LCN: Lichen interaction network study

 $MK \ Lau$

Results

```
### REML
### We know from Lamit's dissertation work that lichen communities are
### heritable, largely driven by bark roughness
### Do we find similar patterns?
## Create a list to generate a results table
h2.tab <- matrix("", 8, 4)
colnames(h2.tab) <- c("Response", "H2", "R2", "p-value")</pre>
## Total cover ~ genotype
ptc.reml <- lme4::lmer(I(PC^(1/2)) ~ (1 | geno),
                       data = onc.dat, REML = TRUE)
ptc.reml.pval <- RLRsim::exactRLRT(ptc.reml)</pre>
ptc.reml.pval
##
   simulated finite sample distribution of RLRT.
##
##
## (p-value based on 10000 simulated values)
##
## data:
## RLRT = 2.9627, p-value = 0.0358
fligner.test(onc.dat$PC^(1/2), onc.dat$geno)
## Fligner-Killeen test of homogeneity of variances
## data: onc.dat$PC^(1/2) and onc.dat$geno
## Fligner-Killeen:med chi-squared = 13.751, df = 12, p-value =
## 0.3169
shapiro.test(residuals(ptc.reml))
##
## Shapiro-Wilk normality test
## data: residuals(ptc.reml)
## W = 0.95096, p-value = 0.02174
h2.tab[1, "p-value"] <- ptc.reml.pval$"p.value"
h2.tab[1, "H2"] \leftarrow H2(ptc.reml, g = onc.dat$geno)
h2.tab[1, "R2"] <- R2(ptc.reml)
R2(ptc.reml)
## 0.1727875
```

```
h2.tab[1, "Response"] <- "Percent Lichen Cover"
## Species richness ~ genotype
spr.reml \leftarrow lme4::lmer(I(SR^(1/2)) \sim (1 \mid geno),
                        data = onc.dat, REML = TRUE)
spr.reml.pval <- RLRsim::exactRLRT(spr.reml)</pre>
spr.reml.pval
##
##
  simulated finite sample distribution of RLRT.
##
   (p-value based on 10000 simulated values)
##
## data:
## RLRT = 1.0001, p-value = 0.1391
shapiro.test(residuals(spr.reml))
    Shapiro-Wilk normality test
##
##
## data: residuals(spr.reml)
## W = 0.97364, p-value = 0.2467
fligner.test(onc.dat$SR^(1/2), onc.dat$geno)
##
## Fligner-Killeen test of homogeneity of variances
## data: onc.datSR^(1/2) and onc.datgeno
## Fligner-Killeen:med chi-squared = 13.276, df = 12, p-value =
## 0.3493
h2.tab[2, "p-value"] <- spr.reml.pval$"p.value"
h2.tab[2, "H2"] <- H2(spr.reml, g = onc.dat$geno)
h2.tab[2, "R2"] <- R2(spr.reml)
R2(spr.reml)
##
          R<sub>2</sub>c
## 0.09814791
h2.tab[2, "Response"] <- "Lichen Species Richness"
## Bark roughness REML
prb.reml <- lme4::lmer(I(BR^(1/2)) ~ (1 | geno), data = onc.dat, REML = TRUE)
prb.reml.pval <- RLRsim::exactRLRT(prb.reml)</pre>
prb.reml.pval
##
##
    simulated finite sample distribution of RLRT.
##
    (p-value based on 10000 simulated values)
##
##
## data:
## RLRT = 10.69, p-value = 5e-04
```

```
fligner.test(onc.dat$BR^(1/2), onc.dat$geno)
## Fligner-Killeen test of homogeneity of variances
##
## data: onc.dat$BR^(1/2) and onc.dat$geno
## Fligner-Killeen:med chi-squared = 6.1915, df = 12, p-value =
## 0.9061
shapiro.test(residuals(prb.reml))
## Shapiro-Wilk normality test
##
## data: residuals(prb.reml)
## W = 0.97975, p-value = 0.4529
h2.tab[3, "p-value"] <- prb.reml.pval$"p.value"
h2.tab[3, "H2"] <- H2(prb.reml, g = onc.dat$geno)
h2.tab[3, "R2"] <- R2(prb.reml)
R2(prb.reml)
## 0.3783496
h2.tab[3, "Response"] <- "Percent Rough Bark"
## condensed tannins REML
ct.reml <- lme4::lmer(I(CT^(1/4)) ~ (1 | geno), data = onc.dat, REML = TRUE)
ct.reml.pval <- RLRsim::exactRLRT(ct.reml)</pre>
ct.reml.pval
##
## simulated finite sample distribution of RLRT.
## (p-value based on 10000 simulated values)
##
## data:
## RLRT = 4.3224, p-value = 0.0146
fligner.test(onc.dat$CT^(1/4), onc.dat$geno)
##
## Fligner-Killeen test of homogeneity of variances
## data: onc.dat$CT^(1/4) and onc.dat$geno
## Fligner-Killeen:med chi-squared = 7.8941, df = 12, p-value =
## 0.7933
shapiro.test(residuals(ct.reml))
## Shapiro-Wilk normality test
##
## data: residuals(ct.reml)
## W = 0.74892, p-value = 2.431e-08
```

```
## CN ratio REML
cnr.reml <- lme4::lmer(I(CN^(1/1)) ~ (1 | geno), data = onc.dat, REML = TRUE)</pre>
## boundary (singular) fit: see ?isSingular
cnr.reml.pval <- RLRsim::exactRLRT(cnr.reml)</pre>
cnr.reml.pval
##
##
   simulated finite sample distribution of RLRT.
##
## (p-value based on 10000 simulated values)
##
## data:
## RLRT = 0, p-value = 1
fligner.test(onc.dat$CN^(1/1), onc.dat$geno)
##
## Fligner-Killeen test of homogeneity of variances
##
## data: onc.dat$CN^(1/1) and onc.dat$geno
## Fligner-Killeen:med chi-squared = 8.1116, df = 12, p-value =
## 0.7763
shapiro.test(residuals(cnr.reml))
## Shapiro-Wilk normality test
## data: residuals(cnr.reml)
## W = 0.92183, p-value = 0.001754
## Is species richness correlated with percent cover?
cor.test(onc.dat[, "SR"], onc.dat[, "PC"], data = onc.dat)
##
##
  Pearson's product-moment correlation
##
## data: onc.dat[, "SR"] and onc.dat[, "PC"]
## t = 8.3456, df = 55, p-value = 2.393e-11
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.6047186 0.8437321
## sample estimates:
##
         cor
## 0.7475023
## Were these correlated with bark roughness?
ptc.prb.lm \leftarrow lm(I(PC^{(1/2)}) \sim I(BR^{(1/2)}), data = onc.dat)
summary(ptc.prb.lm)
##
## Call:
## lm(formula = I(PC^(1/2)) \sim I(BR^(1/2)), data = onc.dat)
## Residuals:
##
       Min
               1Q Median
                                 3Q
                                        Max
```

```
## -5.9770 -1.6378 0.6333 1.9603 3.4658
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 4.4142
                           1.0901 4.049 0.000162 ***
## I(BR^(1/2))
                0.4942
                           0.1896 2.607 0.011730 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 2.485 on 55 degrees of freedom
## Multiple R-squared: 0.11, Adjusted R-squared: 0.09381
## F-statistic: 6.797 on 1 and 55 DF, p-value: 0.01173
fligner.test(onc.dat$PC, onc.dat$BR)
##
## Fligner-Killeen test of homogeneity of variances
## data: onc.dat$PC and onc.dat$BR
## Fligner-Killeen:med chi-squared = 27.401, df = 24, p-value =
## 0.2861
shapiro.test(residuals(ptc.prb.lm))
##
## Shapiro-Wilk normality test
## data: residuals(ptc.prb.lm)
## W = 0.95045, p-value = 0.02061
spr.prb.lm \leftarrow lm(I(SR^{(1)}) \sim I(BR^{(1/2)}), data = onc.dat)
summary(spr.prb.lm)
##
## Call:
## lm(formula = I(SR^(1)) \sim I(BR^(1/2)), data = onc.dat)
##
## Residuals:
               1Q Median
                               3Q
## -3.0420 -1.3123 -0.1178 1.2308 4.3519
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
                2.5015
                           0.8002
                                    3.126 0.00283 **
## (Intercept)
                           0.1392
                0.1709
## I(BR^(1/2))
                                   1.228 0.22456
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.824 on 55 degrees of freedom
## Multiple R-squared: 0.0267, Adjusted R-squared: 0.009003
## F-statistic: 1.509 on 1 and 55 DF, p-value: 0.2246
fligner.test(onc.dat$SR^(1), onc.dat$BR)
## Fligner-Killeen test of homogeneity of variances
```

