Multivariate Analysis of Ecological Communities in R: vegan tutorial

Jari Oksanen

June 10, 2015

Abstract

This tutorial demonstrates the use of ordination methods in R package vegan. The tutorial assumes familiarity both with R and with community ordination. Package vegan supports all basic ordination methods, including non-metric multidimensional scaling. The constrained ordination methods include constrained analysis of proximities, redundancy analysis and constrained correspondence analysis. Package vegan also has support functions for fitting environmental variables and for ordination graphics.

Contents

1 Introduction 2

2 Ordination: basic method 3
   2.1 Non-metric Multidimensional scaling .................. 3
   2.2 Community dissimilarities ............................ 5
   2.3 Comparing ordinations: Procrustes rotation .......... 8
   2.4 Eigenvector methods .................................. 8
   2.5 Detrended correspondence analysis ................... 11
   2.6 Ordination graphics ................................... 12

3 Environmental interpretation 14
   3.1 Vector fitting ........................................... 14
   3.2 Surface fitting ......................................... 16
   3.3 Factors .................................................. 16

4 Constrained ordination 18
   4.1 Model specification ..................................... 19
   4.2 Permutation tests ....................................... 21
   4.3 Model building .......................................... 23
   4.4 Linear combinations and weighted averages .......... 28
   4.5 Biplot arrows and environmental calibration .......... 29
   4.6 Conditioned or partial models ......................... 30
1 Introduction

This tutorial demonstrates typical work flows in multivariate ordination analysis of biological communities. The tutorial first discusses basic unconstrained analysis and environmental interpretation of their results. Then it introduces constrained ordination using constrained correspondence analysis as an example: alternative methods such as constrained analysis of proximities and redundancy analysis can be used (almost) similarly. Finally the tutorial describes analysis of species–environment relations without ordination, and briefly touches classification of communities.

The examples in this tutorial are tested: This is a Sweave document. The original source file contains only text and R commands: their output and graphics are generated while running the source through Sweave. However, you may need a recent version of vegan. This document was generated using vegan version 2.3-0 and R Under development (unstable) (2015-06-09 r68498).

The manual covers ordination methods in vegan. It does not discuss many other methods in vegan. For instance, there are several functions for analysis of biodiversity: diversity indices (diversity, renyi, fisher.alpha), extrapolated species richness (specpool, estimateR), species accumulation curves (specaccumm), species abundance models (radfit, fisherfit, prestonfit) etc. Neither is vegan the only R package for ecological community ordination. Base R has standard statistical tools, labdsv complements vegan with some advanced methods and provides alternative versions of some methods, and ade4 provides an alternative implementation for the whole gamme of ordination methods.

The tutorial explains only the most important methods and shows typical work flows. I see ordination primarily as a graphical tool, and I do not show too much exact numerical results. Instead, there are small vignettes of plotting results in the margins close to the place where you see a plot command. I suggest that you repeat the analysis, try different alternatives and inspect the results more thoroughly at your leisure. The functions are explained only briefly, and it is very useful to check the corresponding help pages for a more thorough explanation of methods. The methods also are only briefly explained. It is best to consult a textbook on ordination methods, or my lectures, for firmer theoretical background.

5 Dissimilarities and environment

5.1 adonis: Multivariate ANOVA based on dissimilarities . . . 32
5.2 Homogeneity of groups and beta diversity . . . . . . . . . 34
5.3 Mantel test . . . . . . . . . . . . . . . . . . . . . . . . . 35
5.4 Procrust: Procrustes test . . . . . . . . . . . . . . . . . . 36

6 Classification

6.1 Cluster analysis . . . . . . . . . . . . . . . . . . . . . . . 37
6.2 Display and interpretation of classes . . . . . . . . . . . . 38
6.3 Classified community tables . . . . . . . . . . . . . . . . . 40
2 Ordination: basic method

2.1 Non-metric Multidimensional scaling

Non-metric multidimensional scaling can be performed using isoMDS function in the MASS package. This function needs dissimilarities as input. Function vegdist in vegan contains dissimilarities which are found good in community ecology. The default is Bray-Curtis dissimilarity, nowadays often known as Steinhaus dissimilarity, or in Finland as Sørensen index. The basic steps are:

```r
> library(vegan)
> library(MASS)
> data(varespec)
> vare.dis <- vegdist(varespec)
> vare.mds0 <- isoMDS(vare.dis)
```

initial value 18.026495
iter 5 value 10.095483
final value 10.020469 converged

The default is to find two dimensions and use metric scaling (cmdscale) as the starting solution. The solution is iterative, as can be seen from the tracing information (this can be suppressed setting `trace = F`).

