Todo list

Describe experiment																											2	4
---------------------	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	---	---

EDUARD SZÖCS

QUANTITATIVE ECOTOXICOLOGY

WITH R!

This document was created using LATEX, knitr and the tufte book class.

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Contents

1	Introduction 9
2	The Measurement Process 11 2.1 Winsorized Mean and Standard Deviation 11 2.2 Probability Plotting 12
3	Bioaccumulation 13
4	Tests for Detection of Chronic Lethal and Sublethal Stress 15
5	Lethal and Other Quantal Responses to Stress 17 5.1 Fitting dose-response models 17
6	Population and Metapopulation Effects 19
7	Community Effects 21 7.1 Species Richness 21 7.2 Species Diversity 23 7.3 Dissimilarity indices 24 7.4 Ordination techniques - PCA, RDA, NMDS, MDS, dbRDA 24 7.5 Testing hypotheses - PERMANOVA 24
	7.6 Analyzing mesocosm data 24 7.7 Species Sensitivity Distributions 30

R Session Info 31

Bibliography 33

List of Figures

2.1	A histogramm of the so4 data. 11
-	LD50. Source: http://xkcd.com/1260/ 17 Proportion of fish died at different NaCl concentrations. 17
7.1	Species Richness along outfall 22
7.2	A) Rarefaction Curve. B) Rank abundance curve 22
7.3	Diversity indices along outfall 23
7.4	Principal response curves (PRC) of pyrifos data 26
7.5	Responses of <i>G. pulex</i> and <i>C. horaria</i> to chlorpyrifos. 26
7.6	PRC - Explained variance 28

List of Tables

7.1 One restricted permutation

28

1

Introduction

```
require(devtools)
install_github("qetx", "EDiLD")

require(qetx)
```

The Measurement Process

2.1 Winsorized Mean and Standard Deviation

Example data

The following sulfate concentrations (mg/L) were measured during a routine water quality survey of the Savannah River (South Carolina). The data is available in the qetx package ¹:

```
data(so4)
```

```
## [1] 1.3 2.3 2.6 3.3 3.5 3.5 3.6 4.0 4.1 4.5 5.2 5.6
## [13] 5.7 6.1 6.2 6.5 6.9 7.1 7.7 7.9 9.9

length(so4)
## [1] 21

mean(so4)
## [1] 5.119
sd(so4)
## [1] 2.137
```

So there are 21 measurements with a mean of 5.12 mg/L and a standard deviation of 2.14 mg/L.

Winsorization

Suppose we have a detection limit of 2.5 mg/L and want to winsorize values below LOD, i.e. replace the two lowest values by 2.6 mg/L and the two highest values by 7.7 mg/L.

¹ Note that in this case you do not have to assign the data to a name.

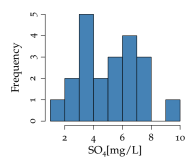


Figure 2.1: A histogramm of the so4 data.

Happily there is function in the qetx-package to do this for us: winsor(). This function takes a vector of values and a second argument specifying how many values should be winsorized (either by giving a LOD-value or the number of values on each side) ².

```
so4_w <- winsor(so4, lod = 2.5)
so4_w

## [1] 2.6 2.6 2.6 3.3 3.5 3.5 3.6 4.0 4.1 4.5 5.2 5.6
## [13] 5.7 6.1 6.2 6.9 6.5 7.1 7.7 7.7 7.7
## attr(,"width")
## [1] 2</pre>
```

This give the expected results, moreover we see that on each end two observations where modified ³.

```
mean(so4_w)
## [1] 5.081

sd(so4_w)
## [1] 1.792

sd_winsor(so4_w)
## [1] 2.24
```

The Winsorized mean (\bar{x}_w) now is 5.08 mg/L, the standard deviation of the modified data set (s) is 1.79 mg/L and the Winsorized standard deviation (s_w) 2.24 mg/L.

2.2 Probability Plotting

Look at the source of this function
 type the function name into the console - to see which computations are done.