##

```
## data: onc.dat$SR^(1) and onc.dat$BR
## Fligner-Killeen:med chi-squared = 26.046, df = 24, p-value =
## 0.3508
shapiro.test(residuals(spr.prb.lm))
## Shapiro-Wilk normality test
##
## data: residuals(spr.prb.lm)
## W = 0.97168, p-value = 0.2008
## COM ~ genotype + Bark roughness + PTC + SPR
set.seed(2)
rcom.ng.perm <- vegan::adonis2(onc.com.rel^(1/1) ~ BR + PC + SR,
                               data = onc.dat, perm = 10000, mrank = TRUE)
set.seed(2)
rcom.perm <- vegan::adonis2(onc.com.rel^(1/1) ~ geno + BR + PC + SR,
                            data = onc.dat, perm = 10000, mrank = TRUE)
set.seed(2)
com.ng.perm <- vegan::adonis2(onc.com^(1/1) ~ BR + PC + SR,
                              data = onc.dat, perm = 10000, mrank = TRUE)
set.seed(2)
com.perm <- vegan::adonis2(onc.com^(1/1) ~ geno + BR + PC + SR,
                           data = onc.dat, perm = 10000, mrank = TRUE)
rcom.ng.perm
## Permutation test for adonis under reduced model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 10000
## vegan::adonis2(formula = onc.com.rel^(1/1) ~ BR + PC + SR, data = onc.dat, permutations = 10000, mra
           Df SumOfSqs
                            R2
                                     F
                                           Pr(>F)
## BR
            1
                0.4398 0.03889 3.7408 0.006399 **
## PC
                3.8618 0.34151 32.8482 9.999e-05 ***
            1
## SR
                0.7754 0.06857 6.5958 0.000200 ***
            1
## Residual 53 6.2309 0.55102
## Total
          56 11.3079 1.00000
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
rcom.perm
## Permutation test for adonis under reduced model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 10000
## vegan::adonis2(formula = onc.com.rel^(1/1) ~ geno + BR + PC + SR, data = onc.dat, permutations = 100
           Df SumOfSqs
                                          Pr(>F)
                            R2
           12
                2.7463 0.24287 1.8221
                                           0.0029 **
## geno
## BR.
            1
                 0.1248 0.01104 0.9938
                                           0.3841
## PC
                2.6711 0.23622 21.2661 9.999e-05 ***
            1
                0.6159 0.05447 4.9036
            1
                                          0.0011 **
                5.1498 0.45541
## Residual 41
```

```
56 11.3079 1.00000
## Total
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
h2.tab[4, "p-value"] <- unlist(rcom.perm)["Pr(>F)1"]
h2.tab[4, "H2"] <- H2(rcom.perm, g = onc.dat$geno)
h2.tab[4, "R2"] \leftarrow R2(rcom.perm)
h2.tab[4, "Response"] <- "Lichen Community Composition"
## Is network similarity correlated with community composition?
ecodist::mantel(cn.d.onc ~ vegdist(onc.com.rel), mrank = TRUE)
                             pval2
                                       pval3 llim.2.5% ulim.97.5%
                  pval1
## 0.09198784 0.08300000 0.91800000 0.13900000 0.05293002 0.13002104
spr.d <- dist(onc.dat$SR)</pre>
ptc.d <- dist(onc.dat$PC)</pre>
prb.d <- dist(onc.dat$BR)</pre>
### rough -> cover -> rich -> net
ecodist::mantel(cn.d.onc ~ vegdist(onc.com.rel) + spr.d + ptc.d + prb.d, mrank = TRUE)
                                       pval3 llim.2.5% ulim.97.5%
                             pval2
     mantelr
                  pval1
## 0.06853395 0.15400000 0.84700000 0.31000000 0.02651488 0.13049408
## Partial Mantels using RFLP distance
ecodist::mantel(cn.mu.d.onc ~ rflp.d)
      mantelr
                    pval1
                                pval2
                                           pval3
                                                  llim.2.5% ulim.97.5%
ecodist::mantel(onc.com.mu.d ~ rflp.d)
                                       pval3 llim.2.5% ulim.97.5%
     mantelr
                  pval1
                             pval2
## 0.1179051 0.2760000 0.7250000 0.4980000 -0.2412808 0.2447645
ecodist::mantel(cn.mu.d.onc ~ onc.com.mu.d)
##
      mantelr
                    pval1
                                pval2
                                           pval3
                                                  llim.2.5% ulim.97.5%
## 0.29000439 0.08900000 0.91200000 0.08900000 -0.02092565 0.42837710
## Was lichen network similarity determined by genotype?
set.seed(1234)
cn.perm <- vegan::adonis2(cn.d.onc ~ geno + BR + PC + SR,</pre>
                         data = onc.dat, permutations = 10000, mrank = TRUE)
set.seed(1234)
cn.perm.ng <- vegan::adonis2(cn.d.onc ~ BR + PC + SR,</pre>
              data = onc.dat, permutations = 10000, mrank = TRUE)
cn.perm.ng
## Permutation test for adonis under reduced model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 10000
## vegan::adonis2(formula = cn.d.onc ~ BR + PC + SR, data = onc.dat, permutations = 10000, mrank = TRUE
           Df SumOfSqs
                           R2
                                    F
                                         Pr(>F)
## BR
            1
                 61.42 0.03968 4.1680
                                        0.03770 *
## PC
            1
                49.47 0.03197 3.3573
                                        0.06779 .
## SR
            1 655.76 0.42373 44.5034 9.999e-05 ***
```

```
## Residual 53 780.96 0.50462
## Total 56 1547.61 1.00000
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
cn.perm
## Permutation test for adonis under reduced model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 10000
## vegan::adonis2(formula = cn.d.onc ~ geno + BR + PC + SR, data = onc.dat, permutations = 10000, mrank
           Df SumOfSqs R2 F
                                         Pr(>F)
## geno
          12 450.52 0.29111 2.6902 0.009399 **
                 29.11 0.01881 2.0858 0.146985
## BR
           1
## PC
                 30.01 0.01939 2.1504 0.144086
            1
           1 465.78 0.30097 33.3755 9.999e-05 ***
## SR
## Residual 41 572.18 0.36972
## Total 56 1547.61 1.00000
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
h2.tab[5, "p-value"] <- as.matrix(cn.perm)[1, "Pr(>F)"]
h2.tab[5, "H2"] <- H2(cn.perm, g = onc.dat[, "geno"], perm =10000)
h2.tab[5, "R2"] <- R2(cn.perm)
h2.tab[5, "Response"] <- "Lichen Network"
                                       # db rda for network similarity
dbr.cn.geno <- vegan::dbrda(cn.d.onc ~ geno, data = onc.dat, distance = "bray")
anova(dbr.cn.geno, permutations = 5000)
## Permutation test for dbrda under reduced model
## Permutation: free
## Number of permutations: 5000
## Model: vegan::dbrda(formula = cn.d.onc ~ geno, data = onc.dat, distance = "bray")
          Df Variance F Pr(>F)
                8.045 1.5057 0.1362
           12
## Model
## Residual 44
                19.591
H2(dbr.cn.geno)
## [1] 0.2911089
## What aspects of networks explained the similiarity?
## L = number of edges, LD = link density, C = connectivity,
## dcen = degree centrality
link.reml \leftarrow lme4::lmer(I(log(L + 0.00000001)) \sim (1 | geno),
                         data = onc.dat, REML = TRUE)
link.reml.pval <- RLRsim::exactRLRT(link.reml, nsim = 50000)</pre>
link.reml.pval
##
## simulated finite sample distribution of RLRT.
##
## (p-value based on 50000 simulated values)
##
```

```
## data:
## RLRT = 2.0484, p-value = 0.06602
fligner.test(log(onc.dat$L + 0.0000001), onc.dat$geno)
## Fligner-Killeen test of homogeneity of variances
## data: log(onc.dat$L + 1e-07) and onc.dat$geno
## Fligner-Killeen:med chi-squared = 11.991, df = 12, p-value =
## 0.4464
shapiro.test(residuals(link.reml))
##
## Shapiro-Wilk normality test
##
## data: residuals(link.reml)
## W = 0.83643, p-value = 2.036e-06
h2.tab[6, "p-value"] <- link.reml.pval$"p.value"
h2.tab[6, "H2"] <- H2(link.reml, g = onc.dat$geno)
h2.tab[6, "R2"] <- R2(link.reml)
R2(link.reml)
##
         R<sub>2</sub>c
## 0.1701568
h2.tab[6, "Response"] <- "Number of Network Links"
                                         # network centrality
cen.reml <- lme4::lmer(I(Cen^(1/2)) ~ (1 | geno),
                       data = onc.dat, REML = TRUE)
cen.reml.pval <- RLRsim::exactRLRT(cen.reml, nsim = 50000)</pre>
cen.reml.pval
##
## simulated finite sample distribution of RLRT.
##
## (p-value based on 50000 simulated values)
##
## data:
## RLRT = 2.7801, p-value = 0.04076
fligner.test(onc.dat$L^(1/1), onc.dat$geno)
## Fligner-Killeen test of homogeneity of variances
## data: onc.dat$L^(1/1) and onc.dat$geno
## Fligner-Killeen:med chi-squared = 14.241, df = 12, p-value =
## 0.2856
shapiro.test(residuals(cen.reml))
##
## Shapiro-Wilk normality test
##
```

```
## data: residuals(cen.reml)
## W = 0.90072, p-value = 0.0002041
h2.tab[7, "p-value"] <- cen.reml.pval$"p.value"
h2.tab[7, "H2"] \leftarrow H2(cen.reml, g = onc.dat$geno)
h2.tab[7, "R2"] <- R2(cen.reml)
R2(cen.reml)
## 0.2016649
h2.tab[7, "Response"] <- "Network Centrality"
                                         # network modularity
mod.reml <- lme4::lmer(I(onc.ns[, "mod.lik"]^(1/4)) ~ (1 | geno),
                       data = onc.dat, REML = TRUE)
mod.reml.pval <- RLRsim::exactRLRT(mod.reml)</pre>
mod.reml.pval
##
## simulated finite sample distribution of RLRT.
##
## (p-value based on 10000 simulated values)
##
## data:
## RLRT = 0.23363, p-value = 0.2809
fligner.test(onc.ns[, "mod.lik"]^(1/4), onc.dat$geno)
##
## Fligner-Killeen test of homogeneity of variances
## data: onc.ns[, "mod.lik"]^(1/4) and onc.dat$geno
## Fligner-Killeen:med chi-squared = 13.439, df = 12, p-value =
## 0.3379
shapiro.test(residuals(mod.reml))
##
## Shapiro-Wilk normality test
## data: residuals(mod.reml)
## W = 0.54001, p-value = 4.252e-12
h2.tab[8, "p-value"] <- mod.reml.pval$"p.value"
h2.tab[8, "H2"] <- H2(mod.reml, g = onc.dat$geno)
h2.tab[8, "R2"] <- R2(mod.reml)
h2.tab[8, "Response"] <- "Network Modularity"</pre>
                                         # network stats in relation to other variables
L.aov \leftarrow aov(I(log(L + 0.000001)) ~ BR + PC + SR, data = onc.dat)
summary(L.aov)
               Df Sum Sq Mean Sq F value
##
                                           Pr(>F)
## BR.
               1 102.3
                           102.3 2.776
                                           0.1016
                           239.6 6.504
## PC
                1 239.6
                                           0.0137 *
## SR
               1 957.0 957.0 25.980 4.71e-06 ***
## Residuals 53 1952.2
                           36.8
```

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
shapiro.test(residuals(L.aov))
## Shapiro-Wilk normality test
##
## data: residuals(L.aov)
## W = 0.9629, p-value = 0.07794
cen.aov \leftarrow aov(I(Cen^(1/2)) ~ BR + PC + SR, data = onc.dat)
summary(cen.aov)
##
              Df Sum Sq Mean Sq F value
                                          Pr(>F)
## BR
                   3.77
                           3.77
                                 2.174
                                           0.146
               1
                           6.46 3.724
## PC
                  6.46
                                           0.059
               1
               1 56.48
                          56.48 32.552 5.31e-07 ***
## SR
              53 91.95
## Residuals
                           1.73
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
shapiro.test(residuals(cen.aov))
##
  Shapiro-Wilk normality test
##
## data: residuals(cen.aov)
## W = 0.97222, p-value = 0.2126
mod.aov \leftarrow aov(I(onc.ns[, "mod.lik"]^(1/4)) \sim BR + PC + SR, data = onc.dat)
summary(mod.aov)
##
              Df Sum Sq Mean Sq F value
                                          Pr(>F)
## BR
               1 0.0442 0.0442 0.787
                                           0.379
## PC
               1 0.0879 0.0879
                                 1.564
                                           0.217
               1 1.3799 1.3799 24.558 7.76e-06 ***
## SR
## Residuals
              53 2.9781 0.0562
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
shapiro.test(residuals((mod.aov)))
##
##
  Shapiro-Wilk normality test
## data: residuals((mod.aov))
## W = 0.9201, p-value = 0.001078
cor.test(onc.ns[, "L"], onc.ns[, "Cen"])
##
## Pearson's product-moment correlation
## data: onc.ns[, "L"] and onc.ns[, "Cen"]
## t = 13.37, df = 55, p-value < 2.2e-16
## alternative hypothesis: true correlation is not equal to 0
```