The results of isoMDS is a list (items points, stress) for the configuration and the stress. Stress $S$ is a statistic of goodness of fit, and it is a function of and non-linear monotone transformation of observed dissimilarities $\theta(d)$ and ordination distances $\tilde{d}$.

NMDS maps observed community dissimilarities nonlinearly onto ordination space and it can handle nonlinear species responses of any shape. We can inspect the mapping using function Shepard in MASS package, or a simple wrapper stressplot in vegan:

```r
> stressplot(vare.mds0, vare.dis)
```

Function stressplot draws a Shepard plot where ordination distances are plotted against community dissimilarities, and the fit is shown as a monotone step line. In addition, stressplot shows two correlation like statistics of goodness of fit. The correlation based on stress is $R^2 = 1 - S^2$. The “fit-based $R^2$” is the correlation between the fitted values $\theta(d)$ and ordination distances $\tilde{d}$, or between the step line and the points. This should be linear even when the fit is strongly curved and is often known as the “linear fit”. These two correlations are both based on the residuals in the Shepard plot, but they differ in their null models. In linear fit, the null model is that all ordination distances are equal, and the fit is a flat horizontal line. This sounds sensible, but you need $N - 1$ dimensions for the null model of $N$ points, and this null model is geometrically impossible in the ordination space. The basic stress uses the null model where all observations are put in the same point, which is geometrically possible. Finally a word of warning: you sometimes see that people use correlation between community dissimilarities and ordination distances. This is dangerous and misleading since NMDS is a nonlinear method: an improved
2.1 Non-metric Multidimensional scaling

ordination with more nonlinear relationship would appear worse with this criterion.

Functions scores and ordiplot in vegan can be used to handle the results of NMDS:

```r
> ordiplot(vare.mds0, type = "t")
```

Only site scores were shown, because dissimilarities did not have information about species.

The iterative search is very difficult in nmds, because of nonlinear relationship between ordination and original dissimilarities. The iteration easily gets trapped into local optimum instead of finding the global optimum. Therefore it is recommended to use several random starts, and select among similar solutions with smallest stresses. This may be tedious, but vegan has function metaMDS which does this, and many more things. The tracing output is long, and we suppress it with trace = 0, but normally we want to see that something happens, since the analysis can take a long time:

```r
> vare.mds <- metaMDS(varespec, trace = FALSE)
> vare.mds
Call: metaMDS(comm = varespec, trace = FALSE)

global Multidimensional Scaling using monoMDS

Data: wisconsin(sqrt(varespec))
Distance: bray

Dimensions: 2
Stress: 0.1826
Stress type 1, weak ties
No convergent solutions - best solution after 20 tries
Scaling: centring, PC rotation, halfchange scaling
Species: expanded scores based on ‘wisconsin(sqrt(varespec))’
```
2. Function used Bray–Curtis dissimilarities.

3. Function run isoMDS with several random starts, and stopped either after a certain number of tries, or after finding two similar configurations with minimum stress. In any case, it returned the best solution.

4. Function rotated the solution so that the largest variance of site scores will be on the first axis.

5. Function scaled the solution so that one unit corresponds to halving of community similarity from the replicate similarity.

6. Function found species scores as weighted averages of site scores, but expanded them so that species and site scores have equal variances. This expansion can be undone using `shrink = TRUE` in display commands.

The help page for `metaMDS` will give more details, and point to explanation of functions used in the function.

### 2.2 Community dissimilarities

Non-metric multidimensional scaling is a good ordination method because it can use ecologically meaningful ways of measuring community dissimilarities. A good dissimilarity measure has a good rank order relation to distance along environmental gradients. Because nmds only uses rank information and maps ranks non-linearly onto ordination space, it can handle non-linear species responses of any shape and effectively and robustly find the underlying gradients.

