calare", "Rhichr", "Xanlin", "Xanpli",
"Xanele", "GrBr.cr")
ind.spp$p.value <- p.adjust(ind.spp$p.value, method = "fdr")
do.call(cbind, ind.spp)
```

```
##           p.value          t
## Acacon    0.0129109 -3.3776290
## Acasup    0.0129109 -3.2420906
## Canros    0.0129109 -3.5818840
## Lobalp    0.1638226 -2.0413612
## Phydub    0.1674381 -1.9226188
## Acabop    0.3847787 -1.0747324
## Sterile.sp 0.3847787 -1.0000000
## Calare    0.2082316 -1.6076070
## Rhichr    0.2082316 -1.5803288
## Xanlin    0.5872370 -0.6169756
## Xanpli    0.7968379 -0.2598207
## Xanele    0.2082316 -1.5662320
## GrBr.cr   0.3847787  1.0000000
```

## Exploring causal pathways

I suggest that we develop these into a structural equation model. Here, I've analyzed several "causal pathways" in parts using correlation tests of the differences (susceptible - resistant) for each variable. Here are two hypothesized causal pathways, where lichen refers to the lichen community abundance, richness, diversity and composition:

- moth decreases light which decreases lichen
- moth increases litter which covers/decreases rocks which support/increase lichen

Only the litter pathway is supported. Although light is impacted by moth susceptibility, it isn't correlated with lichen. Litter, however, is positively affected by the moth infection. In addition, litter is negatively correlated with rock abundance and rock abundance is positively correlated with lichen. This is also supported by the indirect pathway between moth and lichen, which are negatively correlated, as would be expected by the product of the signs along the pathway (i.e. positive \* negative \* positive = negative).

```
### Test for correlations among effects
cor.test(tapply(l.dat[, "Light...average"], l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Big.rock.."], l.dat[, "Tree.pairs"], diff), method = "k")
```

```
##
## Kendall's rank correlation tau
##
## data: tapply(l.dat[, "Light...average"], l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Big.rock.."], l.dat[, "Tree.pairs"], diff)
## T = 229, p-value = 0.6972
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
## tau
## 0.05287356
```

```
cor.test(tapply(l.dat[, "Litter.."], l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Big.rock.."], l.dat[, "Tree.pairs"], diff), method = "k")
```

```
##
## Kendall's rank correlation tau
##
## data: tapply(l.dat[, "Litter.."], l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Big.rock.."], l.dat[, "Tree.pairs"], diff)
## T = 43, p-value = 3.161e-13
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
## tau
## -0.8022989
```

```
cor.test(tapply(l.dat[, "Litter.."], l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Light...average"], l.dat[, "Tree.pairs"], diff), method = "k")
```

```
##
## Kendall's rank correlation tau
##
## data: tapply(l.dat[, "Litter.."], l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Light...average"], l.dat[, "Tree.pairs"], diff)
## T = 193, p-value = 0.395
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
## tau
## -0.1126437
```

```
## Abun
cor.test(tapply(abun, l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Light...average"], l.dat[, "Tree.pairs"], diff), method = "k")
```

```
## Warning in cor.test.default(tapply(abun, l.dat[, "Tree.pairs"], diff),
## tapply(l.dat[, "Light...average"], l.dat[, "Tree.pairs"], diff), method = "k"): Cannot compute exact p-value with ties
```

```
##
## Kendall's rank correlation tau
##
## data: tapply(abun, l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Light...average"], l.dat[, "Tree.pairs"], diff)
```



```

## z = 0.92788, p-value = 0.3535
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
##      tau
## 0.1196779

cor.test(tapply(abun, l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Litter.."] , l.dat[, "Tree.pairs"], diff), method = "k")

## Warning in cor.test.default(tapply(abun, l.dat[, "Tree.pairs"], diff),
## tapply(l.dat[, : Cannot compute exact p-value with ties