```
## 95 percent confidence interval:
## 0.7950728 0.9244074
## sample estimates:
##
         cor
## 0.8744752
                                         # are these metrics correlated with network similarity
L.d <- dist(onc.dat$L)
cen.d <- dist(onc.dat$Cen)</pre>
mod.d <- dist(cn.mod.onc)</pre>
cn.L.cen.perm <- adonis2(cn.d.onc ~ L + Cen, data = onc.dat, mrank = TRUE)</pre>
## So, are there patterns in the centrality of individual lichen species?
sppcen.test <- apply(cen.spp[, apply(cen.spp, 2, sum) >= 2], 2, function(x)
    lme4::lmer(I(x^(1/2)) \sim (1 \mid geno), data = onc.dat, REML = TRUE))
## boundary (singular) fit: see ?isSingular
sppcen.pval <- lapply(sppcen.test, RLRsim::exactRLRT)</pre>
sppcen.tab <- do.call(rbind, lapply(sppcen.pval, function(x)</pre>
    c(x[["statistic"]], x[["p.value"]])))
sppcen.h2 <- round(unlist(lapply(sppcen.test, H2)), 3)</pre>
sppcen.h2
            Cs
                  Ls
                        Ch
                               Χm
                                     Pm
## 0.000 0.127 0.000 0.258 0.201 0.000 0.000
## Mean centrality of species
sort(apply(cen.spp, 2, mean), decreasing = TRUE)
##
           Cs
                      Ch
                                  Ls
                                             R.s
                                                                    Pm
## 0.73204678 0.54157218 0.39722829 0.18378675 0.14553120 0.07914127
##
                      Pu
## 0.06376775 0.02105263 0.00000000
## Ordinations
### nits = 10,
### iconf = random
### epsilon = 1e-12 = acceptable change in stress
### maxit = 500 = maximum number of iterations
ord.com <- nmds.min(nms.com, 3)</pre>
## Minimum stress for given dimensionality: 0.1008923
## r^2 for minimum stress configuration: 0.9357192
## Minimum stress for given dimensionality: 0.1008923
## r^2 for minimum stress configuration: 0.9357192
ord.cn <- nmds.min(nms.cn, 2)
## Minimum stress for given dimensionality: 0.01065177
## r^2 for minimum stress configuration: 0.9993026
## Minimum stress for given dimensionality: 0.01065177
## r^2 for minimum stress configuration: 0.9993026
## checking variance explained by ordinations
ord1.cn.reml <- lme4::lmer(I(ord.cn[, 1]^(1/1)) ~ (1 | geno),
```

```
data = onc.dat, REML = TRUE)
ord2.cn.reml <- lme4::lmer(I(ord.cn[, 2]^(1/1)) \sim (1 | geno),
                       data = onc.dat, REML = TRUE)
ord1.cn.reml.pval <- RLRsim::exactRLRT(ord1.cn.reml)</pre>
ord2.cn.reml.pval <- RLRsim::exactRLRT(ord2.cn.reml)</pre>
ord1.cn.reml.pval
##
## simulated finite sample distribution of RLRT.
##
## (p-value based on 10000 simulated values)
##
## data:
## RLRT = 1.0221, p-value = 0.134
ord2.cn.reml.pval
##
##
   simulated finite sample distribution of RLRT.
##
## (p-value based on 10000 simulated values)
##
## RLRT = 0.5618, p-value = 0.2049
fligner.test(ord.cn[, 1]^(1/1), onc.dat$geno)
##
## Fligner-Killeen test of homogeneity of variances
## data: ord.cn[, 1]^(1/1) and onc.dat$geno
## Fligner-Killeen:med chi-squared = 16.805, df = 12, p-value =
## 0.1571
fligner.test(ord.cn[, 2]^(1/1), onc.dat$geno)
##
## Fligner-Killeen test of homogeneity of variances
## data: ord.cn[, 2]^(1/1) and onc.dat$geno
## Fligner-Killeen:med chi-squared = 9.9165, df = 12, p-value =
## 0.6233
ord1.com.reml <- lme4::lmer(I(ord.com[, 1]^(1/1)) ~ (1 | geno),
                       data = onc.dat, REML = TRUE)
ord2.com.reml <- lme4::lmer(I(ord.com[, 2]^(1/1)) ~ (1 | geno),
                       data = onc.dat, REML = TRUE)
ord1.com.reml.pval <- RLRsim::exactRLRT(ord1.com.reml)</pre>
ord2.com.reml.pval <- RLRsim::exactRLRT(ord2.com.reml)</pre>
ord1.com.reml.pval
##
##
   simulated finite sample distribution of RLRT.
##
##
    (p-value based on 10000 simulated values)
##
## data:
```

```
## RLRT = 0.1669, p-value = 0.3035
ord2.com.reml.pval
##
##
   simulated finite sample distribution of RLRT.
##
##
   (p-value based on 10000 simulated values)
##
## data:
## RLRT = 0.98197, p-value = 0.1381
fligner.test(ord.com[, 1]^(1/1), onc.dat$geno)
##
## Fligner-Killeen test of homogeneity of variances
## data: ord.com[, 1]^(1/1) and onc.dat$geno
## Fligner-Killeen:med chi-squared = 9.3187, df = 12, p-value =
## 0.6755
fligner.test(ord.com[, 2]^(1/1), onc.dat$geno)
##
## Fligner-Killeen test of homogeneity of variances
## data: ord.com[, 2]^(1/1) and onc.dat$geno
## Fligner-Killeen:med chi-squared = 16.947, df = 12, p-value =
## 0.1516
fligner.test(ord.com[, 3]^(1/1), onc.dat$geno)
## Fligner-Killeen test of homogeneity of variances
## data: ord.com[, 3]^(1/1) and onc.dat$geno
## Fligner-Killeen:med chi-squared = 14.943, df = 12, p-value =
## 0.2446
summary(lm(ord.cn[, 1] ~ SR + PC, data = onc.dat))
##
## Call:
## lm(formula = ord.cn[, 1] ~ SR + PC, data = onc.dat)
## Residuals:
       Min
                 1Q
                      Median
                                   3Q
                                            Max
## -10.6007 -1.7887
                       0.1726
                               2.2110
                                         6.7059
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
                                   3.593 0.000706 ***
## (Intercept) 3.77927
                          1.05175
## SR
              -2.89115
                          0.39475 -7.324 1.23e-09 ***
## PC
               0.10728
                          0.02215
                                   4.844 1.11e-05 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 3.596 on 54 degrees of freedom
```

```
## Multiple R-squared: 0.5025, Adjusted R-squared: 0.4841
## F-statistic: 27.27 on 2 and 54 DF, p-value: 6.508e-09
summary(lm(ord.cn[, 2] ~ SR + PC, data = onc.dat))
##
## Call:
## lm(formula = ord.cn[, 2] ~ SR + PC, data = onc.dat)
##
## Residuals:
##
      Min
               1Q Median
                               3Q
## -2.4080 -0.9426 -0.6151 1.3669 2.9279
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
                          0.420811 -2.716 0.008846 **
## (Intercept) -1.143124
               0.561645
                          0.157944 3.556 0.000793 ***
## PC
              -0.013722
                          0.008862 -1.548 0.127384
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.439 on 54 degrees of freedom
## Multiple R-squared: 0.2223, Adjusted R-squared: 0.1935
## F-statistic: 7.718 on 2 and 54 DF, p-value: 0.001127
summary(lm(ord.com[, 1] ~ SR + PC, data = onc.dat))
##
## Call:
## lm(formula = ord.com[, 1] ~ SR + PC, data = onc.dat)
## Residuals:
       Min
                 1Q
                     Median
                                   3Q
## -0.18241 -0.09091 -0.01606 0.05475 0.65204
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) -0.5145496 0.0395271 -13.018 < 2e-16 ***
               0.0527258 0.0148358
                                      3.554 0.000798 ***
## PC
               0.0058018 0.0008324
                                     6.970 4.61e-09 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.1351 on 54 degrees of freedom
## Multiple R-squared: 0.8048, Adjusted R-squared: 0.7976
## F-statistic: 111.3 on 2 and 54 DF, p-value: < 2.2e-16
summary(lm(ord.com[, 2] ~ SR + PC, data = onc.dat))
##
## Call:
## lm(formula = ord.com[, 2] ~ SR + PC, data = onc.dat)
## Residuals:
       Min
                      Median
                                   3Q
                                           Max
                 1Q
## -0.54228 -0.11829 0.03558 0.16463 0.50365
```

```
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
                          0.068196 -3.287 0.00178 **
## (Intercept) -0.224171
                                    0.607 0.54634
## SR.
               0.015539
                         0.025596
## PC
                0.002973 0.001436
                                    2.070 0.04328 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.2331 on 54 degrees of freedom
## Multiple R-squared: 0.2151, Adjusted R-squared: 0.1861
                 7.4 on 2 and 54 DF, p-value: 0.001444
## F-statistic:
## Lichen size distribution
## X. gallericulata thalli are about 0.22 +/- 0.003 cm^2 on average
## with an average median size of 0.12 +/- 0.001 \text{ cm}^2
## and, size does not vary significantly with genotype.
xgs.reml <- lme4::lmer(I(mean.thallus) ~ (1 | geno),</pre>
                       data = xgs.data[xgs.data$geno %in% names(which(table(xgs.data$geno) > 2)), ],
                       REML = TRUE)
xgs.median.reml <- lme4::lmer(median.thallus ~ (1 | geno),
                       data = xgs.data[xgs.data$geno %in% names(which(table(xgs.data$geno) > 2)), ],
                       REML = TRUE)
RLRsim::exactRLRT(xgs.reml)
##
##
  simulated finite sample distribution of RLRT.
##
   (p-value based on 10000 simulated values)
##
## data:
## RLRT = 2.4792, p-value = 0.0474
RLRsim::exactRLRT(xgs.median.reml)
##
##
   simulated finite sample distribution of RLRT.
##
   (p-value based on 10000 simulated values)
##
## data:
## RLRT = 0.092023, p-value = 0.3418
fligner.test(xgs.data$mean.thallus, xgs.data$geno)
##
## Fligner-Killeen test of homogeneity of variances
## data: xgs.data$mean.thallus and xgs.data$geno
## Fligner-Killeen:med chi-squared = 13.244, df = 17, p-value =
fligner.test(xgs.data$median.thallus, xgs.data$geno)
## Fligner-Killeen test of homogeneity of variances
```

##