³ stored within the attribute 'width' of the resulting vector. **TODO: verbatim within sidenote.**

3 Bioaccumulation

4
Tests for Detection of Chronic Lethal and Sublethal Stress

5

Lethal and Other Quantal Responses to Stress

5.1 Fitting dose-response models

Example data

Newman and Aplin (1992) exposed mosquitofish *Gambusia holbrooki* to a series of NaCl concentrations for 96h. The data is available from the qetx package.

```
data(salt)
str(salt)

## 'data.frame': 6 obs. of 3 variables:
## $ dead : int 16 22 40 69 78 77

## $ total: int 76 79 77 76 78 77

## $ conc : num 10.3 10.8 11.6 13.2 15.8 20.1
```

There are three columns:

dead Number of fish died.

total Total number of fish exposed.

cocn NaCl concentration (g/L).

First we calculate the proportion of died fish and safe it as a new column in our data.frame:

```
salt$prop <- salt$dead/salt$total
```

As always we first take a look at the data, to produce Fig. 5.2:

Introduction

1



THE LD₅₀ OF TOXICITY DATA IS 2 KILOGRAMS PER KILOGRAM.

Figure 5.1: LD50. Source: http://xkcd.com/1260/

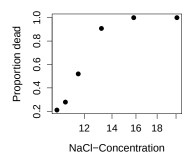


Figure 5.2: Proportion of fish died at different NaCl concentrations. The x axis is on a log scale.

¹ Ritz, 2010

Population and Metapopulation Effects

7

Community Effects

7.1 Species Richness

Example data

Ten species were sampled at eight sites around an outfall and the number of individuals per species noted. The data is available from the qetx package has a usual species x sites format and the rownames give the distance to the outfall¹:

```
data(abu)
abu
##
       Sp1 Sp2 Sp3 Sp4 Sp5 Sp6 Sp7 Sp8 Sp9 Sp10
                                  2
## -1
        12 11 10
                              2
                                      1
                                          0
## -0.5 12 12 10
                                          1
                   2
## 0
        58 21
               3
                      1
                                          0
## 0.5
       18 16 15
                  5 4
                           0
                              0
                                 0
                                     0
                                          0
                   8 7
## 1
        11 11 11
                                     1
                                          0
## 1.7
        8 12 13
                   3
## 2.7
       12 10
              8 11
                       8
                           4
                              1
                                  1
                                      2
                                          0
## 5.3 10 11
                8
```

¹ Most functions for multivariate techniques need the data in this format - rows = samples, columns = species.

For our analyses we will use two packages:

vegan A package for Community Ecology.

BiodiversityR A package forbiodiversity, suitability and community ecology analysis²

```
require(vegan)
require(BiodiversityR)
```

² Advisable further reading: Kindt and Coe (2005) freely availabe at http://www.worldagroforestry. org/resources/databases/ tree-diversity-analysis

Rarefaction

The total number of individuals per sample can be calculated as the row sums

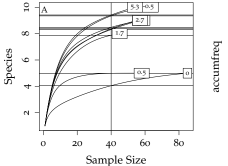
```
rowSums (abu)
##
      -1 -0.5
                      0.5
                                 1.7
                                        2.7
                                             5.3
                              1
##
      59
           62
                 85
                       58
                             55
                                   45
                                         57
                                               54
```

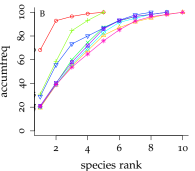
As the total number of individuals varies we will use the rarefaction method to estimate species richness at a sample size of 40 individuals.

```
rarefy(abu, 40, se = TRUE)
##
          -1 -0.5
                        0
                              0.5
                                        1
                                             1.7
                                                    2.7
## S 8.4757 9.347 4.0496 4.99092 8.3665 7.8767 8.3122
## se 0.6225 0.687 0.7458 0.09492 0.6788 0.3313 0.6962
##
         5.3
## S 9.4139
## se 0.6572
## attr(,"Subsample")
## [1] 40
```

As can be seen species richness drops at the outfall, but increases again downstream (Fig. 7.1). Rarefaction curves can be easily created as well as species accumulation curves:

```
rarecurve(abu, sample = 40)
env <- data.frame(dist = rownames(abu))
rankabuncomp(abu, env, 'dist', scale = 'accumfreq')</pre>
```