##
## Kendall's rank correlation tau
##
## data:  tapply(abun, l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Litter.."], l.dat[, "Tree.pairs"]
## z = -2.9978, p-value = 0.00272
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
##      tau
## -0.3866516

cor.test(tapply(abun, l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Big.rock.."] , l.dat[, "Tree.pairs"], diff), method = "k")

## Warning in cor.test.default(tapply(abun, l.dat[, "Tree.pairs"], diff),
## tapply(l.dat[, : Cannot compute exact p-value with ties

##
## Kendall's rank correlation tau
##
## data:  tapply(abun, l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Big.rock.."], l.dat[, "Tree.pa
## z = 3.319, p-value = 0.0009035
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
##      tau
## 0.4280785

## rich
cor.test(tapply(rich, l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Light...average"] , l.dat[, "Tree.pairs"], diff), method = "k")

## Warning in cor.test.default(tapply(rich, l.dat[, "Tree.pairs"], diff),
## tapply(l.dat[, : Cannot compute exact p-value with ties

##
## Kendall's rank correlation tau
##
## data:  tapply(rich, l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Light...average"], l.dat[, "Tre
## z = 1.3285, p-value = 0.184
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
##      tau
## 0.1756536

cor.test(tapply(rich, l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Litter.."] , l.dat[, "Tree.pairs"], diff), method = "k")

```

```

## Warning in cor.test.default(tapply(rich, l.dat[, "Tree.pairs"], diff),
## tapply(l.dat[, : Cannot compute exact p-value with ties
##
## Kendall's rank correlation tau
##
## data: tapply(rich, l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Litter.."], l.dat[, "Tree.pairs"]
## z = -4.0214, p-value = 5.785e-05
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
## tau
## -0.5317081
cor.test(tapply(rich, l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Big.rocks.."] , l.dat[, "Tree.pairs"], diff), method = "k")

## Warning in cor.test.default(tapply(rich, l.dat[, "Tree.pairs"], diff),
## tapply(l.dat[, : Cannot compute exact p-value with ties
##
## Kendall's rank correlation tau
##
## data: tapply(rich, l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Big.rocks.."], l.dat[, "Tree.pa
## z = 4.0573, p-value = 4.964e-05
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
## tau
## 0.5364555
## shan
cor.test(tapply(shan, l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Light...average"] , l.dat[, "Tree.pairs"], diff), method = "k")

## Warning in cor.test.default(tapply(shan, l.dat[, "Tree.pairs"], diff),
## tapply(l.dat[, : Cannot compute exact p-value with ties
##
## Kendall's rank correlation tau
##
## data: tapply(shan, l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Light...average"], l.dat[, "Tre
## z = 1.5352, p-value = 0.1247
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
## tau
## 0.1983864
cor.test(tapply(shan, l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Litter.." ] , l.dat[, "Tree.pairs"], diff), method = "k")

## Warning in cor.test.default(tapply(shan, l.dat[, "Tree.pairs"], diff),
## tapply(l.dat[, : Cannot compute exact p-value with ties
##
## Kendall's rank correlation tau
##
## data: tapply(shan, l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Litter.."], l.dat[, "Tree.pairs"]
## z = -3.4632, p-value = 0.0005338
## alternative hypothesis: true tau is not equal to 0

```

```

## sample estimates:
##      tau
## -0.4475229
cor.test(tapply(shan, l.dat[, "Tree.pairs"], diff),
         tapply(l.dat[, "Big.rock.."] , l.dat[, "Tree.pairs"], diff), method = "k")

## Warning in cor.test.default(tapply(shan, l.dat[, "Tree.pairs"], diff),
## tapply(l.dat[, : Cannot compute exact p-value with ties

##
## Kendall's rank correlation tau
##
## data:  tapply(shan, l.dat[, "Tree.pairs"], diff) and tapply(l.dat[, "Big.rock.."], l.dat[, "Tree.pa
## z = 3.3918, p-value = 0.0006944
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
##      tau
## 0.4382956

```