GradAnalysis.r

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```
rm(list=ls())
setwd('~/19. UBC 2016 Winter Term 2/STAT 550/Case/Code')
library('ggplot2') # for plotting
## Warning: package 'ggplot2' was built under R version 3.2.4
library('lme4') # for mixed-effects models
## Warning: package 'lme4' was built under R version 3.2.5
## Loading required package: Matrix
library('reshape2') # for ...
library('dplyr') # for %>%
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
       filter, lag
## The following objects are masked from 'package:base':
##
       intersect, setdiff, setequal, union
##
library('MASS') # for ...
##
## Attaching package: 'MASS'
## The following object is masked from 'package:dplyr':
##
##
       select
library('base') # for ...
# Import the MWD data.
soil <- read.table('MWD.csv', sep=',', header=T)</pre>
soil.unreleveled <- soil</pre>
soil$Treatment <- relevel(soil$Treatment, ref='con')</pre>
head(soil)
```

```
Sample Date Block Treatment Transect MWD
## 1
          1 April
                      b1
                                bio
                                            1 1.05
## 2
           2 April
                       b1
                                bio
                                            2 0.95
## 3
                                            3 0.99
           3 April
                                bio
                      b1
## 4
           4 April
                       b3
                                bio
                                            1 0.82
## 5
           5 April
                                            2 0.84
                      b3
                                bio
## 6
           6 April
                                            3 0.96
                      b3
                                bio
str(soil)
                     96 obs. of 6 variables:
## 'data.frame':
## $ Sample : int 1 2 3 4 5 6 7 8 9 10 ...
                : Factor w/ 4 levels "April", "Aug", ...: 1 1 1 1 1 1 1 1 1 1 ...
## $ Date
## $ Block
                : Factor w/ 4 levels "b1", "b2", "b3", ...: 1 1 1 3 3 3 2 2 2 4 ...
## $ Treatment: Factor w/ 2 levels "con", "bio": 2 2 2 2 2 2 2 2 2 ...
## $ Transect : int 1 2 3 1 2 3 1 2 3 1 ...
    $ MWD
                : num 1.05 0.95 0.99 0.82 0.84 0.96 1.11 1.25 1.11 1.23 ...
with(soil, table(Treatment,Date))
##
             Date
## Treatment April Aug June Oct
##
          con
               12 12
                           12 12
                 12 12
##
          bio
                           12 12
# Import the cover value data.
pc <- read.csv('plant_cover.csv', header=T)</pre>
pc$Block <- as.factor(pc$Block)</pre>
head(pc)
##
                           Project
                                              Date Block Treatment Transect Plot
                                                       1 Biosolids
## 1 OK Ranch Biosolids Resample June 21, 2016
                                                                            1
## 2 OK Ranch Biosolids Resample June 21, 2016
                                                       1 Biosolids
                                                                            1
                                                                                  2
## 3 OK Ranch Biosolids Resample June 21, 2016
                                                                                 2
                                                       1 Biosolids
                                                                            1
## 4 OK Ranch Biosolids Resample June 21, 2016 1 Biosolids ## 5 OK Ranch Biosolids Resample June 21, 2016 1 Biosolids ## 6 OK Ranch Biosolids Resample June 21, 2016 1 Biosolids
                                                                                 2
                                                                            1
     Species Cover.class Cover.value
## 1
        ACMI
                         1
## 2
         ALCE
                                    2.5
                         1
## 3
        ASAG
                         1
                                   2.5
## 4
        BIOS
                         2
                                   15.0
## 5
        HECO
                         4
                                   62.5
                         5
                                   85.0
## 6
        LITT
str(pc)
## 'data.frame':
                      2669 obs. of 9 variables:
## $ Project : Factor w/ 1 level "OK Ranch Biosolids Resample": 1 1 1 1 1 1 1 1 1 1 ...
## $ Date
                  : Factor w/ 2 levels "June 21, 2016",..: 1 1 1 1 1 1 1 1 1 1 ...
                  : Factor w/ 4 levels "1", "2", "3", "4": 1 1 1 1 1 1 1 1 1 1 ...
## $ Block
```

```
## $ Treatment : Factor w/ 2 levels "Biosolids", "Control": 1 1 1 1 1 1 1 1 1 1 ...
## $ Transect : int 1 1 1 1 1 1 1 1 1 ...
              : int 2 2 2 2 2 2 2 2 3 ...
## $ Species : Factor w/ 50 levels "ACMI", "ACRI", ...: 1 3 9 11 29 33 39 47 48 3 ...
## $ Cover.class: int 1 1 1 2 4 5 3 3 1 2 ...
## $ Cover.value: num 2.5 2.5 2.5 15 62.5 85 37.5 37.5 2.5 15 ...
with(pc, table(Treatment,Block))
##
             Block
## Treatment
               1 2 3 4
   Biosolids 301 301 239 267
    Control 420 406 353 382
##
tapply(pc$Cover.value, pc$Treatment, table)
## $Biosolids
##
## 2.5
        15 37.5 62.5
                        85 97.5
## 405 265 111 78 91 158
## $Control
##
## 2.5
                        85 97.5
        15 37.5 62.5
## 676 510 243
######################################
# Grad Analysis:
######################################
# Obtain the relevant species.
# Note: The client said they are interested in POPR, PSSP, POJU, HECO,
       ALCE, ANDI, TAOF, SOIL and BRYO, as well as anything with a cover
        greater than 5% across all the sites.
#
     : None of the species occurred in all of the 400 sites (4 blocks x
#
       2 treatments × 5 transects × 10 samples per transect), although HECO
#
        and LITT were close. The rest occurred in below half of the sites.
#
       So, we change our criterion to choose species whose averages over
        the (occurring) transects and plots, for every block-treatment
        combination, are all greater than 5%.
summary(pc$Species)
## ACMI ACRI ALCE ANDI ANMI ARCA ARFR ARHO ASAG ASMI BIOS BRTE BRYO CAFI CAMI
          6 134
                  81 103
                              1
                                  90
                                       12 187
                                                 11
## CAOB CAPE CASP CEAR COUM DEPI ELTR ERCO FECE FEOC GAAR GABO GETR HECO JUBA
                         5
                             10
                                   1
                                       19
                                            25
                                                  3
                                                       3
                                                                1 353
          1
               6
                    1
                                                           3
## KOMA LILE LITT LOMA ORLU OXSE PEPR POHI POJU POPR POSE PSSP ROAC SECA SIAL
          9 398
                            3
                                  3
                                      1 62 120
                                                     84 48
                    2 1
## SOIL TAOF TRDU TRPR ZIVE
## 186 107 60
                    5
```

```
##
          Block
## Species
            1
                 2
                     3
                         4
##
                        28
      ACMI
           17
                 6
                   15
##
      ACRI
             4
                 0
                     0
                         2
##
      ALCE 83
                46
                         1
                     4
##
      ANDI 16
                28
                    19
                        18
##
      ANMI 27
                37
                    28
                        11
##
      ARCA
            1
                 0
                     0
                         0
##
      ARFR 12
                27
                    18
                        33
##
      ARHO
             2
                7
                         1
                     2
##
      ASAG 50
                30
                    51
                        56
      ASMI 10
##
                 0
                     1
                         0
##
      BIOS
            6
                17
                     2
                        11
##
      BRTE
             0
                 0
                     0
                         2
##
      BRYO 50
                52
                    47
                        50
##
      CAFI
             0
                 0
                         0
                     1
##
      CAMI
                 0
                         3
             0
                     0
##
      CAOB
                 2
                     0
                         0
             0
##
      CAPE
             1
                 0
                     0
                         0
##
      CASP
                 0
                     5
                         1
             0
##
      CEAR
             0
                 0
                     1
                         0
##
      COUM
             5
                 0
                     0
                         0
##
      DEPI
             0
                 0
                    10
                         0
      ELTR
##
             0
                 0
                     1
                         0
##
      ERCO 16
                 0
                     0
                         3
##
      FECE
                23
                     0
                         2
             0
##
      FEOC
                 0
                     2
                         1
             0
      GAAR
##
             2
                 0
                     0
                         1
      GABO
                     1
                         0
##
             2
                 0
##
      GETR
                 0
                     1
                         0
             0
##
      HECO 89
                95
                    91
                        78
##
      JUBA
             0
                 0
                        17
                     4
##
      KOMA 37
                61
                    37
                        48
      LILE
##
             9
                 0
                         0
##
      LITT 100 100
                    99
                        99
##
      LOMA
             2
                 0
                     0
                         0
##
      ORLU
                 0
                     0
                         0
             1
##
      OXSE
                 0
                     2
                         0
             1
##
      PEPR
                     0
                         0
                 0
             3
##
      POHI
             0
                 0
                     0
                         1
##
      POJU 12
                41
                     4
                         5
##
      POPR 34
                14
                    30 42
      POSE 15
                28
                        34
##
                     7
##
      PSSP 21
                18
                         8
                     1
##
      ROAC
            0
                 0
                     0
                         5
##
      SECA
             0
                 0
                     0
                         1
##
                 2
      SIAL
                     0
                         0
            1
##
      SOIL 50 65 34 37
```