```
## data: xgs.data$median.thallus and xgs.data$geno
## Fligner-Killeen:med chi-squared = 19.374, df = 17, p-value =
## 0.3075
mean(xgs.data$mean.thallus)
## [1] 0.1808442
sd(xgs.data$mean.thallus) / (length(xgs.data$mean.thallus) - 1)
## [1] 0.001845945
mean(xgs.data$median.thallus)
## [1] 0.1170852
sd(xgs.data$median.thallus) / (length(xgs.data$median.thallus) - 1)
## [1] 0.001223999
                                          # ONC and Wild Stand (Uintah)
                                          # Network distances
cn.all <- cn.wild
for (i in 1:length(cn.wild)){
    cn.all[[i]] <- cn.wild[[i]][match(rownames(cn.onc[[1]]), rownames(cn.wild[[i]])),</pre>
                                 match(colnames(cn.onc[[1]]), colnames(cn.wild[[i]]))]
}
cn.all <- append(cn.all, cn.onc)</pre>
cn.d.all <- netDist(cn.all, method = "bc")</pre>
cn.nms.geno <- c(rep("wild", length(cn.wild)), onc.geno)</pre>
if (!exists("cn.nms.all")){cn.nms.all <- nmds.min(nmds(cn.d.all, 2, 2))}</pre>
vec.all <- envfit(cn.nms.all[cn.nms.geno != "wild",], vec.env[, c("SR", "PC", "BR")], na.rm = TRUE)</pre>
Tables
```

```
h2.tab[, "H2"] <- round(as.numeric(h2.tab[, "H2"]), digits = 5)
h2.tab[, "R2"] <- round(as.numeric(h2.tab[, "R2"]), digits = 5)
h2.tab[, "p-value"] <- round(as.numeric(h2.tab[, "p-value"]), digits = 5)
h2.tab <- h2.tab[order(h2.tab[, "H2"], decreasing = TRUE), ]
h2.xtab <- xtable::xtable(h2.tab, caption =
    "Genotypic effects of cottonwood trees on the associated lichen community.",
                           label = "tab:h2_table")
print(h2.xtab,
      type = "latex",
      include.rownames = FALSE,
      include.colnames = TRUE
\% latex table generated in R 3.5.2 by xtable 1.8-3 package \% Mon Mar 25 16:21:58 2019
                                         # community permanova
rcom.ng.perm.xtab <- xtable::xtable(rcom.ng.perm, caption =</pre>
    "PerMANOVA Pseudo-F Table showing the predictors of community similarity.",
                          label = "tab:com_ng_perm")
print(rcom.ng.perm.xtab,
      type = "latex",
```

Response	H2	R2	p-value
Percent Rough Bark	0.37835	0.37835	5e-04
Network Centrality	0.20166	0.20166	0.04076
Percent Lichen Cover	0.17279	0.17279	0.0358
Number of Network Links	0.17016	0.17016	0.06602
Lichen Community Composition	0.16093	0.24287	0.0029
Lichen Species Richness	0.09815	0.09815	0.1391
Lichen Network	0.06252	0.29111	0.0094
Network Modularity	0.05731	0.05731	0.2809

Table 1: Genotypic effects of cottonwood trees on the associated lichen community.

```
include.rownames = TRUE,
include.colnames = TRUE
)
```

% latex table generated in R 3.5.2 by xtable 1.8-3 package % Mon Mar 25 16:21:58 2019

	Df	SumOfSqs	R2	F	Pr(>F)
BR	1	0.44	0.04	3.74	0.0064
PC	1	3.86	0.34	32.85	0.0001
SR	1	0.78	0.07	6.60	0.0002
Residual	53	6.23	0.55		
Total	56	11.31	1.00		

Table 2: PerMANOVA Pseudo-F Table showing the predictors of community similarity.

% latex table generated in R 3.5.2 by xtable 1.8-3 package % Mon Mar 25 16:21:58 2019

	Df	SumOfSqs	R2	F	Pr(>F)
geno	12	2.75	0.24	1.82	0.0029
BR	1	0.12	0.01	0.99	0.3841
PC	1	2.67	0.24	21.27	0.0001
SR	1	0.62	0.05	4.90	0.0011
Residual	41	5.15	0.46		
Total	56	11.31	1.00		

Table 3: PerMANOVA Pseudo-F Table showing the predictors of community similarity.

```
include.colnames = TRUE
)
```

% latex table generated in R 3.5.2 by x table 1.8-3 package % Mon Mar 25 16:21:58 2019

	Df	SumOfSqs	R2	F	Pr(>F)
BR	1	61.42	0.04	4.17	0.0377
PC	1	49.47	0.03	3.36	0.0678
SR	1	655.76	0.42	44.50	0.0001
Residual	53	780.96	0.50		
Total	56	1547.61	1.00		

Table 4: PerMANOVA Pseudo-F Table showing the predictors of network similarity.

%latex table generated in R3.5.2 by x
table 1.8-3 package % Mon Mar25 16:21:58
 2019

	Df	SumOfSqs	R2	F	Pr(>F)
geno	12	450.52	0.29	2.69	0.0094
BR	1	29.11	0.02	2.09	0.1470
PC	1	30.01	0.02	2.15	0.1441
SR	1	465.78	0.30	33.38	0.0001
Residual	41	572.18	0.37		
Total	56	1547.61	1.00		

Table 5: PerMANOVA Pseudo-F Table showing the predictors of network similarity.

% latex table generated in R 3.5.2 by x table 1.8-3 package % Mon Mar 25 16:21:58 2019

	Df	Sum Sq	Mean Sq	F value	Pr(>F)
BR	1	102.25	102.25	2.78	0.1016
PC	1	239.57	239.57	6.50	0.0137
SR	1	956.96	956.96	25.98	0.0000
Residuals	53	1952.23	36.83		

Table 6: ANOVA F Table showing the predictors of the number of network links.

% latex table generated in R 3.5.2 by xtable 1.8-3 package % Mon Mar 25 16:21:58 2019

	Df	Sum Sq	Mean Sq	F value	Pr(>F)
BR	1	3.77	3.77	2.17	0.1463
PC	1	6.46	6.46	3.72	0.0590
SR	1	56.48	56.48	32.55	0.0000
Residuals	53	91.95	1.73		

Table 7: ANOVA F Table showing the predictors of network centralization.

% latex table generated in R 3.5.2 by xtable 1.8-3 package % Mon Mar 25 16:21:58 2019

	Df	SumOfSqs	R2	F	Pr(>F)
L	1	1330.80	0.86	734.67	0.0010
Cen	1	118.99	0.08	65.69	0.0010
Residual	54	97.82	0.06		
Total	56	1547.61	1.00		

Table 8: PerMANOVA Pseudo-F Table showing the predictors of network similarity.

Plots

Figure: Genotype barplots Community composition NMDS with vectors

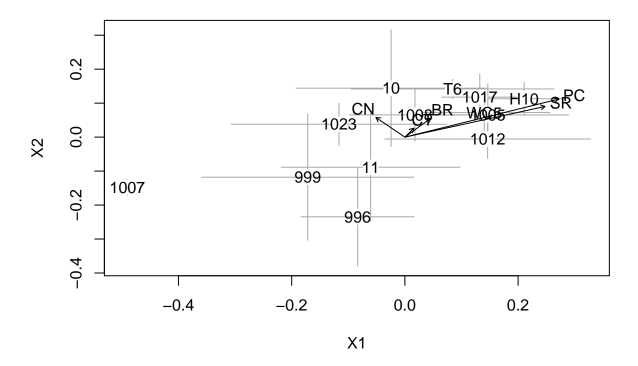


Figure: Lichen networks