The most natural dissimilarity measure is Euclidean distance which is inherently used by eigenvector methods of ordination. It is the distance in species space. Species space means that each species is an axis orthogonal to all other species, and sites are points in this multidimensional hyperspace. However, Euclidean distance is based on squared differences and strongly dominated by single large differences. Most ecologically meaningful dissimilarities are of Manhattan type, and use differences instead of squared differences. Another feature in good dissimilarity indices is that they are proportional: if two communities share no species, they have a maximum dissimilarity = 1. Euclidean and Manhattan dissimilarities will vary according to total abundances even though there are no shared species.

Package `vegan` has function `vegdist` with Bray–Curtis, Jaccard and Kulczyński indices. All these are of the Manhattan type and use only first order terms (sums and differences), and all are relativized by site total and reach their maximum value (1) when there are no shared species between two compared communities. Function `vegdist` is a drop-in replacement for standard R function `dist`, and either of these functions can be used in analyses of dissimilarities.

There are many confusing aspects in dissimilarity indices. One is that same indices can be written with very different looking equations: two alternative formulations of Manhattan dissimilarities in the margin serve

\[
d_{jk} = \sum_{i=1}^{N} (x_{ij} - x_{ik})^2 \quad \text{Euclidean}
\]

\[
d_{jk} = \sum_{i=1}^{N} |x_{ij} - x_{ik}| \quad \text{Manhattan}
\]

\[
A = \sum_{i=1}^{N} x_{ij} \quad B = \sum_{i=1}^{N} x_{ik}
\]

\[
J = \sum_{i=1}^{N} \min(x_{ij}, x_{ik})
\]

\[
d_{jk} = A + B - 2J \quad \text{Manhattan}
\]

\[
d_{jk} = \frac{A + B - 2J}{A + B} \quad \text{Bray}
\]

\[
d_{jk} = \frac{A + B - 2J}{A + B - J} \quad \text{Jaccard}
\]

\[
d_{jk} = 1 - \frac{1}{2} \left( \frac{J}{A} + \frac{J}{B} \right) \quad \text{Kulczyński}
\]
as an example. Another complication is naming. Function \texttt{vegdist} uses colloquial names which may not be strictly correct. The default index in \texttt{vegan} is called Bray (or Bray–Curtis), but it probably should be called Steinhaus index. On the other hand, its correct name was supposed to be Czekanowski index some years ago (but now this is regarded as another index), and it is also known as Sørensen index (but usually misspelt). Strictly speaking, Jaccard index is binary, and the quantitative variant in \texttt{vegan} should be called Ružička index. However, \texttt{vegan} finds either quantitative or binary variant of any index under the same name.

These three basic indices are regarded as good in detecting gradients. In addition, \texttt{vegdist} function has indices that should satisfy other criteria. Morisita, Horn–Morisita, Raup–Cric, Binomial and Mountford indices should be able to compare sampling units of different sizes. Euclidean, Canberra and Gower indices should have better theoretical properties.

Function \texttt{metaMDS} used Bray-Curtis dissimilarity as default, which usually is a good choice. Jaccard (Ružička) index has identical rank order, but has better metric properties, and probably should be preferred. Function \texttt{rankindex} in \texttt{vegan} can be used to study which of the indices best separates communities along known gradients using rank correlation as default. The following example uses all environmental variables in data set \texttt{varechem}, but standardizes these to unit variance:

```r
> data(varechem)
> rankindex(scale(varechem), varespec, c("euc","man","bray","jac","kul"))

               euc         man         bray        jac        kul
[1,] 0.2395929 0.2735314 0.2837898 0.2837898 0.2840057
```

are non-linearly related, but they have identical rank orders, and their rank correlations are identical. In general, the three recommended indices are fairly equal.

I took a very practical approach on indices emphasizing their ability to recover underlying environmental gradients. Many textbooks emphasize metric properties of indices. These are important in some methods, but not in \texttt{nMDS} which only uses rank order information. The metric properties simply say that

\[
\begin{align*}
\text{for } A = B & \quad d_{AB} = 0 \\
\text{for } A \neq B & \quad d_{AB} > 0 \\
d_{AB} &= d_{BA} \\
d_{AB} &\leq d_{Ax} + d_{xB}
\end{align*}
\]

1. if two sites are identical, their distance is zero,
2. if two sites are different, their distance is larger than zero,
3. distances are symmetric, and
4. the shortest distance between two sites is a line, and you cannot improve by going through other sites.

These all sound very natural conditions, but they are not fulfilled by all dissimilarities. Actually, only Euclidean distances – and probably Jaccard index – fulfill all conditions among the dissimilarities discussed here, and are metrics. Many other dissimilarities fulfill three first conditions and are semimetrics.