```
## TAOF 29 4 47 27
## TRDU 9 2 26 23
## TRPR 3 2 0 0
## ZIVE 1 0 1 0
```

with(pc, table(Species, Treatment))

##	-	[reatment	
##	Species	Biosolids	Control
##	ACMI	39	27
##	ACRI	2	4
##	ALCE	55	79
##	ANDI	0	81
##	ANMI	5	98
##	ARCA	0	1
##	ARFR	24	66
##	ARHO	1	11
##	ASAG	60	127
##	ASMI	1	10
##	BIOS	36	0
##	BRTE	2	0
##	BRYO	6	193
##	CAFI	1	0
##	CAMI	3	0
##	CAOB	0	2
##	CAPE	0	1
##	CASP	4	2
##	CEAR	0	1
##	COUM	0	5
##	DEPI	10	0
##	ELTR	1	0
##	ERCO	0	19
##	FECE	10	15
##	FEOC	0	3
##	GAAR	0	3
##	GABO GETR	3	0
##	HECO	160	193
##	JUBA	190	193
##	KOMA	38	145
##	LILE	0	9
##	LITT	199	199
##	LOMA	1	1
##	ORLU	0	1
##	OXSE	1	2
##	PEPR	0	3
##	POHI	1	0
##	POJU	53	9
##	POPR	115	5
##	POSE	13	71
##	PSSP	38	10
##	ROAC	5	0
##	SECA	0	1
##	SIAL	3	0

```
SOIL
                          135
##
                  51
##
      TAOF
                 101
                            6
      TRDU
##
                  43
                           17
                            2
##
      TRPR
                   3
##
      ZIVE
                   0
                            2
species.vec <- NULL</pre>
for (ii in 1:length(all.species))
  # Obtain the subset of the data corresponding to this species.
  species.try <- all.species[ii]</pre>
  pc.sub.try <- subset(pc, Species==species.try)</pre>
  pc.sub.try \leftarrow pc.sub.try[ ,c(3,4,5,6,9)]
  # Check if this species occurrs in every block-treatment combination.
  # Note: entries.try is the table of counts for each block, by treatment
          group, for this species.
  entries.try <- table(pc.sub.try$Block, pc.sub.try$Treatment)</pre>
  which.entries.miss.try <- which(entries.try==0, arr.ind=TRUE)
  is.in.all.combs <- ifelse(test=(nrow(which.entries.miss.try)==0), yes=1,
                             no=0)
  # Obtain averages over the (occurring) transects and plots, for every
  # (occurring) block-treatment combination.
  by_blockTrt.try <- group_by(pc.sub.try, Block, Treatment)</pre>
  #dat.avq.try <- summarise(by_blockTrt.try, y.avq=sum(Cover.value)/50)</pre>
  dat.avg.try2 <- summarise(by_blockTrt.try, y.avg=mean(Cover.value))</pre>
  print(species.try)
  print(dat.avg.try2)
  all.occurring.greater.than5 <- ifelse(test=(sum(dat.avg.try2$y.avg<5)==0),</pre>
                                         yes=1, no=0)
  all.greater.than5 <- is.in.all.combs * all.occurring.greater.than5
  # Determine whether the species should be included.
  if ((species.try%in%interesting) || (all.greater.than5))
    species.vec <- c(species.vec, species.try)</pre>
  }
}
## [1] "ACMI"
## Source: local data frame [7 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                           y.avg
##
     (fctr)
               (fctr)
                           (dbl)
         1 Biosolids 5.441176
## 1
## 2
          2 Biosolids 2.500000
## 3
          2 Control 2.500000
## 4
          3 Biosolids 8.750000
## 5
          3 Control 10.961538
          4 Biosolids 5.666667
## 6
## 7
              Control 2.500000
## [1] "ACRI"
```

```
## Source: local data frame [3 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          1 Biosolids 26.25
## 2
              Control 15.00
## 3
              Control 20.00
          4
## [1] "ALCE"
## Source: local data frame [7 x 3]
## Groups: Block [?]
##
      Block Treatment
##
                           y.avg
##
     (fctr)
               (fctr)
                           (dbl)
## 1
          1 Biosolids 21.060606
## 2
              Control 19.700000
## 3
          2 Biosolids 6.875000
## 4
              Control
                       4.903846
## 5
          3 Biosolids 2.500000
## 6
              Control 2.500000
## 7
          4
              Control 2.500000
## [1] "ANDI"
## Source: local data frame [4 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                          y.avg
##
     (fctr)
               (fctr)
                          (dbl)
## 1
              Control 2.500000
          1
## 2
          2
              Control 6.071429
## 3
              Control 3.157895
          3
## 4
          4
              Control 3.194444
## [1] "ANMI"
## Source: local data frame [7 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                          y.avg
##
     (fctr)
               (fctr)
                          (dbl)
## 1
          1 Biosolids 2.50000
## 2
          1
              Control 15.00000
## 3
          2 Biosolids 22.50000
## 4
          2
              Control 13.38235
## 5
          3
              Control 12.41071
## 6
          4 Biosolids 15.00000
## 7
              Control 8.75000
          4
## [1] "ARCA"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          1
              Control
                          15
## [1] "ARFR"
## Source: local data frame [8 x 3]
## Groups: Block [?]
```

```
##
##
     Block Treatment
                          y.avg
##
     (fctr)
               (fctr)
                           (dbl)
## 1
          1 Biosolids 14.375000
## 2
              Control 10.000000
          2 Biosolids 8.750000
## 3
## 4
              Control 4.705882
          3 Biosolids 2.500000
## 5
## 6
          3
              Control 11.029412
## 7
          4 Biosolids 11.666667
## 8
              Control 10.729167
## [1] "ARHO"
## Source: local data frame [5 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          1 Biosolids
                        2.5
## 2
              Control
                        2.5
          1
## 3
          2
              Control
                        2.5
## 4
          3
              Control
                        2.5
## 5
          4
              Control
                        2.5
## [1] "ASAG"
## Source: local data frame [8 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                          y.avg
##
     (fctr)
               (fctr)
                           (db1)
## 1
          1 Biosolids 11.300000
## 2
              Control 6.500000
          1
## 3
          2 Biosolids 3.636364
## 4
          2
              Control 3.815789
## 5
          3 Biosolids 11.250000
## 6
              Control 10.000000
          3
## 7
          4 Biosolids 7.857143
## 8
          4
              Control 5.119048
## [1] "ASMI"
## Source: local data frame [2 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          1
              Control 8.75
## 2
          3 Biosolids 2.50
## [1] "BIOS"
## Source: local data frame [4 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                         y.avg
##
     (fctr)
               (fctr)
                          (dbl)
## 1
          1 Biosolids 6.666667
## 2
          2 Biosolids 6.176471
## 3
          3 Biosolids 2.500000
## 4
          4 Biosolids 7.045455
```

```
## [1] "BRTE"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
          4 Biosolids 8.75
## [1] "BRYO"
## Source: local data frame [6 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                         y.avg
##
     (fctr)
               (fctr)
                          (dbl)
## 1
              Control 40.60000
          1
## 2
          2 Biosolids 8.75000
              Control 24.73958
## 3
          2
## 4
          3
              Control 33.13830
## 5
          4 Biosolids 26.25000
## 6
              Control 30.26042
## [1] "CAFI"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
     Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          3 Biosolids
                        2.5
## [1] "CAMI"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
     Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          4 Biosolids
                        2.5
## [1] "CAOB"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
     Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
              Control
          2
                        2.5
## [1] "CAPE"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
      Block Treatment y.avg
##
##
     (fctr)
               (fctr) (dbl)
## 1
              Control
          1
## [1] "CASP"
## Source: local data frame [3 x 3]
## Groups: Block [?]
##
##
     Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
         3 Biosolids 15.0
```