```
par(mfrow = c(2, 2), mar = c(0, 0.1, 1.0, 0.1))
set.seed(123)
net.col <- sign(netMean(cn.mu.onc))</pre>
net.col[net.col == -1] <- 2
net.col[net.col == 1] <- 1</pre>
coord <- gplot(abs(netMean(cn.mu.onc)), gmode = "digraph",</pre>
      displaylabels = TRUE,
      edge.lwd = abs(netMean(cn.mu.onc)) * 20,
      edge.col = net.col,
      vertex.col = "black",
      vertex.cex = 0.5,
      arrowhead.cex = 0.5,
      label.cex = 1,
      main = "All Genotypes")
cn.mu.plot <- cn.mu.onc[names(cn.mu.onc) %in% c("996", "11", "1008")]</pre>
for (i in 1:length(cn.mu.plot)){
        net.col <- sign(cn.mu.plot[[i]])</pre>
        net.col[net.col == -1] <- 2
        net.col[net.col == 1] <- 1
        set.seed(123)
        gplot(abs(cn.mu.plot[[i]]), gmode = "digraph",
              displaylabels = TRUE,
              coord = coord,
              edge.lwd = abs(cn.mu.plot[[i]]) * 20,
              edge.col = net.col,
              vertex.col = "black",
              vertex.cex = 0.5,
              arrowhead.cex = 0.5,
              label.cex = 1,
```

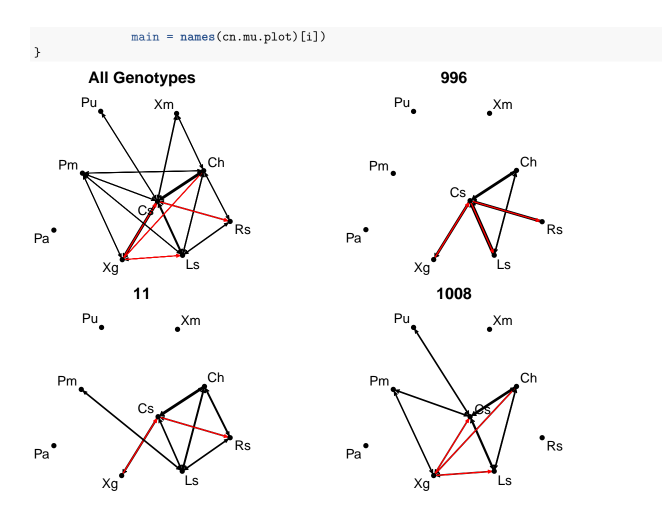


Figure: Genotype network similarity by genotype

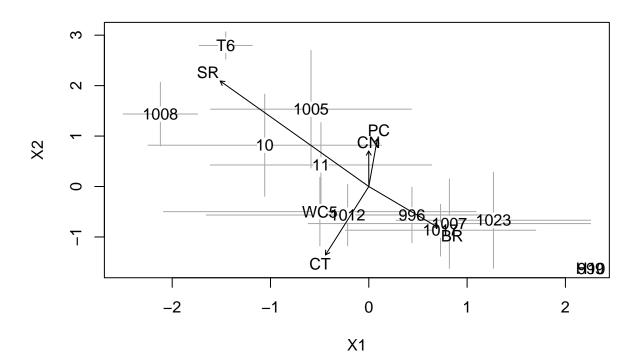
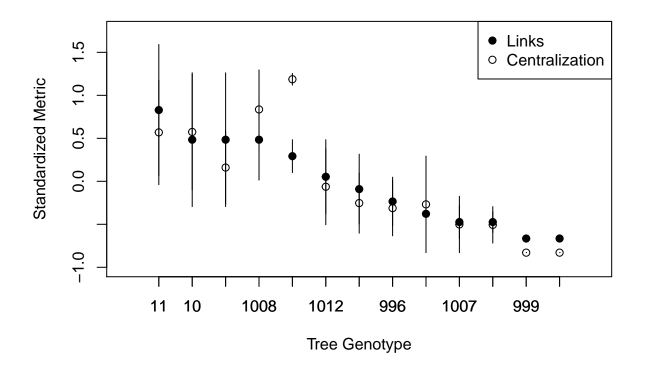


Figure: (A) Linkage and centrality by genotype and (B) Total cover and species richness predict L and Cen



Supplementary Figure: Lichen size distribution

0

-0.1

0.0

0.1

Median Lichen Thallus Area (cm^2)

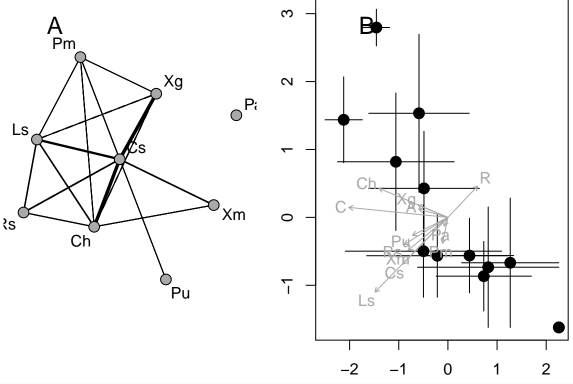
0.2

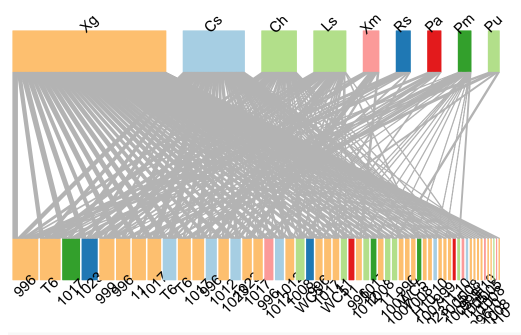
0.3

0.4

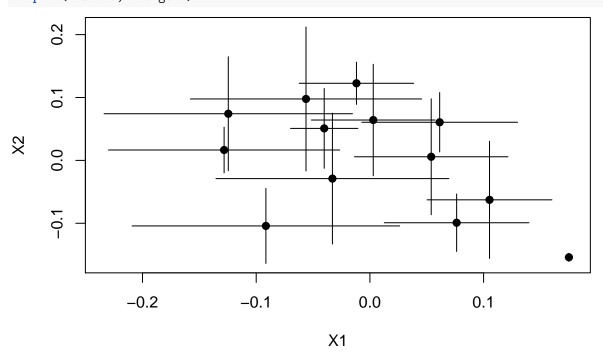
0.5

Figure 2





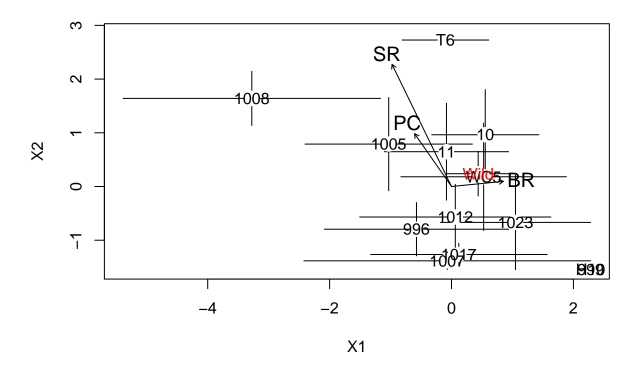
ch.plot(nms.onc, onc.geno)



Х2 ## Х1 ## 10 -0.091479884 -0.104244255 ## 1005 -0.124540472 0.074171596 **##** 1007 0.175012652 -0.154102531 ## 1008 0.061349598 0.060637950 ## 1012 -0.056201035 0.097682862 ## 1017 -0.040197275 0.050863036 ## 1023 0.053916695 0.005735269 ## 11 -0.032978298 -0.028980365 ## 996 0.076271011 -0.099167595 ## 999 0.105185439 -0.062733516

```
## H10 -0.011830997 0.122603983
## T6
          0.002941633 0.064173827
        -0.128224482 0.016507373
## WC5
## plot(cv.onc, col = "grey30")
## legend("topleft", legend = "A")
g.order <- tapply(ns.onc[, "C"], onc.geno, mean)</pre>
g.order <- names(g.order)[order(g.order, decreasing = TRUE)]</pre>
onc.g <- factor(onc.geno, levels = g.order)</pre>
plot(ns.onc[, "C"] ~ onc.g, xlab = "Tree Genotype", ylab = "Lichen Network Connectance (C)")
Lichen Network Connectance (C)
      0.15
                                   0
       0.10
                                                          0
       0.05
       0.00
                                 1008
                                            1012
                                                         996
                                                                    1007
                                                                                999
                 11
                       10
                                             Tree Genotype
```

Which wild uintah trees are similar to garden trees?



Send results to manuscript

```
manuscript.dir <- "../../lcn_manuscript"</pre>
### Send tables and figures to manuscript directory
if (exists("manuscript.dir")){
    tabs.figs <- dir(manuscript.dir)</pre>
    tab.fig.update <- dir("../results/lcn_notebook_files/figure-latex/",</pre>
                           full.names = TRUE)[
                               dir("../results/lcn_notebook_files/figure-latex/") %in% tabs.figs]
    tab.fig.update <- c(tab.fig.update,</pre>
                         dir("../docs", full.names = TRUE)[dir("../docs") %in% tabs.figs])
    sapply(tab.fig.update, file.copy, to = manuscript.dir, overwrite = TRUE)
                                          # supplementary figures
    si.dir <- paste0(manuscript.dir, "/supplement")</pre>
    si <- dir(si.dir)</pre>
    si.update <- dir("../results/lcn_notebook_files/figure-latex/",</pre>
                      full.names = TRUE)[
                          dir("../results/lcn_notebook_files/figure-latex/") %in% si]
    si.update <- c(si.update, dir("../docs", full.names = TRUE)[dir("../docs") %in% si])
    sapply(si.update, file.copy, to = si.dir,
           overwrite = TRUE)
```

named list()

Loading and pre-processing data

```
## Check for supporting packages
pkg.list <- c("MuMIn", "lme4", "RLRsim", "vegan", "ecodist", "bipartite", "RColorBrewer", "enaR", "devt
                                          # Install packages that are not installed
if (any(!(pkg.list %in% installed.packages()[, 1]))){
    sapply(pkg.list[which(!(pkg.list %in% installed.packages()[, 1]))],
           install.packages, dependencies = TRUE, repos='http://cran.us.r-project.org')
}
                                          # Check non-CRAN packages
if (!("ComGenR" %in% installed.packages()[, 1])){
    devtools::install_github("CommunityGeneticsAnalyses/ComGenR")
if (!("coNet" %in% installed.packages()[, 1])){
    devtools::install_github("CommunityGeneticsAnalyses/coNet")
}
                                          # Load libraries
sapply(c(pkg.list, "ComGenR", "coNet"), library, quietly = TRUE, character.only = TRUE)
### Loading data objects
## onc.com = "community" occurrences summed across all cells for each tree
## onc.q = occurrence matrices separated out for each tree
## onc.geno = genotypes
## prb.onc = percent rough bark (averaged between the upper and lower)
### Data notes:
## Trees were removed from the analysis genotype RL6 and N1.31
## No physciods
## Lecanoras merged
                                          # Loading data
xgal.size <- gdata::read.xls("../data/lichen_networks/ONC_Xgal_SizeData_May2011.xls")</pre>
garden.data <- read.csv('../data/lichen_networks/LCO_data_ONC_PIT.csv')</pre>
                                          # remove genotype RL6 and N1.31
garden.data <- garden.data[garden.data$Geno!='RL6',]</pre>
garden.data <- garden.data[garden.data$Tree!='N1.31',]</pre>
                                          #separate onc
garden.data[,1] <- as.character(garden.data[,1])</pre>
g1 <- substr(garden.data[,1],2,2)</pre>
g1[g1!='P'] <- 'onc'
onc <- garden.data[g1 == 'onc',]</pre>
colnames(onc)[which(colnames(onc) == "Ls")] <- "Lh"</pre>
pit <- garden.data[g1 == 'P',]</pre>
                     #tree overlap between years
unique(onc$Tree[onc$Year == '2010']) %in% unique(onc$Tree[onc$Year == '2011'])
unique(onc$Tree[onc$Year == '2011']) %in% unique(onc$Tree[onc$Year == '2010'])
                                         # Checking the data
if (!(all(table(onc[,1]) == 100))){for (i in 1:1000){print('Warning: check input data!!!')}}
                                          # Separate trees
                                          # onc
colnames(onc)[7:ncol(onc)] <- substr(colnames(onc)[7:ncol(onc)],1,2)</pre>
onc.q <- split(onc,paste(onc[,1],onc[,2]))</pre>
onc.q <- lapply(onc.q,function(x) x[,7:ncol(x)])</pre>
                                          # pit
```