There is a school that says that we should use metric indices, and most naturally, Euclidean distances. One of their drawbacks was that
they have no fixed limit, but two sites with no shared species can vary in dissimilarities, and even look more similar than two sites sharing some species. This can be cured by standardizing data. Since Euclidean distances are based on squared differences, a natural transformation is to standardize sites to equal sum of squares, or to their vector norm using function `decostand`:

```r
> dis <- vegdist(decostand(varespec, "norm"), "euclid")
```

This gives chord distances which reach a maximum limit of $\sqrt{2}$ when there are no shared species between two sites. Another recommended alternative is Hellinger distance which is based on square roots of sites standardized to unit total:

```r
> dis <- vegdist(decostand(varespec, "hell"), "euclidean")
```

Despite standardization, these still are Euclidean distances with all their good properties, but for transformed data. Actually, it is often useful to transform or standardize data even with other indices. If there is a large difference between smallest non-zero abundance and largest abundance, we want to reduce this difference. Usually square root transformation is sufficient to balance the data. Wisconsin double standardization often improves the gradient detection ability of dissimilarity indices; this can be performed using command `wisconsin` in `vegan`. Here we first divide all species by their maxima, and then standardize sites to unit totals. After this standardization, many dissimilarity indices become identical in rank ordering and should give equal results in `nmds`.

You are not restricted to use only `vegdist` indices in `vegan`: `vegdist` returns similar dissimilarity structure as standard R function `dist` which also can be used, as well as any other compatible function in any package. Some compatible functions are `dsvdis` (labdsv package), `daisy` (cluster package), and `distance` (analogue package), and beta diversity indices in `betadiver` in `vegan`. Moreover, `vegan` has function `designdist` where you can define your own dissimilarity indices by writing its equation using either the notation for $A$, $B$ and $J$ above, or with binary data, the $2 \times 2$ contingency table notation where $a$ is the number of species found on both compared sites, and $b$ and $c$ are numbers of species found only in one of the sites. The following three equations define the same Sørensen index where the number of shared species is divided by the average species richness of compared sites:

```r
> d <- vegdist(varespec, "bray", binary = TRUE)
> d <- designdist(varespec, "((A+B-2*J)/(A+B))")
> d <- designdist(varespec, "((b+c)/(2*a+b+c))", abcd=TRUE)
```

Function `betadiver` defines some more binary dissimilarity indices in `vegan`.

Most published dissimilarity indices can be expressed as `designdist` formulæ. However, it is much easier and safer to use the canned alternatives in existing functions: it is very easy to make errors in writing the dissimilarity equations.
2.3 Comparing ordinations: Procrustes rotation

Two ordinations can be very similar, but this may be difficult to see, because axes have slightly different orientation and scaling. Actually, in nmds the sign, orientation, scale and location of the axes are not defined, although metaMDS uses simple method to fix the last three components. The best way to compare ordinations is to use Procrustes rotation. Procrustes rotation uses uniform scaling (expansion or contraction) and rotation to minimize the squared differences between two ordinations. Package vegan has function procrustes to perform Procrustes analysis.

How much did we gain with using metaMDS instead of default isoMDS?

```r
> tmp <- wisconsin(sqrt(varespec))
> dis <- vegdist(tmp)
> vare.mds0 <- isoMDS(dis, trace = 0)
> pro <- procrustes(vare.mds, vare.mds0)
> pro
Call: procrustes(X = vare.mds, Y = vare.mds0)
Procrustes sum of squares:
0.156
> plot(pro)
```

In this case the differences were fairly small, and mainly concerned two points. You can use identify function to identify those points in an interactive session, or you can ask a plot of residual differences only:

```r
> plot(pro, kind = 2)
```

The descriptive statistic is “Procrustes sum of squares” or the sum of squared arrows in the Procrustes plot. Procrustes rotation is nonsymmetric, and the statistic would change with reversing the order of ordinations in the call. With argument symmetric = TRUE, both solutions are first scaled to unit variance, and a more scale-independent and symmetric statistic is found (often known as Procrustes $m^2$).