```
## 2
          3
              Control
                         2.5
## 3
          4
              Control
                         2.5
## [1] "CEAR"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
      Block Treatment y.avg
               (fctr) (dbl)
##
     (fctr)
## 1
          3
              Control 37.5
## [1] "COUM"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          1
              Control
## [1] "DEPI"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          3 Biosolids
## [1] "ELTR"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          3 Biosolids
## [1] "ERCO"
## Source: local data frame [2 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                         y.avg
##
     (fctr)
               (fctr)
                         (db1)
## 1
              Control 3.28125
## 2
          4
              Control 2.50000
## [1] "FECE"
## Source: local data frame [4 \times 3]
## Groups: Block [?]
##
##
      Block Treatment
                           y.avg
##
     (fctr)
               (fctr)
                           (dbl)
## 1
          2 Biosolids 6.388889
## 2
              Control 6.964286
          2
## 3
          4 Biosolids 2.500000
              Control 37.500000
          4
## [1] "FEOC"
## Source: local data frame [2 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
               (fctr) (dbl)
##
     (fctr)
```

```
3
              Control 8.75
## 2
          4
              Control 2.50
## [1] "GAAR"
## Source: local data frame [2 x 3]
## Groups: Block [?]
##
##
     Block Treatment y.avg
##
               (fctr) (dbl)
     (fctr)
## 1
          1
              Control
                        2.5
                        2.5
## 2
          4
              Control
## [1] "GABO"
## Source: local data frame [2 x 3]
## Groups: Block [?]
##
##
     Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          1 Biosolids 8.75
## 2
          3 Biosolids 37.50
## [1] "GETR"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
     Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          3 Biosolids
                        2.5
## [1] "HECO"
## Source: local data frame [8 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                         y.avg
##
     (fctr)
               (fctr)
                         (dbl)
## 1
          1 Biosolids 35.37500
## 2
          1
              Control 24.13265
## 3
          2 Biosolids 31.08696
          2 Control 29.48980
## 4
## 5
          3 Biosolids 58.39286
## 6
          3
              Control 44.23469
## 7
          4 Biosolids 42.34375
## 8
          4
              Control 30.05435
## [1] "JUBA"
## Source: local data frame [3 x 3]
## Groups: Block [?]
##
      Block Treatment
                          y.avg
     (fctr)
               (fctr)
                          (dbl)
## 1
          3 Biosolids 11.875000
## 2
          4 Biosolids 8.166667
## 3
          4
              Control 2.500000
## [1] "KOMA"
## Source: local data frame [8 x 3]
## Groups: Block [?]
##
##
     Block Treatment
                          y.avg
##
                          (dbl)
     (fctr)
               (fctr)
```

```
1 Biosolids 8.500000
              Control 7.129630
          1
## 3
          2 Biosolids 12.125000
## 4
              Control 9.451220
          2
## 5
          3 Biosolids 6.666667
## 6
              Control 8.308824
          3
## 7
          4 Biosolids 7.500000
              Control 12.093023
## 8
          4
## [1] "LILE"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
              Control
                        2.5
          1
## [1] "LITT"
## Source: local data frame [8 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                         y.avg
##
     (fctr)
               (fctr)
                          (dbl)
          1 Biosolids 93.70000
## 2
              Control 30.30000
          1
## 3
          2 Biosolids 82.85000
## 4
              Control 28.35000
          2
## 5
          3 Biosolids 94.23469
## 6
          3
              Control 46.30000
          4 Biosolids 86.15000
## 8
              Control 33.36735
          4
## [1] "LOMA"
## Source: local data frame [2 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          1 Biosolids
                        2.5
## 2
          1
              Control
                        2.5
## [1] "ORLU"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
               (fctr) (dbl)
     (fctr)
## 1
              Control
                        2.5
          1
## [1] "OXSE"
## Source: local data frame [2 x 3]
## Groups: Block [?]
##
      Block Treatment y.avg
##
##
     (fctr)
               (fctr) (dbl)
## 1
          1 Biosolids 85.00
## 2
          3
              Control 8.75
## [1] "PEPR"
## Source: local data frame [1 x 3]
```

```
## Groups: Block [?]
##
##
      Block Treatment
                         y.avg
                         (dbl)
##
     (fctr)
               (fctr)
## 1
          1
              Control 6.666667
## [1] "POHI"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
     (fctr)
               (fctr) (dbl)
## 1
         4 Biosolids
## [1] "POJU"
## Source: local data frame [6 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                         y.avg
##
     (fctr)
               (fctr)
                         (dbl)
## 1
          1 Biosolids 14.54545
## 2
              Control 15.00000
          1
## 3
          2 Biosolids 33.71622
## 4
              Control 8.75000
## 5
              Control 2.50000
          3
## 6
          4 Biosolids 12.50000
## [1] "POPR"
## Source: local data frame [7 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                         y.avg
     (fctr)
##
               (fctr)
                         (dbl)
## 1
          1 Biosolids 37.82258
## 2
          1
              Control 2.50000
## 3
          2 Biosolids 34.82143
## 4
          3 Biosolids 46.29310
              Control 2.50000
## 5
## 6
          4 Biosolids 52.92683
## 7
          4
              Control 15.00000
## [1] "POSE"
## Source: local data frame [6 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                          y.avg
##
     (fctr)
               (fctr)
                          (dbl)
## 1
              Control 4.166667
         1
## 2
          2 Biosolids 14.545455
## 3
          2
              Control 9.705882
## 4
          3
              Control 6.071429
## 5
          4 Biosolids 2.500000
## 6
          4
              Control 7.890625
## [1] "PSSP"
## Source: local data frame [7 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                          y.avg
```

```
(fctr)
               (fctr)
                           (dbl)
## 1
          1 Biosolids 44.264706
## 2
          1
              Control 11.875000
## 3
          2 Biosolids 21.666667
## 4
              Control 2.500000
## 5
          3 Biosolids 37.500000
## 6
          4 Biosolids 19.500000
              Control 6.666667
## 7
          4
## [1] "ROAC"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          4 Biosolids
                        9.5
## [1] "SECA"
## Source: local data frame [1 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
          4
              Control
## [1] "SIAL"
## Source: local data frame [2 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          1 Biosolids
                        2.5
## 2
          2 Biosolids
                        2.5
## [1] "SOIL"
## Source: local data frame [8 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                          y.avg
##
     (fctr)
               (fctr)
                           (db1)
          1 Biosolids 3.888889
## 1
## 2
          1
              Control 6.341463
## 3
          2 Biosolids 4.880952
## 4
              Control 11.022727
          2
## 5
          3 Biosolids 3.636364
## 6
          3
              Control 4.673913
## 7
          4 Biosolids 2.500000
## 8
              Control 12.314815
          4
## [1] "TAOF"
## Source: local data frame [7 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                           y.avg
##
     (fctr)
               (fctr)
                           (dbl)
## 1
          1 Biosolids 5.648148
## 2
          1
              Control 2.500000
## 3
          2 Biosolids 2.500000
## 4
          3 Biosolids 12.555556
```

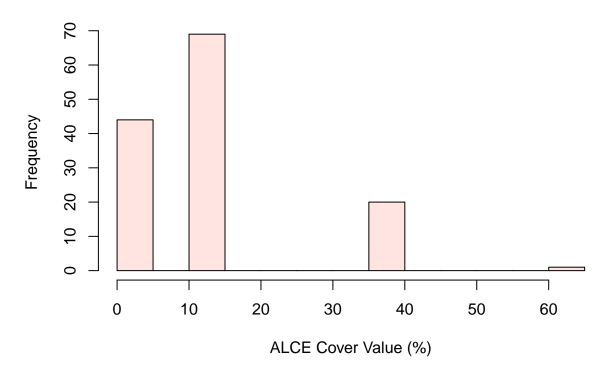
```
## 5
              Control 2.500000
          4 Biosolids 6.400000
## 7
              Control 2.500000
## [1] "TRDU"
## Source: local data frame [6 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                          y.avg
##
     (fctr)
               (fctr)
                          (dbl)
## 1
          1 Biosolids 2.500000
## 2
          2 Biosolids 2.500000
## 3
          3 Biosolids 8.815789
## 4
              Control 4.285714
## 5
          4 Biosolids 5.384615
          4
              Control 2.500000
## [1] "TRPR"
## Source: local data frame [2 x 3]
## Groups: Block [?]
##
##
      Block Treatment
                          y.avg
##
     (fctr)
               (fctr)
                          (dbl)
## 1
          1 Biosolids 10.83333
## 2
              Control 2.50000
          2
## [1] "ZIVE"
## Source: local data frame [2 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
          1
              Control 15.0
## 2
          3
              Control
                       2.5
print(species.vec)
## [1] "ALCE" "ANDI" "BRYO" "HECO" "KOMA" "LITT" "POJU" "POPR" "PSSP" "SOIL"
## [11] "TAOF"
# Obtain the MWD averages (to be used in calculating the correlations).
a <- group_by(soil.unreleveled, Block, Treatment) %>%
     summarise(mean(MWD))
b <- a mean (MWD)
mean.MWD <- b
# Loop over all the relevant species.
model.avg.list <-list()</pre>
p.vals.vec <- rep(NA, times=length(species.vec))</pre>
coefs.vec <- rep(NA, times=length(species.vec))</pre>
nonparam.model.list <-list()</pre>
nonparam.p.vals.vec <- rep(NA, times=length(species.vec))</pre>
nonparam.coefs.vec <- rep(NA, times=length(species.vec))</pre>
cor.vec <- rep(NA, times=length(species.vec))</pre>
for (ii in 1:length(species.vec))
```