```
colnames(pit)[7:ncol(pit)] <- substr(colnames(pit)[7:ncol(pit)],1,2)</pre>
pit.q <- split(pit,paste(pit[,1],pit[,2]))</pre>
pit.q <- lapply(pit.q,function(x) x[,7:ncol(x)])</pre>
                                            # Get genotype
onc.geno <- unlist(sapply(names(onc.q),function(x) strsplit(x,split=' ')[[1]][2]))</pre>
pit.geno <- unlist(sapply(names(pit.q),function(x) strsplit(x,split=' ')[[1]][2]))</pre>
                                           # Xgal size data
xgs \leftarrow xgal.size[-1:-4, -(ncol(xgal.size) - 1):-ncol(xgal.size)]
xgs.cols <- xgal.size[4, -(ncol(xgal.size) - 1):-ncol(xgal.size)]</pre>
colnames(xgs) <- gsub("\\#", "", as.character(unlist(xgs.cols)))</pre>
xgs <- xgs[, 1:13]
xgs <- apply(xgs, 2, gsub, pattern = "\\,", replacement = "")</pre>
xgs.dim <- xgs[, "Measurement"]</pre>
xgs.geno <- xgs[, "Genotype"]</pre>
xgs.tree <- xgs[, "Tree"]</pre>
xgs <- xgs[, grep("Thallus", colnames(xgs))]</pre>
                                            # fix qenotypes
                                            # t6
xgs.geno[grep("T6", xgs.geno)] <- "T6"
xgs.geno[grep("H10", xgs.geno)] <- "H-10"
                                           # Coercing to numeric
xgs <- apply(xgs, 2, as.numeric)</pre>
                                           # Dealing with NA values
xgs.geno <- xgs.geno[grep("Dimension", xgs.dim)]</pre>
xgs.tree <- xgs.tree[grep("Dimension", xgs.dim)]</pre>
xgs <- xgs[grep("Dimension", xgs.dim), ]</pre>
xgs.dim <- xgs.dim[grep("Dimension", xgs.dim)]</pre>
                                           # Convert to cm
xgs <- xgs * 0.1
xgs.ellipse <- pi * xgs[xgs.dim == "Dimension 1", ] * xgs[xgs.dim == "Dimension 2", ]</pre>
xgs.geno <- xgs.geno[xgs.dim == "Dimension 1"]</pre>
xgs.tree <- xgs.tree[xgs.dim == "Dimension 1"]</pre>
                                           # package all xgs related data
xgs.data <- data.frame(tree = xgs.tree, geno = xgs.geno,
                         mean.thallus = apply(xgs.ellipse, 1, mean, na.rm = TRUE),
                         median.thallus = apply(xgs.ellipse, 1, median, na.rm = TRUE),
                         xgs.ellipse)
                                            # remove trees not done (i.e. all NA)
xgs.data <- xgs.data[apply(xgs.data[, grep("Thallus", colnames(xgs.data))], 1, function(x) !(all(is.na(
                                            # Roughness in the Garden
rough <- read.csv('../data/lichen_networks/ONC_raw_roughness.csv')</pre>
                                            # Isolate roughness
rough <- rough[, 1:5]</pre>
                                            # Isolate north quadrats
rough <- rough[grepl("North", rough[,3]), ]</pre>
                                            # Average roughness
avg.rough <- tapply(rough[,5], rough[,1], mean)</pre>
r.tree <- names(avg.rough)</pre>
r.tree <- sub('-', '\\.', r.tree)
r.tree <- sub('\\.0', '\\.', r.tree)</pre>
names(avg.rough) <- r.tree</pre>
                                            # match roughness to to ses values
```

```
load('../data/lichen_networks/lcn_onc_ses.rda')
onc.ses <- unlist(os[,1])</pre>
onc.ses[is.na(onc.ses)] <- 0
names(onc.ses) <- rownames(os)</pre>
if (!(all(names(onc.ses) == names(onc.q)))){print('Holy crap!')}
ses.tree <- as.character(sapply(names(onc.ses),function(x) unlist(strsplit(x,split=' '))[1]))</pre>
onc.rough <- avg.rough[match(ses.tree, r.tree)]</pre>
if (!(all(ses.tree == names(onc.rough)))){print('Holy Crap!')}
                                           #RFLP distance values from Zink from Martinsen
rflp.d <- readLines('.../data/acn/rflp_queller_goodnight.txt')</pre>
rflp.d <- lapply(rflp.d, strsplit,split='\t')</pre>
rflp.d <- lapply(rflp.d, function(x) x[[1]])</pre>
rflp.d[[61]] <- c(rflp.d[[61]], "")
rflp.d <- do.call(rbind, rflp.d)
rflp.n \leftarrow rflp.d[1, -1]
rflp.d \leftarrow rflp.d[-1, -1]
diag(rflp.d) <- 1</pre>
rflp.d <- matrix(as.numeric(rflp.d),nrow=nrow(rflp.d))</pre>
rownames(rflp.d) <- colnames(rflp.d) <- rflp.n
rflp.d <- rflp.d[rownames(rflp.d) %in% unique(onc.geno),
                  colnames(rflp.d) %in% unique(onc.geno)]
rflp.d <- rflp.d[match(unique(onc.geno), rownames(rflp.d)),</pre>
                  match(unique(onc.geno),rownames(rflp.d))]
if (!(all(rownames(rflp.d) == unique(onc.geno)))){
    print('Holy crap, rflp.d names match error')
}
## Duplicate by genotype? Would this work to make the RFLPs values replicated?
                                           # Coerce to distance matrices
rflp.d <- as.dist(rflp.d)
                                           # Lichen Network Models
                                           # onc
cn.onc <- lapply(split(onc[, -1:-6], onc[, "Tree"]), coNet,</pre>
                  ci.p = 95, cond = TRUE)
cn.sign.onc <- lapply(split(onc[, -1:-6], onc[, "Tree"]), coNet,</pre>
                        ci.p = 95)
cn.d.onc <- netDist(cn.onc, method = "euclidean")</pre>
                                           # pit
cn.pit <- lapply(split(pit[, -1:-6], pit[, "Tree"]), coNet, ci.p = 95)</pre>
cn.sign.pit <- lapply(split(pit[, -1:-6], pit[, "Tree"]), coNet, ci.p = 95)</pre>
cn.d.pit <- netDist(cn.pit, method = "bc")</pre>
                                           # genotype means and mean distances
onc.tree <- do.call(rbind, strsplit(names(onc.geno), " "))[, 1]</pre>
cn.mu.onc <- list()</pre>
for (i in 1:length(unique(onc.geno))){
    cn.mu.onc[[i]] <- netMean(cn.onc[onc.geno == unique(onc.geno)[i]])</pre>
names(cn.mu.onc) <- unique(onc.geno)</pre>
cn.mu.d.onc <- netDist(cn.mu.onc, method = "bc")</pre>
                                           # mean bark roughness calculations
prb.mu.onc <- tapply(onc.rough, onc.geno, mean)</pre>
prb.mu.d.onc <- dist(prb.mu.onc)</pre>
                                           # network statistics
```

```
ns.onc <- lapply(lapply(cn.onc, function(x)</pre>
    abs(sign(x))), enaR:::structure.statistics)
ns.onc <- do.call(rbind, ns.onc)</pre>
                                           # Ratio P / N
ns.rpn <- unlist(lapply(cn.onc, function(x)</pre>
    mean(x[x > 0]) / mean(x[x < 0]))
                                           # modularity
cn.mod.onc <- matrix(nrow = length(cn.onc), ncol = 2)</pre>
for (i in 1:length(cn.onc)){
    if (sum(sign(cn.onc[[i]])) >= 3){
        ## Networks with modules = 2 9 14 19 20 25 27 28 30 31 42 44 54 57
        mod.tmp <- computeModules(cn.onc[[i]])</pre>
        cn.mod.onc[i, 1] <- slot(mod.tmp, "likelihood") ## module likelihood</pre>
        cn.mod.onc[i, 2] <- nrow(slot(mod.tmp, "modules")) - 1 ## number modules</pre>
    }else{cn.mod.onc[i] <- NA}</pre>
}
cn.mod.onc[is.na(cn.mod.onc)] <- 0</pre>
names(cn.mod.onc) <- c("mod.lik", "mod.n")</pre>
                                           # graph level centralization
dcen.onc <- unlist(lapply(cn.onc, function(x)</pre>
    sna::centralization(x, FUN = sna::degree, normalize = FALSE)))
onc.ns <- cbind(ns.onc, Cen = dcen.onc,
                 mod.lik = cn.mod.onc[, 1], mod.n = cn.mod.onc[, 2])
if (!(all(onc.tree == names(cn.onc)))){print("Danger Will Robinson!")}
                                           # species centralities
cen.spp <- lapply(cn.onc, sna::degree, rescale = FALSE)</pre>
cen.spp <- do.call(rbind, cen.spp)</pre>
colnames(cen.spp) <- colnames(cn.onc[[1]])</pre>
                                           # Community data
onc.com <- do.call(rbind,lapply(onc.q,function(x) apply(x,2,sum)))</pre>
onc.R <- apply(sign(onc.com),1,sum)</pre>
onc.H <- vegan::diversity(onc.com)</pre>
onc.com.gm <- apply(onc.com, 2, function(x, g) tapply(x, g, mean), g = onc.geno)
onc.com.gm.rel <- apply(onc.com.gm, 2, function(x) x/max(x))</pre>
onc.com.rel <- apply(onc.com, 2, function(x) x/max(x))</pre>
onc.com.rel <- cbind(onc.com.rel, ds = rep(min(onc.com.rel[onc.com.rel != 0]) / 1000, nrow(onc.com.rel)
onc.com <- cbind(onc.com, ds = rep(min(onc.com[onc.com != 0]) / 1000, nrow(onc.com)))
                                           # pit genotype mean community
pit.com <- do.call(rbind,lapply(pit.q,function(x) apply(x,2,sum)))</pre>
pit.com.gm <- apply(pit.com, 2, function(x, g) tapply(x, g, mean), g = pit.geno)
pit.com.gm.rel <- apply(pit.com.gm, 2, function(x) x/max(x))</pre>
pit.com.gm.rel[is.na(pit.com.gm.rel)] <- 0</pre>
                                           # Lichen community metrics
                                           # Percent Total Cover
ptc.onc <- unlist(lapply(onc.q, function(x) sum(apply(x, 1, function(x) sign(sum(x))))))</pre>
                                           # Species richness
spr.onc <- apply(onc.com[, colnames(onc.com) != "ds"], 1, function(x) sum(sign(x)))</pre>
                                           # Vectors for network similarity
## ns.vec.onc <- envfit(ord, data.frame(onc.ns[, c("L", "Cen")], R = onc.rough, Cov = ptc.onc))</pre>
## Creating distance matrices that match rflp
## this is for the "mean" distance matrices
```