2.4 Eigenvector methods

Non-metric multidimensional scaling was a hard task, because any kind of dissimilarity measure could be used and dissimilarities were nonlinearly mapped into ordination. If we accept only certain types of dissimilarities and make a linear mapping, the ordination becomes a simple task of rotation and projection. In that case we can use eigenvector methods. Principal components analysis (PCA) and correspondence analysis (CA) are the most important eigenvector methods in community ordination. In addition, principal coordinates analysis a.k.a. metric scaling (MDS) is used occasionally. PCA is based on Euclidean distances, CA is based on Chi-square distances, and principal coordinates can use any dissimilarities (but with Euclidean distances it is equal to PCA).

PCA is a standard statistical method, and can be performed with base R functions prcomp or princomp. Correspondence analysis is not as ubiquitous, but there are several alternative implementations for that also. In
this tutorial I show how to run these analyses with vegan functions rda and cca which actually were designed for constrained analysis.

Principal components analysis can be run as:

```r
> vare.pca <- rda(varespec)
> vare.pca
```

```
Call: rda(X = varespec)
Inertia Rank
Total 1826
Unconstrained 1826 23
Inertia is variance
Eigenvalues for unconstrained axes:
PC1 PC2 PC3 PC4 PC5 PC6 PC7 PC8
983 464 132 74 48 37 26 20
(Showed only 8 of all 23 unconstrained eigenvalues)

> plot(vare.pca)
```

The output tells that the total inertia is 1826, and the inertia is variance. The sum of all 23 (rank) eigenvalues would be equal to the total inertia. In other words, the solution decomposes the total variance into linear components. We can easily see that the variance equals inertia:

```r
> sum(apply(varespec, 2, var))
```

```
[1] 1826
```

Function apply applies function var or variance to dimension 2 or columns (species), and then sum takes the sum of these values. Inertia is the sum of all species variances. The eigenvalues sum up to total inertia. In other words, they each “explain” a certain proportion of total variance. The first axis “explains” 983/1826 = 53.8 % of total variance.

The standard ordination plot command uses points or labels for species and sites. Some people prefer to use biplot arrows for species in PCA and possibly also for sites. There is a special biplot function for this purpose:

```r
> biplot(vare.pca, scaling = -1)
```

For this graph we specified scaling = -1. The results are scaled only when they are accessed, and we can flexibly change the scaling in plot, biplot and other commands. The negative values mean that species scores are divided by the species standard deviations so that abundant and scarce species will be approximately as far away from the origin.

The species ordination looks somewhat unsatisfactory: only reindeer lichens (Cladina) and Pleurozium schreberi are visible, and all other species are crowded at the origin. This happens because inertia was variance, and only abundant species with high variances are worth explaining (but we could hide this in plot by setting negative scaling). Standardizing all species to unit variance, or using correlation coefficients instead of covariances will give a more balanced ordination:

```r
> vare.pca <- rda(varespec, scale = TRUE)
> vare.pca
```
Call: rda(X = varespec, scale = TRUE)

Inertia Rank
Total 44
Unconstrained 44 23
Inertia is correlations

Eigenvalues for unconstrained axes:
PC1 PC2 PC3 PC4 PC5 PC6 PC7 PC8
8.90 4.76 4.26 3.73 2.96 2.88 2.73 2.18
(Showed only 8 of all 23 unconstrained eigenvalues)

Now inertia is correlation, and the correlation of a variable with itself is one. Thus the total inertia is equal to the number of variables (species). The rank or the total number of eigenvectors is the same as previously. The maximum possible rank is defined by the dimensions of the data: it is one less than smaller of number of species or number of sites:

> dim(varespec)
[1] 24 44

If there are species or sites similar to each other, rank will be reduced even from this.

The percentage explained by the first axis decreased from the previous pca. This is natural, since previously we needed to “explain” only the abundant species with high variances, but now we have to explain all species equally. We should not look blindly at percentages, but the result we get.

Correspondence analysis is very similar to pca:

> vare.ca <- cca(varespec)
> vare.ca

Call: cca(X = varespec)

Inertia Rank
Total 2.08
Unconstrained 2.08 23
Inertia is mean squared contingency coefficient

Eigenvalues for unconstrained axes:
CA1 CA2 CA3 CA4 CA5 CA6 CA7 CA8
0.525 0.357 0.234 0.195 0.178 0.122 0.115 0.089
(Showed only 8 of all 23 unconstrained eigenvalues)

Now the inertia is called mean squared contingency coefficient. Correspondence analysis is based on Chi-squared distance, and the inertia is the Chi-squared statistic of a data matrix standardized to unit total:

> chisq.test(varespec/sum(varespec))

Pearson’s Chi-squared test
data: varespec/sum(varespec)
X-squared = 2.1, df = 990, p-value = 1
You should not pay any attention to \( P \)-values which are certainly misleading, but notice that the reported \( X^2 \)-squared is equal to the inertia above.