```
# Obtain the subset of the data corresponding to this species.
this.species <- species.vec[ii]</pre>
pc.this.species <- subset(pc, Species==this.species)</pre>
pc.this.species <- pc.this.species[ ,c(3,4,9)]</pre>
print(this.species)
# Table of counts for each class of cover value, by treatment group, for
# this species.
#print(tapply(pc.this.species$Cover.value, pc.this.species$Treatment, table))
# Table of counts for each block, by treatment group, for this species.
entries <- table(pc.this.species$Block, pc.this.species$Treatment)</pre>
which.entries.miss <- which(t(entries)==0)</pre>
which.entries.miss.ind <- which(entries==0, arr.ind=TRUE)
# Obtain averages over the transects and plots.
by_blockTrt.this.species <- group_by(pc.this.species, Block, Treatment)
dat.avg.this.species <- summarise(by_blockTrt.this.species,</pre>
                                   y.avg=sum(Cover.value)/50)
if (length(which.entries.miss) > 0)
 for (jj in 1:length(which.entries.miss))
    which.row <- which.entries.miss[jj]</pre>
    which.block <- toString(which.entries.miss.ind[jj,1])</pre>
    which.trt <- ifelse(test=(which.entries.miss.ind[jj,2]==1), yes='Biosolids',
                         no='Control')
    new.row <- c(which.block, which.trt, 0)</pre>
    dat.avg.this.species <- rbind(dat.avg.this.species[1:(which.row-1),],</pre>
                                   new.row,
                                   dat.avg.this.species[-(1:(which.row-1)),])
 }
}
dat.avg.this.species$y.avg <- as.numeric(dat.avg.this.species$y.avg)</pre>
print(dat.avg.this.species)
############################
# Plots:
#############################
# Histogram of cover value for this species.
hist(pc.this.species$Cover.value,
     main=paste('Histogram of cover value for ',this.species,sep=''),
     xlab=paste(this.species,' Cover Value (%)',sep=''), col='mistyrose')
# Normal Q-Q plot of cover value for this species.
# Note: Strange behaviour due to discreteness.
#qqnorm(pc.this.species$Cover.value)
# Boxplots of cover value for this species, for the two treatments.
#print(qqplot(aes(y=Cover.value, x=Treatment, fill=Treatment, alpha=0.4),
              data=pc.this.species) +
       geom_boxplot() +
```

```
geom_point())
# Boxplot of cover value for this species, for the two treatments,
# across the four blocks.
#print(ggplot(aes(y=Cover.value, x=Treatment, fill=Treatment, alpha=0.4),
              data=pc.this.species) +
#
       geom_boxplot() +
#
       geom point() +
       facet wrap(~Block))
# Boxplot of average cover value for this species, for the two treatments.
print(ggplot(aes(y=y.avg, x=Treatment, fill=Treatment, alpha=0.4),
                 data=dat.avg.this.species) +
      geom_boxplot() +
      geom_point() +
      labs(y=paste(this.species,' Cover Value (%)',sep=''),
           title=paste('Boxplot of ',this.species,' Cover Value',sep='')))
# Interaction plot for block and treatment, with cover value for this
# species as the response.
#print(ggplot(aes(x=Block, y=Cover.value, group=Treatment,
                  colour=Treatment),
#
              data=pc.this.species) +
#
       stat_summary(fun.y='mean', geom='line') +
       labs(x='Block', y=paste(this.species,' Cover Value', sep=''),
#
            title=paste('Change in ',this.species,'
                         Cover Value over Different Blocks'), sep=''))
# Interaction plot for block and treatment, with average cover value for
# this species as the response.
#print(qqplot(aes(x=Block, y=y.avq, group=Treatment, colour=Treatment),
              data=dat.avq.this.species) +
#
       geom_point() +
#
       geom_line() +
       labs(x='Block', y=paste(this.species,' Cover Value (%)',sep=''),
            title=paste('Comparison of ',this.species,
#
                         ' Cover Values between Biosolids and Control',
                        sep='')))
############################
# Modelling cover value:
##############################
# Linear regression of average cover value for this species, with treatment
# effect.
dat.avg.this.species$Treatment <- relevel(dat.avg.this.species$Treatment,</pre>
                                           ref='Control')
model_avg.this.species <- lm(as.numeric(y.avg)~Treatment,</pre>
                              data=dat.avg.this.species)
print(summary(model_avg.this.species))
model.avg.list[[ii]] <- model_avg.this.species</pre>
p.vals.vec[ii] <- summary(model_avg.this.species)$coefficients[2,4]</pre>
coefs.vec[ii] <- summary(model_avg.this.species)$coefficients[2,1]</pre>
```