```
cn.mu.d <- as.matrix(cn.mu.d.onc)</pre>
prb.mu.d <- as.matrix(prb.mu.d.onc)</pre>
prb.mu.d <- prb.mu.d[match(rownames(cn.mu.d), rownames(prb.mu.d)),</pre>
                      match(rownames(cn.mu.d), rownames(prb.mu.d))]
prb.mu.d <- as.dist(prb.mu.d)</pre>
onc.com.mu <- apply(onc.com[, -ncol(onc.com)], 2,</pre>
                     function(x, g) tapply(x, g, mean), g = onc.geno)
onc.com.mu <- onc.com.mu[match(rownames(cn.mu.d), rownames(onc.com.mu)), ]</pre>
onc.com.mu.d <- vegdist(onc.com.mu)</pre>
if (!(all(c(all(rownames(as.matrix(rflp.d)) == rownames(as.matrix(cn.mu.d.onc))),
            all(rownames(as.matrix(rflp.d)) == rownames(as.matrix(prb.mu.d))),
            all(rownames(as.matrix(rflp.d)) == rownames(as.matrix(onc.com.mu.d))))))){
    warning("Warning: distance matrices are not aligned.")
}else{
    print("Distance matrices good to go!")
                                          # Bipartite analysis
nperm <- 20
if (!(file.exists("../data/lichen_networks/nest_rel_onc.rda"))){
    nest.onc <- nestedness(onc.com.rel[, colnames(onc.com.rel) != "ds"], n.nulls = 999)</pre>
    dput(nest.onc, "../data/lichen_networks/nest_rel_onc.rda")
}else{
    nest.onc <- dget("../data/lichen_networks/nest_rel_onc.rda")</pre>
if (!(file.exists("../data/lichen networks/null mod onc.csv"))){
    obs.mod.onc <- bipartite::computeModules(onc.com.rel[, colnames(onc.com.rel) != "ds"])
    mods.onc <- tail(apply(slot(obs.mod.onc, "modules"), 2,</pre>
                            function(x) sum(sign(x[2:length(x)]) *
                                                  (1:(length(x) - 1))),
                      sum(dim(onc.com[, colnames(onc.com) != "ds"])))
    mods.onc <- list(sp = tail(mods.onc, ncol(onc.com[, colnames(onc.com) != "ds"])),</pre>
                      tree = head(mods.onc, nrow(onc.com)))
    sim.onc \leftarrow lapply(1:nperm, sim.com, x = onc.q)
    sim.onc <- lapply(sim.onc, function(x) x / max(x))</pre>
    nul.mod.onc <- lapply(sim.onc, function(x) bipartite::computeModules(x))</pre>
    nul.mod.onc <- unlist(lapply(nul.mod.onc, slot, "likelihood"))</pre>
    dput(mods.onc, "../data/lichen_networks/mod_list_onc.rda")
    write.csv(slot(obs.mod.onc, "likelihood"),
               "../data/lichen_networks/obs_mod_onc.csv",
              row.names = FALSE)
    write.csv(nul.mod.onc,
               "../data/lichen_networks/null_mod_onc.csv",
              row.names = FALSE)
n}else{
    obs.mod.onc <- read.csv("../data/lichen_networks/obs_mod_onc.csv")[1]</pre>
    nul.mod.onc <- read.csv("../data/lichen_networks/null_mod_onc.csv")[,1]</pre>
    z.mod.onc <- (obs.mod.onc - mean(nul.mod.onc)) / sd(nul.mod.onc)</pre>
    mods.onc <- dget("../data/lichen_networks/mod_list_onc.rda")</pre>
pval.mod.onc <- length(nul.mod.onc[nul.mod.onc >= obs.mod.onc]) / length(nul.mod.onc)
if (pval.mod.onc == 0){pval.mod.onc <- 1/nperm}</pre>
z.mod.onc <- (obs.mod.onc - mean(nul.mod.onc)) / sd(nul.mod.onc)
```

```
bp.mod.onc <- round(unlist(c(nperm = nperm, obs = obs.mod.onc,</pre>
                 mu.sim = mean(nul.mod.onc), sd.sim = sd(nul.mod.onc),
                 z = z.mod.onc, p.value = pval.mod.onc)), 5)
## NMDS ordinations
                                          # community ordination
if (!file.exists("../data/lichen_networks/onc_nmds.csv")){
    nms.info.onc <- capture.output(nms.onc <- nmds.min(nmds(</pre>
        vegdist(onc.com.rel), 2, 2)))
    write.csv(nms.onc, "../data/lichen_networks/onc_nmds.csv", row.names = FALSE)
    write.table(nms.info.onc,
                 "../data/lichen_networks/onc_nmds_info.txt",
                 col.names = FALSE, row.names = FALSE)
}else{nms.onc <- read.csv("../data/lichen_networks/onc_nmds.csv")}</pre>
                                          # Network ordination
if (!(file.exists("../data/lichen_networks/conet_nmds.csv"))){
    cn.nmds.stats.onc <- capture.output(cn.nms.onc <- nmds.min(nmds(cn.d.onc, 2, 2)))</pre>
    write.csv(cn.nms.onc, "../data/lichen_networks/conet_nmds.csv", row.names = FALSE)
    write.table(cn.nmds.stats.onc,
                 "../data/lichen_networks/conet_nmds_info.txt",
                 col.names = FALSE, row.names = FALSE)
}else{cn.nms.onc <- read.csv("../data/lichen_networks/conet_nmds.csv")}</pre>
                                          # Vector fitting
nv.onc <- envfit(cn.nms.onc, data.frame(onc.com[, colnames(onc.com) != 'ds'],</pre>
                                          R = onc.rough,
                                          C = onc.ns[, c("C")],
                                          A = ptc.onc))
cv.onc <- envfit(nms.onc, data.frame(onc.com[, colnames(onc.com) != 'ds'],</pre>
                                          R = onc.rough,
                                          C = onc.ns[, c("C")],
                                       A = ptc.onc))
                                          #qenotype means
omu <- apply(onc.com[,colnames(onc.com) != 'ds'], 2,</pre>
             function(x,g) tapply(x,g,mean),g=onc.geno)
oms <- tapply(onc.ses, onc.geno, mean)</pre>
oms.d <- dist(oms[match(rownames(as.matrix(rflp.d)),names(oms))])</pre>
                                          #bark roughness means
oprbmu <- tapply(onc.rough,onc.geno,mean)</pre>
oprbmu <- oprbmu[match(rownames(as.matrix(rflp.d)),names(oprbmu))]</pre>
                                          #get araujo coordinates
coord <- read.csv('../data/lichen_networks/lcn_coord_onc.csv')</pre>
rownames(coord) <- coord[,1]</pre>
coord <- coord[,-1]</pre>
                                          # packing into a dataframe
tree <- onc.geno
for (i in 1:length(unique(onc.geno))){
    tree[onc.geno == unique(onc.geno)[i]] <- 1:length(tree[onc.geno == unique(onc.geno)[i]])
tree <- factor(tree)</pre>
tree.id <- do.call(rbind, strsplit(names(ptc.onc), split = " "))[, 1]</pre>
                                          # add chemistry data
onc.nc <- read.csv("../data/lichen_networks/ONC_phytochem_NC.csv")</pre>
onc.tan <- read.csv("../data/lichen_networks/ONC_phytochem_tannin.csv")</pre>
```

```
onc.nc[, 1] <- as.character(paste0("N", gsub("-", "\\.", onc.nc[, 1])))
onc.tan[, 1] <- as.character(paste0("N", gsub("-", "\\.", onc.tan[, 1])))
                                          # rename headers
                                          # mass is in mq
colnames(onc.nc)[1:4] <- c("tree.id", "sample.mass", "N", "C")</pre>
colnames(onc.tan)[1] <- "tree.id"</pre>
colnames(onc.tan)[grep("X.CT", colnames(onc.tan))] <- "CT"</pre>
                                          # add C:N ratio
onc.nc$rCN <- onc.nc$N / onc.nc$C
                                          # remove trees not in onc.dat
                                          # makes missing samples NA
onc.dat <- data.frame(onc.dat,</pre>
                    C = onc.nc[match(onc.dat[, "tree.id"],
                         onc.nc[, "tree.id"]), "C"],
                    N = onc.nc[match(onc.dat[, "tree.id"],
                         onc.nc[, "tree.id"]), "N"],
                    CN = onc.nc[match(onc.dat[, "tree.id"],
                         onc.nc[, "tree.id"]), "rCN"],
                    CT = onc.tan[match(onc.dat[, "tree.id"],
                        onc.tan[, "tree.id"]), "CT"])
                                          # collect into a single df
onc.dat <- data.frame(tree.id, PC = ptc.onc, SR = spr.onc,</pre>
                       geno = factor(onc.geno), tree = tree,
                       BR = onc.rough, onc.ns[, c("L", "Cen")])
                                          # Plot calculations
pw.onc <- onc.com.rel[, colnames(onc.com.rel) != "ds"]</pre>
pw.onc <- pw.onc[order(apply(pw.onc, 1, sum), decreasing = TRUE),</pre>
                  order(apply(pw.onc, 2, sum), decreasing = TRUE)]
rownames(pw.onc) <- onc.geno</pre>
col.pal <- RColorBrewer::brewer.pal((max(unlist(mods.onc))), "Paired")</pre>
                                          # Figure ordinations
                                          # Communities
if (file.exists("../data/lichen_networks/nms_com_onc.rda")){
    nms.com <- dget(file = "../data/lichen_networks/nms_com_onc.rda")</pre>
}else{
    set.seed(12345)
    nms.com <- nmds(vegdist(onc.com.rel), 2, 3)</pre>
    dput(nms.com, file = "../data/lichen_networks/nms_com_onc.rda")
}
                                          # Networks
if (file.exists("../data/lichen_networks/nms_cn_onc.rda")){
    nms.cn <- dget(file = "../data/lichen_networks/nms_cn_onc.rda")</pre>
}else{
    set.seed(12345)
    nms.cn <- nmds(cn.d.onc, 1, 2)
    dput(nms.cn, file = "../data/lichen_networks/nms_cn_onc.rda")
ord.com <- nmds.min(nms.com, 3)
ord.cn <- nmds.min(nms.cn, 2)
                                          # Vectors for plotting
                                          # Composition
vec.env <- onc.dat[, c("BR", "PC", "SR")]</pre>
colnames(vec.env) <- c("BR", "PC", "SR")</pre>
```