Correspondence analysis is a weighted averaging method. In the graph above species scores were weighted averages of site scores. With different scaling of results, we could display the site scores as weighted averages of species scores:

\[
> \text{plot(vare.ca, scaling = 1)}
\]

We already saw an example of \( \text{scaling} = 3 \) or symmetric scaling in PCA. The other two integers mean that either species are weighted averages of sites (2) or sites are weighted averages of species (1). When we take weighted averages, the range of averages shrinks from the original values. The shrinkage factor is equal to the eigenvalue of \( \text{ca} \), which has a theoretical maximum of 1.

### 2.5 Detrended correspondence analysis

Correspondence analysis is a much better and more robust method for community ordination than principal components analysis. However, with long ecological gradients it suffers from some drawbacks or “faults” which were corrected in detrended correspondence analysis (DCA):

- Single long gradients appear as curves or arcs in ordination (arc effect): the solution is to detrend the later axes by making their means equal along segments of previous axes.

- Sites are packed more closely at gradient extremes than at the centre: the solution is to rescale the axes to equal variances of species scores.

- Rare species seem to have an unduly high influence on the results: the solution is to downweight rare species.

All these three separate tricks are incorporated in function \( \text{decorana} \) which is a faithful port of Mark Hill’s original programme with the same name. The usage is simple:

\[
> \text{vare.dca <- decorana(varespec)}
> \text{vare.dca}
\]

Call:

\[
\text{decorana(veg = varespec)}
\]

Detrended correspondence analysis with 26 segments.
Rescaling of axes with 4 iterations.

<table>
<thead>
<tr>
<th></th>
<th>DCA1</th>
<th>DCA2</th>
<th>DCA3</th>
<th>DCA4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eigenvalues</td>
<td>0.524</td>
<td>0.325</td>
<td>0.2001</td>
<td>0.1918</td>
</tr>
<tr>
<td>Decorana values</td>
<td>0.525</td>
<td>0.157</td>
<td>0.0967</td>
<td>0.0608</td>
</tr>
<tr>
<td>Axis lengths</td>
<td>2.816</td>
<td>2.205</td>
<td>1.5465</td>
<td>1.6486</td>
</tr>
</tbody>
</table>

\[
> \text{plot(vare.dca, display="sites")}
\]
2.6 Ordination graphics

Function `decorana` finds only four axes. Eigenvalues are defined as shrinkage values in weighted averages, similarly as in `cca` above. The “Decorana values” are the numbers that the original programme returns as “eigenvalues” — I have no idea of their possible meaning, and they should not be used. Most often people comment on axis lengths, which sometimes are called “gradient lengths”. The etymology is obscure: these are not gradients, but ordination axes. It is often said that if the axis length is shorter than two units, the data are linear, and `pca` should be used. This is only folklore and not based on research which shows that `ca` is at least as good as `pca` with short gradients, and usually better.

The current data set is homogeneous, and the effects of `dca` are not very large. In heterogeneous data with a clear arc effect the changes often are more dramatic. Rescaling may have larger influence than detrending in many cases.

The default analysis is without downweighting of rare species: see help pages for the needed arguments. Actually, `downweight` is an independent function that can be used with `cca` as well.

There is a school of thought that regards `dca` as the method of choice in unconstrained ordination. However, it seems to be a fragile and vague back of tricks that is better avoided.

2.6 Ordination graphics

We have already seen many ordination diagrams in this tutorial with one feature in common: they are cluttered and labels are difficult to read. Ordination diagrams are difficult to draw cleanly because we must put a large number of labels in a small plot, and often it is impossible to draw clean plots with all items labelled. In this chapter we look at producing cleaner plots. For this we must look at the anatomy of plotting functions in `vegan` and see how to gain a better control of default functions.