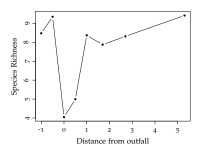


Figure 7.1: Species Richness along outfall

Figure 7.2: A) Rarefaction Curve. B) Rank abundance curve

7.2 Species Diversity

Function diversity() from the vegan package can calculate the Shannon and Simpson diversity indices ³:

```
diversity(abu, base = 2)
                                         0.5
          - 1
                   -0.5
                                0
## 2.8673512 2.9862404 1.2475503 2.1119181 2.7892213
         1.7
                    2.7
## 2.5720769 2.8102132 3.0159575
diversity(abu, index = "simpson")
##
                                   0
                                             0.5
           - 1
                     -0.5
                                                           1
## 0.84860672 0.85691988 0.47141869 0.74851367 0.83768595
                      2.7
## 0.79802469 0.84148969 0.86145405
```

Brillouin's index is not implemented so have to calculate it by hand:

```
N <- rowSums(abu)
log(factorial(N)/apply(factorial(abu), 1, prod), base = 2)/N
          - 1
                   -0.5
                                0
                                        0.5
## 2.5566733 2.6578637 1.1447406 1.9283402 2.4710966
         1.7
                    2.7
                              5.3
## 2.2413558 2.4986847 2.6527313
```

Or use the function brillouin from the qetx package, which runs the two above lines of code for us:

```
brillouin(abu, 2)
```

Species Eveness

Pielou's J can be easyily calculated from the shannon index by dividing it with the number of species in each sample:

```
diversity(abu)/log(rowSums(abu > 0))
                    -0.5
                                            0.5
## 0.90454859 0.89894792 0.53729068 0.90955361 0.87990134
##
          1.7
                     2.7
                                 5.3
## 0.85735897 0.88652356 0.90789368
```

³ The logarithm with base 2 is used for the Shannon index

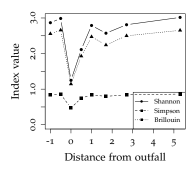


Figure 7.3: Diversity indices along outfall

7.3 Dissimilarity indices

```
data(oil)
oil
      Abra nitida Abra tenuis Acanthocardia echinata
##
## 1
                 0
                               1
                               0
                                                        2
## 2
                 0
## 3
                 0
                               0
                                                        0
                        0
                                               0
## 19
## 20
```

7.4 Ordination techniques - PCA, RDA, NMDS, MDS, dbRDA

⁴ Further reading: Borcard et al. (2011)

7.5 Testing hypotheses - PERMANOVA

7.6 Analyzing mesocosm data

Example data

Here we will analyze the pyrifos data set from Van den Brink and Ter Braak (1999) which is shipped with the vegan package.

Describe experiment

```
require(vegan)
data(pyrifos)
head(pyrifos[, c(1:8)])
          Simve Daplo Cerpu Alogu Aloco Alore Aloaf Copsp
## w.4.c1 3.951
                            0
                                  0
                                               0
                     0
                                                     0 2.773
## w.4.c2 2.303
                     0
                           0
                                  0
                                               0
                                                     0 2.079
## w.4.c3 4.595
                     0
                           0
                                  0
                                        0
                                               0
                                                     0 3.761
## w.4.c4 2.398
                     0
                           0
                                  0
                                        0
                                               0
                                                     0 3.296
## w.4.c5 4.025
                     0
                           0
                                  0
                                        0
                                               0
                                                     0 3.466
## w.4.c6 2.303
                           0
                                  0
                                                     0 2.197
                     0
```

So rows correspond to samples and columns are the species (with abbreviated names), a usual species x sites matrix. The column names code treatment and time, but we will create a separate data.frame with information about experimental ditch, sampling time and treatment:

```
ditch <- gl(12, 1, length = 132)
week \leftarrow gl(11, 12, labels = c(-4, -1, 0.1, 1, 2, 4, 8, 12,
    15, 19, 24))
dose <- factor(rep(c(0.1, 0, 0, 0.9, 0, 44, 6, 0.1, 44,
    0.9, 0, 6), 11))
pyrifos_env <- data.frame(ditch, week, dose)</pre>
```

Introduction

Principle Response Curves (PRC)⁵ are commonly used for analyzing ecotoxicological mesocosm experiments. PRC analyses the change of a community due to a treatment over time and is a special form of Redundancy Analysis (RDA) ⁶.