```
############################
  # Non-parametric test:
  ##############################
  # Parametric tests:
  \#t.test(x=dat.avg.this.species\$y.avg[dat.avg.this.species\$Treatment=='Biosolids'],
          y=dat.avg.this.species\$y.avg[dat.avg.this.species\$Treatment=='Control'],
          alternative='two.sided', mu=0, paired=FALSE, var.equal=TRUE,
          conf.level=0.95)
  \#t.test(x=dat.avq.this.species\$y.avq[dat.avq.this.species\$Treatment=='Biosolids'],
          y=dat.avg.this.species\$y.avg[dat.avg.this.species\$Treatment=='Control'],
          alternative='two.sided', mu=0, paired=TRUE, var.equal=TRUE,
          conf.level=0.95)
  # Non-parametric test:
  wilcox.this.species <- wilcox.test(x=dat.avg.this.species$y.avg[dat.avg.this.species$Treatment=='Bios
                                      y=dat.avg.this.species$y.avg[dat.avg.this.species$Treatment=='Cont.
                                      alternative='two.sided', mu=0, paired=TRUE,
                                      conf.int=TRUE, conf.level=0.95)
  nonparam.model.list[[ii]] <- wilcox.this.species</pre>
  nonparam.p.vals.vec[ii] <- wilcox.this.species$p.value</pre>
  nonparam.coefs.vec[ii] <- wilcox.this.species$estimate</pre>
  ############################
  # Correlations:
  ##########################
  # Find the correlation between MWD and the average cover value for this
  # species.
  mean.cv.this.species <- as.numeric(as.character(dat.avg.this.species$y.avg))</pre>
  cor.vec[ii] <- cor(mean.MWD, mean.cv.this.species)</pre>
}
## [1] "ALCE"
## Source: local data frame [8 x 3]
## Groups: Block [3]
##
##
      Block Treatment y.avg
##
     (fctr)
               (fctr) (dbl)
## 1
         1 Biosolids 13.90
## 2
          1 Control 19.70
          2 Biosolids 2.75
## 3
## 4
          2 Control 2.55
## 5
          3 Biosolids 0.10
         3 Control 0.10
## 6
## 7
         4 Biosolids 0.00
## 8
          4 Control 0.05
```

Histogram of cover value for ALCE



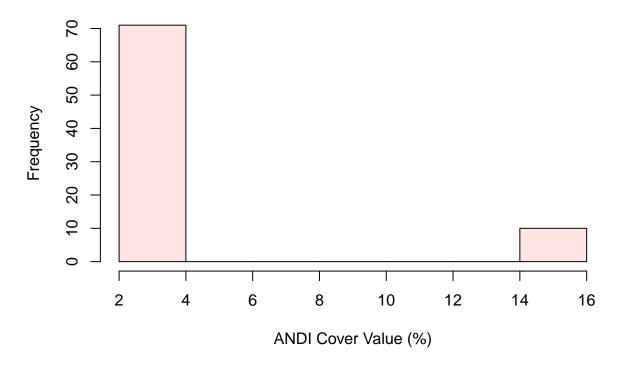
```
##
## Call:
## lm(formula = as.numeric(y.avg) ~ Treatment, data = dat.avg.this.species)
##
## Residuals:
##
     Min
              1Q Median
                            3Q
                                  Max
## -5.550 -4.516 -3.569 1.350 14.100
##
## Coefficients:
##
                      Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                         5.600
                                    4.081
                                            1.372
                                                     0.219
## TreatmentBiosolids
                       -1.413
                                    5.772 -0.245
                                                     0.815
## Residual standard error: 8.163 on 6 degrees of freedom
## Multiple R-squared: 0.009882,
                                    Adjusted R-squared: -0.1551
## F-statistic: 0.05988 on 1 and 6 DF, p-value: 0.8148
## Warning in wilcox.test.default(x = dat.avg.this.species
## $y.avg[dat.avg.this.species$Treatment == : requested conf.level not
## achievable
## Warning in wilcox.test.default(x = dat.avg.this.species
## $y.avg[dat.avg.this.species$Treatment == : cannot compute exact p-value
## with zeroes
## Warning in wilcox.test.default(x = dat.avg.this.species
## $y.avg[dat.avg.this.species$Treatment == : cannot compute exact confidence
## interval with zeroes
```

Boxplot of ALCE Cover Value Treatment Biosolids Control 0.4 0.4

Treatment

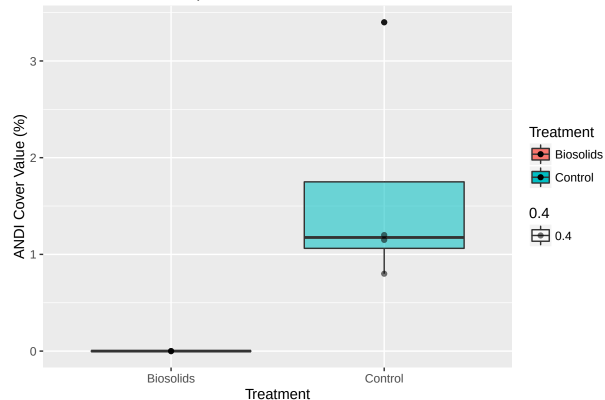
```
## [1] "ANDI"
## Source: local data frame [8 x 3]
## Groups: Block [3]
##
##
     Block Treatment y.avg
     (fctr)
               (fctr) (dbl)
##
         1
              Control 0.80
## 1
## 2
         1 Biosolids 0.00
          2 Biosolids 0.00
## 4
              Control 3.40
## 5
          3 Biosolids 0.00
## 6
              Control 1.20
          3
## 7
          4 Biosolids 0.00
             Control 1.15
## 8
```

Histogram of cover value for ANDI



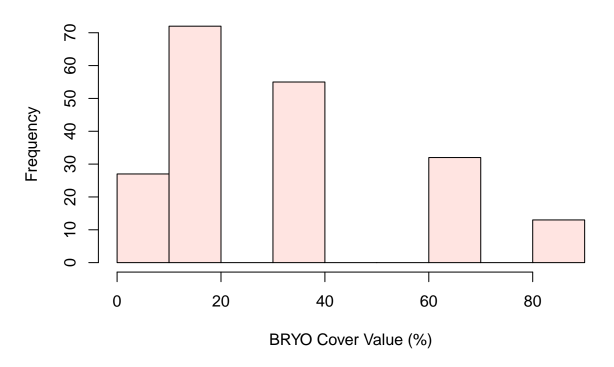
```
##
## Call:
## lm(formula = as.numeric(y.avg) ~ Treatment, data = dat.avg.this.species)
##
## Residuals:
##
      Min
               1Q Median
                               3Q
                                      Max
## -0.8375 -0.4500 0.0000 0.0000 1.7625
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
                                  0.4202
## (Intercept)
                        1.6375
                                           3.897 0.00801 **
## TreatmentBiosolids -1.6375
                                  0.5942 -2.756 0.03304 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.8403 on 6 degrees of freedom
## Multiple R-squared: 0.5586, Adjusted R-squared: 0.4851
## F-statistic: 7.594 on 1 and 6 DF, p-value: 0.03304
## Warning in wilcox.test.default(x = dat.avg.this.species
## $y.avg[dat.avg.this.species$Treatment == : requested conf.level not
## achievable
```

Boxplot of ANDI Cover Value

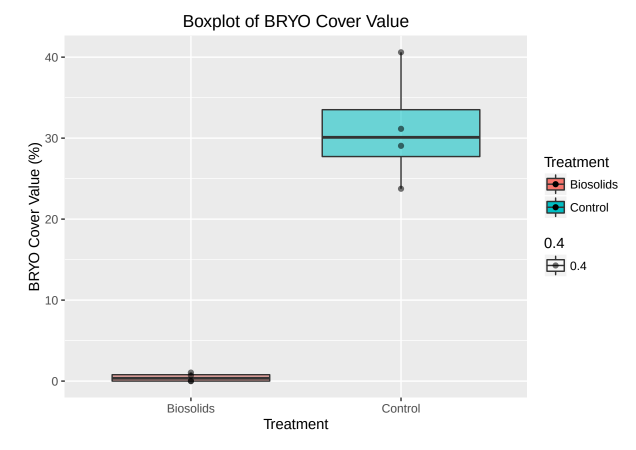


```
## [1] "BRYO"
## Source: local data frame [8 x 3]
## Groups: Block [2]
##
##
     Block Treatment y.avg
     (fctr)
               (fctr) (dbl)
##
         1
              Control 40.60
## 1
## 2
         1 Biosolids 0.00
          2 Biosolids 0.70
## 4
              Control 23.75
          3 Biosolids 0.00
## 5
## 6
              Control 31.15
## 7
          4 Biosolids 1.05
             Control 29.05
## 8
```

Histogram of cover value for BRYO

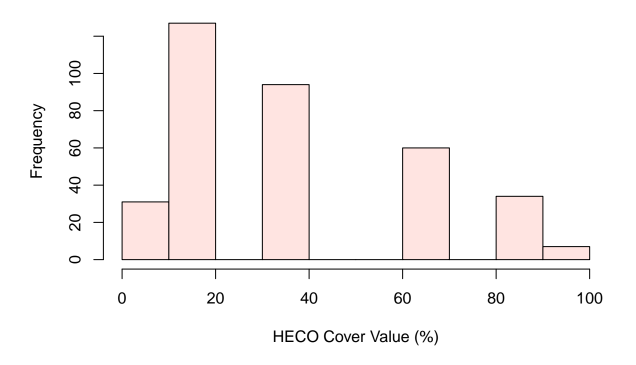


```
##
## Call:
## lm(formula = as.numeric(y.avg) ~ Treatment, data = dat.avg.this.species)
##
## Residuals:
##
      Min
               1Q Median
                               3Q
                                      Max
## -7.3875 -0.8500 -0.2125 0.3500 9.4625
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
                                   2.494 12.484 1.61e-05 ***
## (Intercept)
                        31.137
## TreatmentBiosolids -30.700
                                   3.527 -8.704 0.000127 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 4.988 on 6 degrees of freedom
## Multiple R-squared: 0.9266, Adjusted R-squared: 0.9144
## F-statistic: 75.75 on 1 and 6 DF, p-value: 0.0001271
## Warning in wilcox.test.default(x = dat.avg.this.species
## $y.avg[dat.avg.this.species$Treatment == : requested conf.level not
## achievable
```