Ordination functions in `vegan` have their dedicated `plot` functions which provides a simple plot. For instance, the result of `decorana` is displayed by function `plot.decorana` which behind the scenes is called by our `plot` function. Alternatively, we can use function `ordiplot` which also works with many non-`vegan` ordination functions, but uses points instead of text as default. The `plot.decorana` function (or `ordiplot`) actually works in three stages:

1. It draws an empty plot with labelled axes, but with no symbols for sites or species.
2. It uses functions `text` or `points` to add species to the empty frame. If the user does not ask specifically, the function will use `text` in small data sets and `points` in large data sets.
3. It adds the sites similarly.

For better control of the plots we must repeat these stages by hand: draw an empty plot and then add sites and/or species as desired.

In this chapter we study a difficult case: plotting the Barro Colorado Island ordinations.
> data(BCI)

This is a difficult data set for plotting: it has 225 species and there is no way of labelling them all cleanly – unless we use very large plotting area with small text. We must show only a selection of the species or small parts of the plot. First an ordination with `decorana` and its default plot:

```r
> mod <- decorana(BCI)
> plot(mod)
```

There is an additional problem in plotting species ordination with these data:

```r
> names(BCI)[1:5]
[1] "Abarema.macradenia" "Acacia.melanoceras"
[3] "Acalypha.diversifolia" "Acalypha.macrostachya"
[5] "Adelia.triloba"
```

The data set uses full species names, and there is no way of fitting those in ordination graphs. There is a utility function `make.cepnames` in `vegan` to abbreviate Latin names:

```r
> shnam <- make.cepnames(names(BCI))
> shnam[1:5]
[1] "Abarmacr" "Acacmela" "Acaldive" "Acalmacr" "Adeltril"
```

The easiest way to selectively label species is to use interactive `identify` function: when you click next to a point, its label will appear on the side you clicked. You can finish labelling clicking the right mouse button, or with handicapped one-button mouse, you can hit the `esc` key.

```r
> pl <- plot(mod, dis="sp")
```

All `vegan` ordination plot functions return invisibly an `ordiplot` object which contains information on the points plotted. This invisible result can be caught and used as input to `identify`. The following selectively labels some extreme species as clicked:

```r
> identify(pl, "sp", labels=shnam)
```

There is an “ordination text or points” function `orditorp` in `vegan`. This function will label an item only if this can be done without overwriting previous labels. If an item cannot be labelled with text, it will be marked as a point. Items are processed either from the margin toward the centre, or in decreasing order of priority. The following gives higher priority to the most abundant species:

```r
> stems <- colSums(BCI)
> plot(mod, dis="sp", type="n")
> sel <- orditorp(mod, dis="sp", lab=shnam, priority=stems, pcol = "gray", pch="+")
```

We also can zoom into some parts of the ordination diagrams by setting `xlim` and `ylim`, and we can see more details.

An alternative to `orditorp` is function `ordilabel` which draws text on opaque labels that cover other labels below them. All labels cannot be displayed, but at least the uppermost are readable. Argument `priority` works similarly as in `orditorp` and can be used to select which of the labels are most important to show:
Finally, there is function ordipointlabel which uses both points and labels to these points. The points are in fixed positions, but the labels are iteratively located to minimize their overlap. The Barro Colorado Island data set has much too many names for the ordipointlabel function, but it can be useful in many cases.

In addition to these automatic functions, function orditkplot allows editing of plots. It has points in fixed positions with labels that can be dragged to better places with a mouse. The function uses different graphical toolset (Tcl/Tk) than ordinary R graphics, but the results can be passed to standard R plot functions for editing or directly saved as graphics files. Moreover, the ordipointlabel output can be edited using orditkplot.

Functions identify, orditorp, ordilabel and ordipointlabel may provide a quick and easy way to inspect ordination results. Often we need a better control of graphics, and judiciously select the labelled species. In that case we can first draw an empty plot (with type = "n"), and then use select argument in ordination text and points functions. The select argument can be a numeric vector that lists the indices of selected items. Such indices are displayed from identify functions which can be used to help in selecting the items. Alternatively, select can be a logical vector which is TRUE to selected items. Such a list was produced invisibly from orditorp. You cannot see invisible results directly from the method, but you can catch the result like we did above in the first orditorp call, and use this vector as a basis for fully controlled graphics. In this case the first items were:

```
> sel[1:14]
Abarmacr Acacmela Acaldive Acalmacr Adeltril Aegipana Alchcost   
FALSE FALSE FALSE FALSE FALSE FALSE FALSE
Alchlati Alibedul Allopsil Alseblac Amaicory Anacexce Andiiner   
TRUE FALSE FALSE FALSE TRUE TRUE FALSE
```

### 3 Environmental interpretation

It is often possible to “explain” ordination using ecological knowledge on studied sites, or knowledge on the ecological characteristics of species. Usually it is preferable to use external environmental variables to interpret the ordination. There are many ways of overlaying environmental information onto ordination diagrams. One of the simplest is to change the size of plotting characters according to an environmental variables (argument cex in plot functions). The vegan package has some useful functions for fitting environmental variables.