⁵ Van den Brink and Ter Braak, 1999

Overall pattern

With this a hand we can easily calculate and plot (Figure 7.4) 7 the PRC using the prc() function:

```
pyrifos_prc <- prc(response = pyrifos, treatment = dose,</pre>
    time = week)
pyrifos_prc_sum <- summary(pyrifos_prc, scaling = 1)</pre>
```

```
plot(pyrifos_prc, select = abs(pyrifos_prc_sum$sp) > 1,
    scaling = 1)
```

The plot shows on the x axis the time and on the y-axis the difference from the control treatments. The farther apart from the x-axis the more different are the communities compared to the control (you can say the x axis represents the control).

We see a clearly treatment-related effect: After application at week o the treated communities rapidly change treatment dependent. However to the end of the experiment the treated and the control get similar again, which indicates a 'recovery'.

On the right-hand side we see the species names and their scores. The more extreme the scores the more this species contributed to the plotted pattern. However, you cannot directly infer from these species scores which species are more susceptible. For example Gammarus pulex (gammapule) has a relatively low scores, although it's response pattern (Figure 7.5) shows a strong response, but with no recovery. PRC displays global pattern in the community, but the pattern of G. pulex is different from most other species, therefore it gets a lower species score.

⁷ Note that only species with scores greater or smaller than 1 are displayed to avoid cluttering of the plot

⁶ Legendre and Legendre, 2013

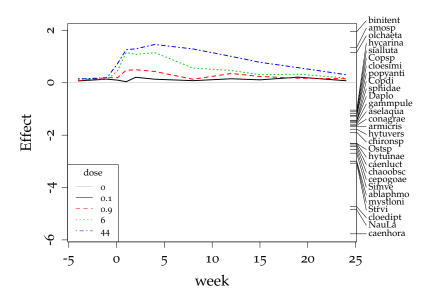


Figure 7.4: Principal response curves (PRC) with species weights for the pyrifos data set indicating effects of the insecticide on the invertebrate community.

We can also look at the numerical output⁸ for this plot using the summary method:

```
pyrifos_prc_sum
```

```
##
## prc(response = pyrifos, treatment = dose, time = week)
## Species scores:
      Simve
##
               0stsp
                         NauLa
                                  Strvi binitent caenhora
##
     -2.688
              -2.312
                        -4.847
                                 -3.070
                                           1.951
                                                    -5.768
##
  caenluct cloedipt hytuinae ablaphmo cepogoae chaoobsc
     -2.376
              -4.734
                        -2.316
                                 -2.993
                                          -2.555
                                                    -2.442
##
## mystloni
     -2.998
##
##
## Coefficients for dose + week:dose interaction
  which are contrasts to dose 0
   rows are dose, columns are week
##
                   - 1
                         0.1
                                    1
## 0.1 0.07218 0.1375 0.1020 0.04068 0.2101 0.1364
## 0.9 0.08106 0.1935 0.1936 0.47699 0.4977 0.4306
## 6 0.16616 0.1232 0.4539 1.15638 1.0835 1.1511
```

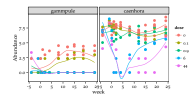


Figure 7.5: Responses of *G. pulex* and *C. hours a* 4956 responses of the pulex and the figure 7.5: Responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *C. hours a* 4956 responses of *G. pulex* and *G. hours a* 4956 response of *G. pulex* and *G. hours a* 4956 responses of *G. pulex* and *G.*

```
## 44 0.13979 0.1958 0.7308 1.26088 1.2978 1.4627
```