```
## [1] "HECO"
## Source: local data frame [8 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
     (fctr)
               (fctr) (dbl)
##
          1 Biosolids 28.30
## 1
## 2
              Control 23.65
          1
          2 Biosolids 28.60
## 4
              Control 28.90
## 5
          3 Biosolids 49.05
## 6
              Control 43.35
## 7
          4 Biosolids 27.10
             Control 27.65
## 8
```

Histogram of cover value for HECO

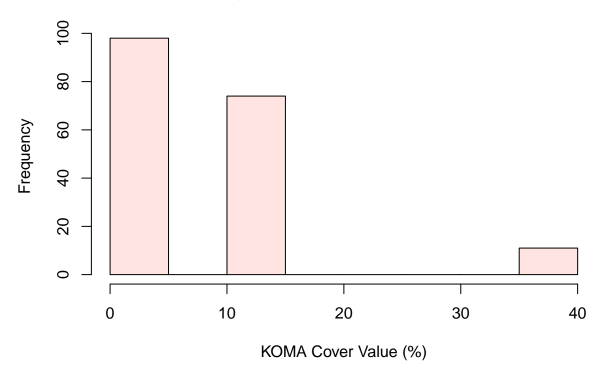


```
##
## Call:
## lm(formula = as.numeric(y.avg) ~ Treatment, data = dat.avg.this.species)
##
## Residuals:
     Min
             1Q Median
                           3Q
## -7.237 -5.263 -3.950 1.625 15.787
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
                                   4.812
## (Intercept)
                        30.887
                                           6.419 0.000675 ***
## TreatmentBiosolids
                        2.375
                                   6.805
                                           0.349 0.739009
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 9.624 on 6 degrees of freedom
## Multiple R-squared: 0.0199, Adjusted R-squared: -0.1435
## F-statistic: 0.1218 on 1 and 6 DF, p-value: 0.739
## Warning in wilcox.test.default(x = dat.avg.this.species
## $y.avg[dat.avg.this.species$Treatment == : requested conf.level not
## achievable
```

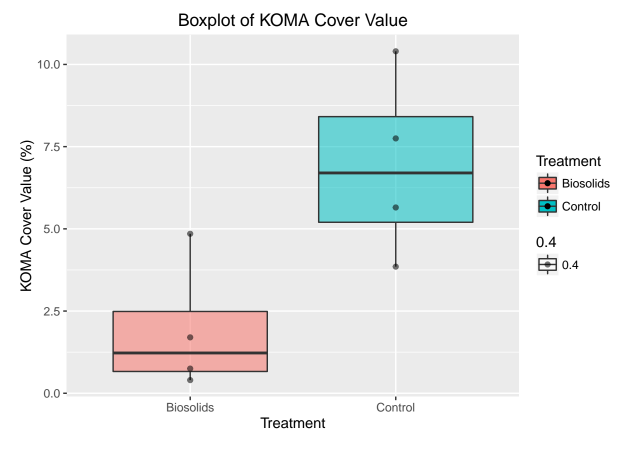
Boxplot of HECO Cover Value Treatment Biosolids Control Treatment O.4 10 O.4 10 Treatment

```
## [1] "KOMA"
## Source: local data frame [8 x 3]
## Groups: Block [?]
##
##
     Block Treatment y.avg
     (fctr)
               (fctr) (dbl)
##
         1 Biosolids 1.70
## 1
## 2
             Control 3.85
         1
         2 Biosolids 4.85
## 4
             Control 7.75
## 5
         3 Biosolids 0.40
## 6
             Control 5.65
## 7
         4 Biosolids 0.75
         4 Control 10.40
## 8
```

Histogram of cover value for KOMA

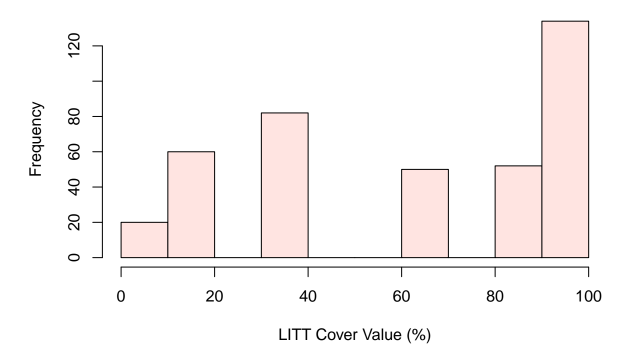