```
vec.com.12 <- envfit(ord.com, env = vec.env, perm = 10000,</pre>
                   choices = c(1,2))
                                          # Network similarity
vec.cn <- envfit(ord.cn, env = vec.env, perm = 10000,</pre>
                   choices = c(1,2))
                                          # onc
if (!("mod_obsval_onc.csv" %in% dir("../data/lichen_networks"))){
        mod.onc <- slot(bipartite::computeModules(rel(onc.com[, -ncol(onc.com)]),</pre>
                                                     deep = FALSE),
                           "likelihood")
        write.csv(mod.onc, file = "../data/mod_obsval_onc.csv", row.names = FALSE)
}else{
    mod.onc <- read.csv(file = "../data/lichen_networks/mod_obsval_onc.csv")[, 1]</pre>
if (!("mod_simvals_onc.csv" %in% dir("../data/lichen_networks"))){
        onc.sweb <- simulate(vegan::nullmodel(onc.com[, -ncol(onc.com)],</pre>
                                                 method = "swsh_samp_c"), 99)
        for (i in 1:dim(onc.sweb)[3]){onc.sweb[,, i] <- rel(onc.sweb[,, i])}</pre>
        onc.smod <- apply(onc.sweb, 3, bipartite::computeModules)</pre>
        mods.onc.sweb <- unlist(lapply(onc.smod, slot, name = "likelihood"))</pre>
        write.csv(mods.onc.sweb, file = "../data/lichen_networks/mod_simvals_onc.csv", row.names = FALS
# nest.onc <- bipartite::nestedness(onc.com.rel)</pre>
}else{
    mods.onc.sweb <- read.csv(file = "../data/lichen_networks/mod_simvals_onc.csv")[, 1]</pre>
                                          # pit
if (!("mod_obsval_pit.csv" %in% dir("../data/lichen_networks"))){
        mod.pit <- slot(bipartite::computeModules(rel(pit.com), deep = FALSE),</pre>
                           "likelihood")
        write.csv(mod.pit, file = "../data/lichen_networks/mod_obsval_pit.csv", row.names = FALSE)
}else{
    mod.pit <- read.csv(file = "../data/lichen_networks/mod_obsval_pit.csv")[, 1]</pre>
}
if (!("mod_simvals_pit.csv" %in% dir("../data/lichen_networks"))){
        pit.sweb <- simulate(vegan::nullmodel(pit.com, method = "swsh_samp_c"), 99)</pre>
for (i in 1:dim(pit.sweb)[3]){pit.sweb[,, i] <- rel(pit.sweb[,, i])}</pre>
        pit.smod <- apply(pit.sweb, 3, bipartite::computeModules)</pre>
        mods.pit.sweb <- unlist(lapply(pit.smod, slot, name = "likelihood"))</pre>
        write.csv(mods.pit.sweb, file = "../data/lichen_networks/mod_simvals_pit.csv", row.names = FALS
# nest.pit <- bipartite::nestedness(pit.com.rel)</pre>
}else{
    mods.pit.sweb <- read.csv(file = "../data/lichen_networks/mod_simvals_pit.csv")[, 1]</pre>
}
### Wild data
###
x <- read.csv('../data/lichen_networks/lco_Apr2012.csv')
                                          #remove notes
x <- x[,colnames(x)!='NOTES.']</pre>
x <- x[,colnames(x)!='dead']</pre>
```

```
x <- na.omit(x)
                                           \#remove\ qnu.44 = FREMONT
x \leftarrow x[x$tree!='gnu.44',]
                                           #remove 11.6, weird tree with super smooth bark
x <- x[x$tree!='11.6',]
x$tree <- factor(as.character(x$tree))</pre>
                                           #condense species
                                           #lecanora, there can be only one!
lec.sp <- apply(x[,c(6,8,10,18)],1,function(x) sign(any(x!=0)))
                                           #no physcioids!
                                           \#phy.spp \leftarrow apply(x[,c(13,14,15,16)],1,function(x) sign(any(x!=
x <- cbind(x,lec=lec.sp)</pre>
x \leftarrow x[,-c(6,8,10,18)]
x <- x[,colnames(x)!='physcioid']</pre>
                                           #break into quadrat list (x,q)
quads <- paste(x$tree,x$quadrat)</pre>
colnames(x)[5:ncol(x)] <- c('Xg','Cs', 'Xm', 'fgb', 'Rs', 'Pm', 'Pa', 'Pu','Ch','Ls')
x <- x[colnames(x)!='fgb']</pre>
x.q <- split(x,quads)</pre>
wild.com <- split(x,x$tree)</pre>
wild.com <- do.call(rbind,lapply(wild.com,function(x) apply(x[,-1:-4],2,sum)))</pre>
wild.com.rel <- apply(wild.com, 2, function(x) x/max(x))</pre>
wild.com.rel[is.na(wild.com.rel)] <- 0</pre>
wild.q <- lapply(split(x,x$tree),function(x) x[,-1:-4])</pre>
                                           #data from lamit
env <- read.csv('.../data/lichen_networks/Uinta2012_all_data_from_Lamit.csv')</pre>
env <- env[is.na(env$Pct.Roughness) == FALSE,]</pre>
env[,1] <- sub('\\?','',sub('\\_','\\.',sub('\\-','\\.',tolower(as.character(env[,1]))</pre>
env[env[,1] == 'll.6_(znu.29)',1] <- 'll.6'
env[env[,1] == 'gnu.85.1ftaway',1] <- 'gnu.85'</pre>
env$Quad.Loc <- as.character(sapply(as.character(env$Quad.Loc),function(x) unlist(strsplit(x,split='_')</pre>
env$Quad.Loc <- sub('\\-','\\.',env$Quad.Loc)</pre>
env$Quad.Loc <- paste('n',env$Quad.Loc,sep='')</pre>
                                           #remove southern aspect
env <- env[env$Aspect!='South',]</pre>
env.tid <- paste(env$Tree.ID,env$Quad.Loc)</pre>
                                           #check that the datasets are compatible
all(names(x.q)%in%env.tid)
                                           #match observations
all(env.tid[match(names(x.q),env.tid)] == names(x.q))
                                           #delimit to co-occurrence dataset and match
env <- env[match(names(x.q),env.tid),]</pre>
x.split <- paste(x$tree,x$quadrat,sep='_')</pre>
env.split <- paste(env$Tree.ID,env$Quad.Loc)</pre>
x.split <- as.character(x$tree)</pre>
env.split <- as.character(env$Tree.ID)</pre>
prb.wild <- tapply(env$Pct.Roughness,env.split,mean) #percent rough bark
                                           #age
age <- read.csv('.../data/lichen_networks/UintaMaster_LichenHeritNL_FallSpring_2012_ForLau.csv')
dbh <- age$DBH.cm_01
age.final <- age$AgeFinal.U
age <- data.frame(tree.id=age[,1],age.final=age$AgeFinal.U)</pre>
age[,1] <- tolower(age[,1])
```

```
age[,1] <- sub('_','\\.',age[,1])
age[,1] <- sub('-','\\.',age[,1])
age[,1] <- sub('\\?','',age[,1])
age[,1] <- sub('\\.0','\\.',age[,1])
age[age[,1] == 'gnu.85.1ftaway',1] <- 'gnu.85'
gnu19.dbh <- dbh[age$tree.id == 'gnu.19']</pre>
new <- data.frame(dbh=seq(min(dbh),max(dbh),by=0.1))</pre>
age.final <- na.omit(age.final)</pre>
pred.age <- predict(lm(age.final~dbh,data=age),new)</pre>
plot(pred.age~new[,1])
gnu19.age <- as.numeric(pred.age[new[,1] == gnu19.dbh])</pre>
tree.age <- age[match(names(prb.wild),age[,1]),2]</pre>
tree.age[is.na(tree.age)] <- gnu19.age</pre>
names(tree.age) <- age[match(names(prb.wild),age[,1]),1]</pre>
age <- tree.age
                                           # networks
cn.wild <- lapply(wild.q, coNet)</pre>
cn.mu.wild <- netMean(cn.wild)</pre>
cn.d.wild <- netDist(cn.wild, method = 'bc')</pre>
                                           #co-occurrence patterns
wco <- do.call(rbind,lapply(wild.q,function(x,t) apply(CoCo(x,type=t),2,sum),t='pos'))</pre>
wch <- do.call(rbind,lapply(wild.q,function(x,t) apply(CoCo(x,type=t),2,sum),t='neg'))</pre>
                                           #get ses values
                                           #"z" "means" "pval" "simulated" "method" "statistic" "alternati
                                           \#ws \leftarrow lapply(wild.q,function(x) oecosimu(x,cs,method='r1',burn
                                           #wses <- unlist(lapply(ws,function(x) x$oecosimu[[1]]))</pre>
                                           #wsmu <- unlist(lapply(ws,function(x) x$oecosimu[[2]]))</pre>
                                           #wsp <- unlist(lapply(ws,function(x) x$oecosimu[[3]]))</pre>
                                           #wsim <- do.call(rbind, lapply(ws, function(x) x$oecosimu[[4]]))</pre>
                                           #rownames(wsim) <- names(wild.q)</pre>
                                           #ws <- cbind(wses,wsmu,wsp)</pre>
                                           #write.csv(ws, file='.../data/wild_ses_21mar2014.csv')
## Araujo Coordinate Values
coord <- read.csv('../data/lichen_networks/lcn_coord_onc.csv')</pre>
rownames(coord) <- coord[,1]</pre>
coord <- coord[,-1]</pre>
                                           # wild
if (!("mod_obsval_wild.csv" %in% dir("../data/lichen_networks"))){
        mod.wild <- slot(bipartite::computeModules(rel(wild.com), deep = FALSE),</pre>
                           "likelihood")
        write.csv(mod.wild, file = "../data/lichen_networks/mod_obsval_wild.csv", row.names = FALSE)
}else{
    mod.wild <- read.csv(file = "../data/lichen_networks/mod_obsval_wild.csv")[, 1]</pre>
if (!("mod_simvals_wild.csv" %in% dir("../data/lichen_networks"))){
        wild.sweb <- simulate(vegan::nullmodel(wild.com, method = "swsh_samp_c"), 99)
for (i in 1:dim(wild.sweb)[3]){wild.sweb[,, i] <- rel(wild.sweb[,, i])}</pre>
        wild.smod <- apply(wild.sweb, 3, bipartite::computeModules)</pre>
        mods.wild.sweb <- unlist(lapply(wild.smod, slot, name = "likelihood"))</pre>
        write.csv(mods.wild.sweb, file = "../data/lichen_networks/mod_simvals_wild.csv", row.names = FA
```

```
# nest.wild <- bipartite::nestedness(wild.com.rel)
}else{
    mods.wild.sweb <- read.csv(file = "../data/lichen_networks/mod_simvals_wild.csv")[, 1]
}
###Rename data objects for simplicity
ws <- read.csv('../data/lichen_networks/wild_ses_21mar2014.csv')</pre>
```