The output of prc() gives us more detailed information about the RDA model:

```
pyrifos_prc
```

```
## Call: prc(response = pyrifos, treatment = dose,
## time = week)
##
##
                   Inertia Proportion Rank
                 288.99201
                               1.00000
## Total
                  63.34934
## Conditional
                               0.21921
                                         10
## Constrained
                  96.68373
                               0.33456
                                         44
## Unconstrained 128.95894
                               0.44624
                                         77
## Inertia is variance
##
## Eigenvalues for constrained axes:
##
      RDA1
              RDA2
                       RDA3
                               RDA4
                                       RDA5
                                                RDA6
## 25.2823
           8.2969
                    6.0442 4.7662 4.1482 3.8568
##
      RDA7
              RDA8
    3.5875 3.3341
##
##
## Eigenvalues for unconstrained axes:
       PC1
               PC2
                        PC3
                                PC4
                                        PC5
                                                 PC<sub>6</sub>
##
## 17.1561
           9.1894
                    7.5849 6.0636 5.7298 4.8434
##
       PC7
               PC8
   4.5184 4.1046
## (Showed only 8 of all 77 unconstrained eigenvalues)
```

We see that 21.9 % of the variance can be attributed to time (Conditional), 33.5 % can be explained by the treatment regime (Constrained) and 44.6 % of residual variance (Unconstrained), which cannot be explained by time and treatment.

The first RDA axis has an eigenvalue of 25.3. If we divide this eigenvalue by the sum of all eigenvalues, we get the proportion of explained variance which is displayed on the first axis9:

```
pyrifos_prc$CCA$eig[1]/sum(pyrifos_prc$CCA$eig) * 100
##
       RDA1
## 26.14946
```

The significance of the PRC diagram can be tested via permutations. However observations from a experimental ditch are not independent, since the same ditch was measured repeatedly during the

⁹ rda() (and therefore also prc()) returns a huge object with all kind of information stored in it. See ?cca.object for the internal structure. Here I directly access the eigenvalues from this object

experiment. We have to take this into account: each ditch represents a time-series. We will permute the whole series of one ditch, keeping the temporal order (see Tab. 7.1).

To setup such a permutation scheme we use the permute package, which is automatically loaded with vegan:

```
control = how(plots = Plots(strata = ditch, type = "free"),
    within = Within(type = "none"), nperm = 99)
```

With this setup we can create a permutation matrix. Each row therein is one permutation, the values are the row numbers of the original data set.

```
set.seed(1234)
permutations <- shuffleSet(nrow(pyrifos), control = control)</pre>
```

This can be passed to permutest, testing the first eigenvalue of our model. 10

We see that our first axis explains a statistically significant proportion of variation (Fig. 7.6). The minimum p-value that we could get is 0.01 (=1/no. of permutations).

Effects per week

After looking at the overall treatment effect, we may want to look at effects at individual time-points. We follow here ¹¹ and use the ln-transformed nominal dose as continuous explanatory variable ¹².

Table 7.1: 3 ditches observed for 4 weeks and a possible permutation.

Week	Ditch	Perm
1	1	3
2	1	3
3	1	3 3 3 1
3 4 1	1	3
1	2	1
2	2	1
3	2	1
3 4	2	1
1	3	2
2	3	2
3	3	2
4	3	2

This sets up our permutation scheme: *plots* We will permute ditches, without any restrictions.

within But within one ditch there will be no permutations.

nperm We want 99 permutations.

¹⁰ vegan is in active development and at the moment the permute-package isn't fully hooked up. Therefore we have to create a permutation matrix beforehand. In the future we will be able to pass the permutation scheme directly into vegan functions.

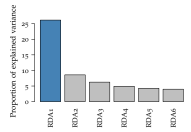


Figure 7.6: Proportion of explained variance of the first 6 RDA-axes. Note that treatment and time were factors and (internally) dummy-coded, therefore we have a total of 44 axes.