```
##
## Call:
## lm(formula = as.numeric(y.avg) ~ Treatment, data = dat.avg.this.species)
##
## Residuals:
##
      Min
              1Q Median
                            3Q
                                 Max
## -3.062 -1.328 -0.700 1.359
                               3.487
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
                                    1.227
## (Intercept)
                         6.913
                                           5.632 0.00134 **
## TreatmentBiosolids
                       -4.988
                                    1.736 -2.874 0.02829 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 2.455 on 6 degrees of freedom
## Multiple R-squared: 0.5792, Adjusted R-squared: 0.509
## F-statistic: 8.257 on 1 and 6 DF, p-value: 0.02829
## Warning in wilcox.test.default(x = dat.avg.this.species
## $y.avg[dat.avg.this.species$Treatment == : requested conf.level not
## achievable
```



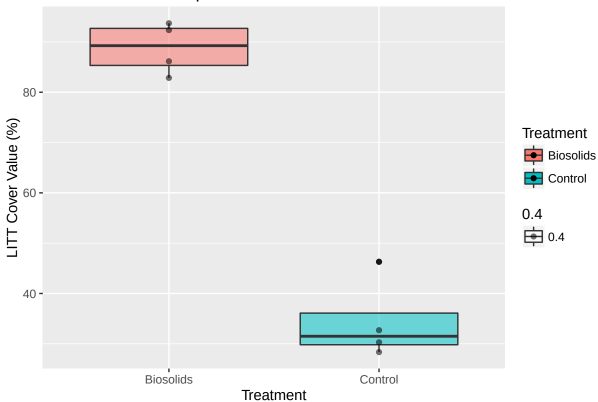
```
## [1] "LITT"
## Source: local data frame [8 x 3]
## Groups: Block [?]
##
##
      Block Treatment y.avg
     (fctr)
               (fctr) (dbl)
##
          1 Biosolids 93.70
## 1
## 2
              Control 30.30
          1
          2 Biosolids 82.85
## 4
              Control 28.35
          3 Biosolids 92.35
## 5
## 6
              Control 46.30
## 7
          4 Biosolids 86.15
              Control 32.70
## 8
```

Histogram of cover value for LITT



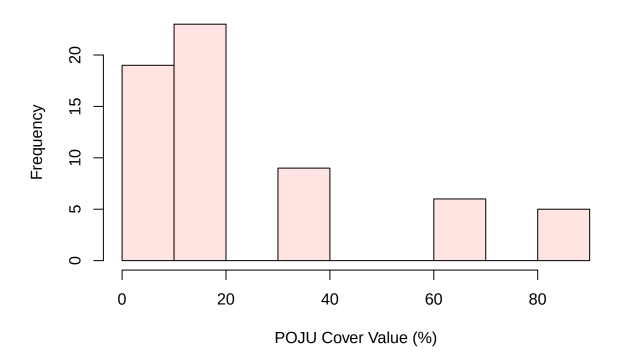
```
##
## Call:
## lm(formula = as.numeric(y.avg) ~ Treatment, data = dat.avg.this.species)
##
## Residuals:
##
      Min
              1Q Median
                            3Q
                                 Max
## -6.062 -4.562 -2.163 3.925 11.887
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
                                    3.397
                                            10.13 5.38e-05 ***
## (Intercept)
                        34.413
## TreatmentBiosolids
                        54.350
                                   4.804
                                            11.31 2.85e-05 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 6.794 on 6 degrees of freedom
## Multiple R-squared: 0.9552, Adjusted R-squared: 0.9478
## F-statistic:
                 128 on 1 and 6 DF, p-value: 2.854e-05
## Warning in wilcox.test.default(x = dat.avg.this.species
## $y.avg[dat.avg.this.species$Treatment == : requested conf.level not
## achievable
```

Boxplot of LITT Cover Value



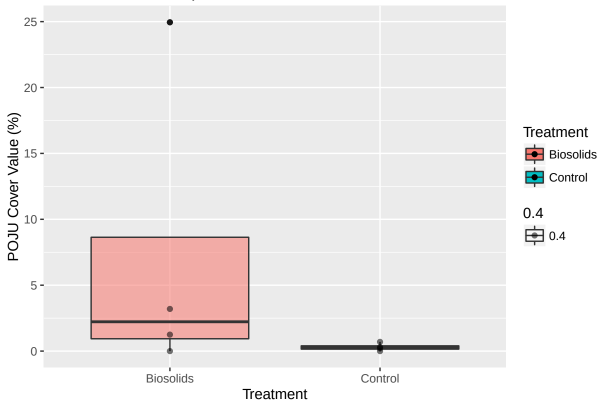
```
## [1] "POJU"
## Source: local data frame [8 x 3]
## Groups: Block [4]
##
##
     Block Treatment y.avg
     (fctr)
               (fctr) (dbl)
##
         1 Biosolids 3.20
## 1
## 2
             Control 0.30
         1
         2 Biosolids 24.95
## 4
             Control 0.70
## 5
          3 Biosolids 0.00
## 6
              Control 0.20
## 7
          4 Biosolids 1.25
             Control 0.00
## 8
```

Histogram of cover value for POJU



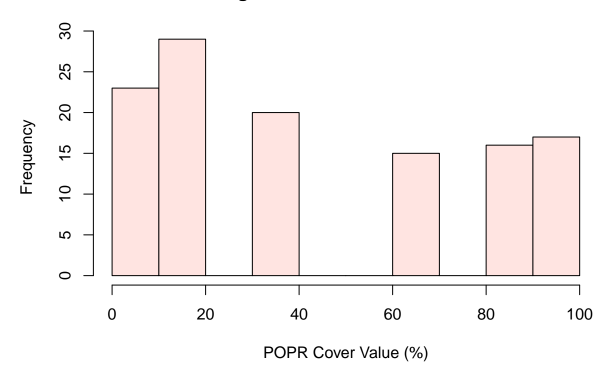
```
##
## Call:
## lm(formula = as.numeric(y.avg) ~ Treatment, data = dat.avg.this.species)
##
## Residuals:
##
     Min
              1Q Median
                            3Q
## -7.350 -4.638 -0.200 0.100 17.600
##
## Coefficients:
##
                      Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                         0.300
                                            0.072
                                                     0.945
                                    4.176
                         7.050
## TreatmentBiosolids
                                    5.905
                                            1.194
                                                     0.278
## Residual standard error: 8.351 on 6 degrees of freedom
## Multiple R-squared: 0.1919, Adjusted R-squared: 0.05727
## F-statistic: 1.425 on 1 and 6 DF, p-value: 0.2776
## Warning in wilcox.test.default(x = dat.avg.this.species
## $y.avg[dat.avg.this.species$Treatment == : requested conf.level not
## achievable
```

Boxplot of POJU Cover Value



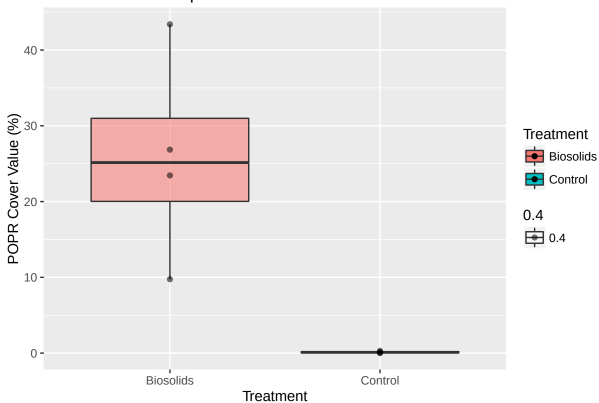
```
## [1] "POPR"
## Source: local data frame [8 x 3]
## Groups: Block [2]
##
##
     Block Treatment y.avg
     (fctr)
               (fctr) (dbl)
##
          1 Biosolids 23.45
## 1
## 2
              Control 0.15
          1
          2 Biosolids 9.75
## 4
              Control 0.00
## 5
          3 Biosolids 26.85
## 6
              Control 0.05
## 7
          4 Biosolids 43.40
             Control 0.30
## 8
```

Histogram of cover value for POPR



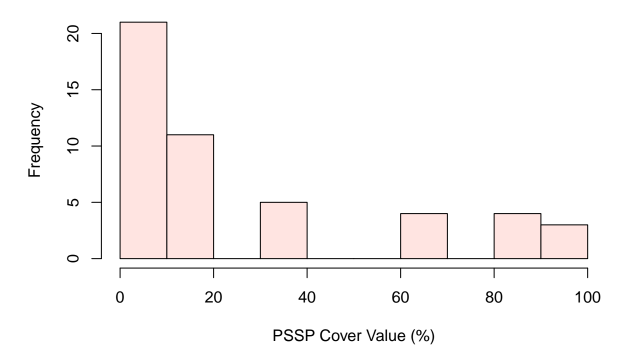
```
##
## Call:
## lm(formula = as.numeric(y.avg) ~ Treatment, data = dat.avg.this.species)
##
## Residuals:
##
       Min
                 1Q
                      Median
                                   3Q
## -16.1125 -0.6969 -0.0250
                               0.3781 17.5375
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
                                   4.891
## (Intercept)
                        0.125
                                           0.026 0.98044
## TreatmentBiosolids
                       25.738
                                   6.916
                                           3.721 0.00984 **
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 9.781 on 6 degrees of freedom
## Multiple R-squared: 0.6977, Adjusted R-squared: 0.6473
## F-statistic: 13.85 on 1 and 6 DF, p-value: 0.009838
## Warning in wilcox.test.default(x = dat.avg.this.species
## $y.avg[dat.avg.this.species$Treatment == : requested conf.level not
## achievable
```

Boxplot of POPR Cover Value



```
## [1] "PSSP"
## Source: local data frame [8 x 3]
## Groups: Block [3]
##
##
     Block Treatment y.avg
     (fctr)
               (fctr) (dbl)
##
         1 Biosolids 15.05
## 1
## 2
             Control 0.95
         1
         2 Biosolids 6.50
## 4
             Control 0.15
## 5
          3 Biosolids 0.75
## 6
             Control 0.00
## 7
          4 Biosolids 1.95
             Control 0.40
## 8
```

Histogram of cover value for PSSP

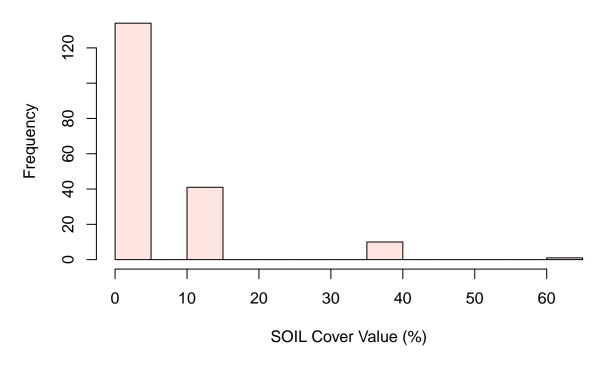


```
##
## Call:
## lm(formula = as.numeric(y.avg) ~ Treatment, data = dat.avg.this.species)
##
## Residuals:
##
       Min
                1Q Median
                                3Q
                                       Max
## -5.3125 -1.3094 -0.1000 0.4719
                                   8.9875
##
## Coefficients:
##
                      Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                         0.375
                                    2.297
                                                     0.876
                                            0.163
                         5.688
## TreatmentBiosolids
                                    3.248
                                            1.751
                                                     0.131
## Residual standard error: 4.594 on 6 degrees of freedom
## Multiple R-squared: 0.3382, Adjusted R-squared: 0.2278
## F-statistic: 3.066 on 1 and 6 DF, p-value: 0.1305
## Warning in wilcox.test.default(x = dat.avg.this.species
## $y.avg[dat.avg.this.species$Treatment == : requested conf.level not
## achievable
```

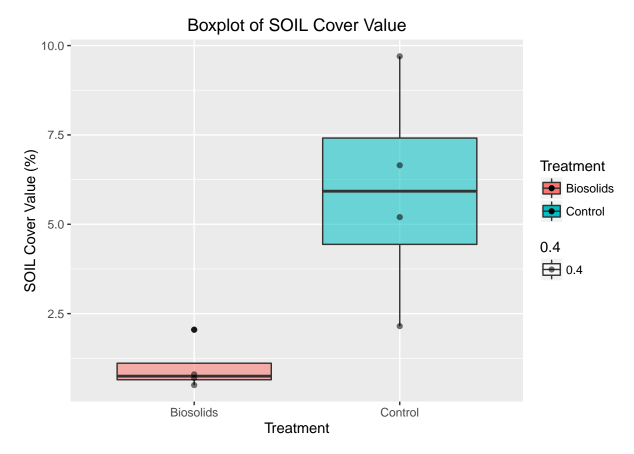
Boxplot of PSSP Cover Value Treatment Biosolids Control Treatment 0.4 15 0.4