¹¹ Van den Brink and Ter Braak, 1999

¹² But before we have to convert dose from a factor to a numeric vector via as.numeric(levels(x))[x]

```
dose_c <- log(20 * as.numeric(levels(dose))[dose] + 1)</pre>
```

No we can write a for-loop and compute for every week a RDA and a permutation test ¹³.

```
rdas <- NULL
for (i in levels(week)) {
  rdas[[i]]$rda <- rda(pyrifos[week == i, ] ~
                          dose_c[week == i])
  rdas[[i]]$anova <- anova(rdas[[i]]$rda, by = 'terms',</pre>
                            step = 199)
}
```

However there is also convenience function in the qetx package:

```
rdas <- rda_per_time(pyrifos, dose_c, week)</pre>
```

This returns a very big list: one list entry per week and each entry itself contains two lists: rda(RDA-Model) and anova (permutation test)).

From this we have to extract the information we need. We can use sapply() to apply a function to every list entry and return results in a vector.

For example to extract the p-values for each week we can use:

```
sapply(rdas, function(x) x$anova[1, 5])
          -4
                    - 1
                             0.1
## 0.4371859296 0.8944723618 0.0050251256 0.00000000000
##
          2
                     4
                               8
15
                    19
## 0.0301507538 0.0100502513 0.1859296482
```

Have a look at the object structure to write a custom function to extract the information you need:

```
str(rdas[[1]]$anova)
```

Other methods

Other methods to analyse mesocosm experiments include:

```
multivariate GLMs <sup>14</sup> In R: mvabund-package.
```

trait-based indicators 15 Currently no package, but look at rspearpackage.

community endpoints ¹⁶ Use vegan for computations.

¹³ First we create an empty object (rdas) that will hold our results. Next we run on a subset of data (based on week) a RDA and permutation test. The results of both are stored as a list entry.

¹⁴ Warton et al., 2011; and Wang et al.,

¹⁵ Liess and Beketov, 2011

¹⁶ Sanchez-Bayo and Goka, 2012

7.7 Species Sensitivity Distributions

```
require(fitdistrplus)
# or
require(drc)
```

R Session Info

```
sessionInfo()
## R version 3.0.2 (2013-09-25)
## Platform: x86_64-pc-linux-gnu (64-bit)
##
## locale:
   [1] LC_CTYPE=en_US.UTF-8
  [2] LC_NUMERIC=C
##
## [3] LC_TIME=en_US.UTF-8
## [4] LC_COLLATE=en_US.UTF-8
## [5] LC_MONETARY=en_US.UTF-8
## [6] LC_MESSAGES=en_US.UTF-8
## [7] LC_PAPER=en_US.UTF-8
## [8] LC_NAME=C
## [9] LC_ADDRESS=C
## [10] LC_TELEPHONE=C
## [11] LC_MEASUREMENT=en_US.UTF-8
## [12] LC_IDENTIFICATION=C
##
## attached base packages:
  [1] splines
                stats4
                          tcltk
                                               stats
                                     grid
  [6] graphics grDevices utils
                                     datasets methods
## [11] base
##
## other attached packages:
## [1] tikzDevice_0.6.3
                           reshape2_1.2.2
## [3] qetx_0.0.1
                           ggplot2_0.9.3.1
## [5] fitdistrplus_1.0-1 survival_2.37-4
## [7] filehash_2.2-1
                           drc_{2.3-7}
## [9] plotrix_3.5-1
                           nlme_3.1-111
## [11] magic_1.5-4
                           gtools_3.1.0
## [13] BiodiversityR_2.3-6 vegan_2.1-35
## [15] lattice_0.20-24
                           permute_0.7-4
## [17] alr3_2.0.5
                           car_2.0-19
```

```
## [19] nnet_7.3-7
                           \texttt{MASS}\_\texttt{7.3-29}
## [21] abind_1.4-0
                           knitr_{-}1.5
##
## loaded via a namespace (and not attached):
## [1] codetools_0.2-8 colorspace_1.2-4
## [3] dichromat_2.0-0 digest_0.6.3
## [5] evaluate_0.5.1 formatR_0.9
## [7] gtable_0.1.2
                          highr_0.2.1
                          munsell_0.4.2
## [9] labeling_0.2
## [11] parallel_3.0.2
                          plyr_1.8
## [13] proto_0.3-10
                          Rcmdr_2.0-0
## [15] RColorBrewer_1.0-5 scales_0.2.3
## [17] stringr_0.6.2 tools_3.0.2
```

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