```
## [1] "SOIL"
## Source: local data frame [8 x 3]
## Groups: Block [?]
##
##
     Block Treatment y.avg
     (fctr)
              (fctr) (dbl)
##
         1 Biosolids 0.70
## 1
## 2
             Control 5.20
         1
         2 Biosolids 2.05
## 4
             Control 9.70
## 5
         3 Biosolids 0.80
## 6
             Control 2.15
         3
## 7
         4 Biosolids 0.50
         4 Control 6.65
## 8
```

Histogram of cover value for SOIL

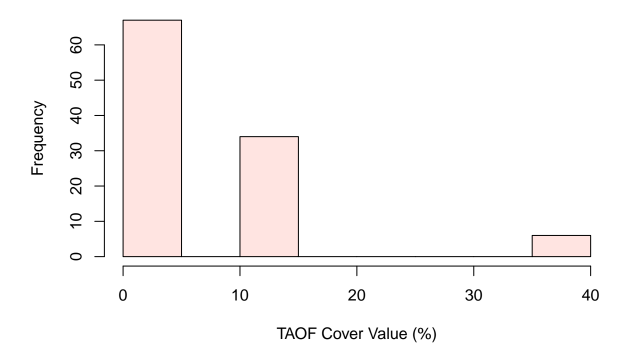


```
##
## Call:
## lm(formula = as.numeric(y.avg) ~ Treatment, data = dat.avg.this.species)
##
## Residuals:
##
      Min
               1Q Median
                               3Q
                                      Max
## -3.7750 -0.5656 -0.2625 0.8031 3.7750
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                        5.925
                                   1.137
                                           5.210 0.00199 **
## TreatmentBiosolids
                       -4.913
                                   1.608 -3.055 0.02238 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 2.274 on 6 degrees of freedom
## Multiple R-squared: 0.6086, Adjusted R-squared: 0.5434
## F-statistic: 9.331 on 1 and 6 DF, p-value: 0.02238
## Warning in wilcox.test.default(x = dat.avg.this.species
## $y.avg[dat.avg.this.species$Treatment == : requested conf.level not
## achievable
```



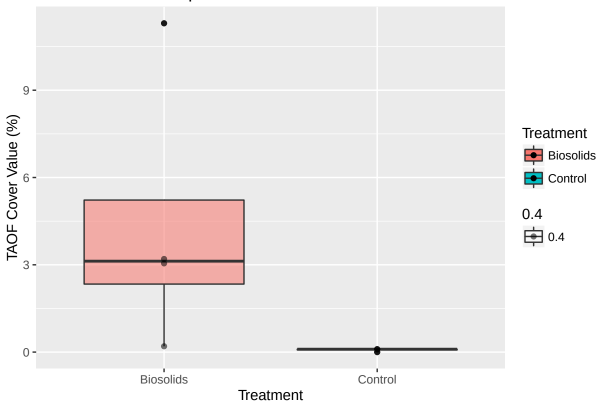
```
## [1] "TAOF"
## Source: local data frame [8 x 3]
## Groups: Block [2]
##
##
     Block Treatment y.avg
     (fctr)
               (fctr) (dbl)
##
         1 Biosolids 3.05
## 1
## 2
              Control 0.10
         1
          2 Biosolids 0.20
## 4
             Control 0.00
## 5
          3 Biosolids 11.30
## 6
              Control 0.10
## 7
          4 Biosolids 3.20
## 8
              Control 0.10
```

Histogram of cover value for TAOF



```
##
## Call:
## lm(formula = as.numeric(y.avg) ~ Treatment, data = dat.avg.this.species)
##
## Residuals:
##
     Min
              1Q Median
                            3Q
                                  Max
## -4.237 -1.275 -0.025 0.025
                                6.862
##
## Coefficients:
##
                      Estimate Std. Error t value Pr(>|t|)
                                                      0.966
## (Intercept)
                         0.075
                                    1.690
                                             0.044
## TreatmentBiosolids
                         4.362
                                    2.389
                                             1.826
                                                      0.118
## Residual standard error: 3.379 on 6 degrees of freedom
## Multiple R-squared: 0.3571, Adjusted R-squared:
## F-statistic: 3.333 on 1 and 6 DF, p-value: 0.1177
## Warning in wilcox.test.default(x = dat.avg.this.species
## $y.avg[dat.avg.this.species$Treatment == : requested conf.level not
## achievable
```

Boxplot of TAOF Cover Value



Print the linear regression p-values and coefficients for each species.

cbind(species.vec, p.vals.vec, coefs.vec)

```
##
         species.vec p.vals.vec
                                             coefs.vec
                                             "-1.4125"
##
    [1,] "ALCE"
                     "0.814834091762861"
                                             "-1.6375"
    [2,] "ANDI"
                     "0.0330367117716462"
##
                     "0.00012709185740304"
##
   [3,] "BRYO"
                                             "-30.7"
                     "0.739008763957111"
                                             "2.375"
##
   [4,] "HECO"
   [5,] "KOMA"
                     "0.0282943011554642"
                                             "-4.9875"
##
##
  [6,] "LITT"
                     "2.85417772499923e-05" "54.35"
                                             "7.05"
##
  [7,] "POJU"
                     "0.27759694571899"
  [8,] "POPR"
                     "0.00983752836594799"
                                             "25.7375"
##
##
   [9,] "PSSP"
                      "0.130533801218531"
                                             "5.6875"
## [10,] "SOIL"
                     "0.0223759429659092"
                                             "-4.9125"
## [11,] "TAOF"
                     "0.117683262573748"
                                             "4.3625"
```

Print the non-parametric p-values and estimates for each species. cbind(species.vec, nonparam.p.vals.vec, nonparam.coefs.vec)

```
##
         species.vec nonparam.p.vals.vec nonparam.coefs.vec
##
    [1,] "ALCE"
                      "0.789268026134281" "-2.29405540546612"
   [2,] "ANDI"
                      "0.125"
                                          "-1.1875"
##
##
    [3,] "BRYO"
                      "0.125"
                                          "-30.3625"
                      "0.625"
                                          "2.375"
##
   [4,] "HECO"
   [5,] "KOMA"
                      "0.125"
                                          "-4.6625"
    [6,] "LITT"
                                          "54.2375"
                      "0.125"
##
```

```
## [7,] "POJU" "0.25" "2.4875"

## [8,] "POPR" "0.125" "25.7375"

## [9,] "PSSP" "0.125" "5.15"

## [10,] "SOIL" "0.125" "-4.9125"

## [11,] "TAOF" "0.125" "3.0625"
```

Print the correlations between MWD and each species' cover value.
cbind(species.vec, cor.vec)

```
##
         species.vec cor.vec
## [1,] "ALCE"
                     "-0.0979609271355758"
## [2,] "ANDI"
                     "-0.408243567137178"
##
  [3,] "BRYO"
                     "-0.408826660600364"
## [4,] "HECO"
                     "-0.436327732101949"
## [5,] "KOMA"
                     "-0.375468054869239"
## [6,] "LITT"
                     "0.619357051173369"
## [7,] "POJU"
                     "0.615528658610572"
## [8,] "POPR"
                     "0.644724082434805"
## [9,] "PSSP"
                     "0.421947867236044"
## [10,] "SOIL"
                     "-0.405705525751975"
## [11,] "TAOF"
                     "0.0317200333125193"
```