#### 3.1 Vector fitting

The most commonly used method of interpretation is to fit environmental vectors onto ordination. The fitted vectors are arrows with the interpretation:
• The arrow points to the direction of most rapid change in the the environmental variable. Often this is called the direction of the gradient.

• The length of the arrow is proportional to the correlation between ordination and environmental variable. Often this is called the strength of the gradient.

Fitting environmental vectors is easy using function **envfit**. The example uses the previous **NMDS** result and environmental variables in the data set **varechem**:

```r
> data(varechem)
> ef <- envfit(vare.mds, varechem, permu = 999)
> ef

***VECTORS

<table>
<thead>
<tr>
<th></th>
<th>NMDS1</th>
<th>NMDS2</th>
<th>r2</th>
<th>Pr(&gt;r)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>-0.057</td>
<td>-0.998</td>
<td>0.25</td>
<td>0.046 *</td>
</tr>
<tr>
<td>P</td>
<td>0.620</td>
<td>0.785</td>
<td>0.19</td>
<td>0.103</td>
</tr>
<tr>
<td>K</td>
<td>0.767</td>
<td>0.642</td>
<td>0.18</td>
<td>0.132</td>
</tr>
<tr>
<td>Ca</td>
<td>0.685</td>
<td>0.728</td>
<td>0.41</td>
<td>0.006 **</td>
</tr>
<tr>
<td>Mg</td>
<td>0.633</td>
<td>0.775</td>
<td>0.43</td>
<td>0.002 **</td>
</tr>
<tr>
<td>S</td>
<td>0.191</td>
<td>0.982</td>
<td>0.18</td>
<td>0.118</td>
</tr>
<tr>
<td>Al</td>
<td>-0.872</td>
<td>0.490</td>
<td>0.53</td>
<td>0.002 **</td>
</tr>
<tr>
<td>Fe</td>
<td>-0.936</td>
<td>0.352</td>
<td>0.44</td>
<td>0.004 **</td>
</tr>
<tr>
<td>Mn</td>
<td>0.799</td>
<td>-0.602</td>
<td>0.52</td>
<td>0.001 ***</td>
</tr>
<tr>
<td>Zn</td>
<td>0.617</td>
<td>0.787</td>
<td>0.19</td>
<td>0.098 .</td>
</tr>
<tr>
<td>Mo</td>
<td>-0.903</td>
<td>0.430</td>
<td>0.06</td>
<td>0.519</td>
</tr>
<tr>
<td>Baresoil</td>
<td>0.925</td>
<td>-0.380</td>
<td>0.25</td>
<td>0.049 *</td>
</tr>
<tr>
<td>Humdepth</td>
<td>0.933</td>
<td>-0.360</td>
<td>0.52</td>
<td>0.002 **</td>
</tr>
<tr>
<td>pH</td>
<td>-0.648</td>
<td>0.762</td>
<td>0.23</td>
<td>0.072 .</td>
</tr>
</tbody>
</table>

---

Signif. codes:  0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1
Permutation: free
Number of permutations: 999
```

The first two columns give direction cosines of the vectors, and r2 gives the squared correlation coefficient. For plotting, the axes should be scaled by the square root of r2. The **plot** function does this automatically, and you can extract the scaled values with **scores(ef, "vectors")**. The significances (Pr>r), or P-values are based on random permutations of the data; if you often get as good or better R2 with randomly permuted data, your values are insignificant.

You can add the fitted vectors to an ordination using **plot** command. You can limit plotting to most significant variables with argument **p.max**. As usual, more options can be found in the help pages.

```r
> plot(vare.mds, display = "sites")
> plot(ef, p.max = 0.1)
```
3.2 Surface fitting

Vector fitting is popular, and it provides a compact way of simultaneously displaying a large number of environmental variables. However, it implies a linear relationship between ordination and environment: direction and strength are all you need to know. This may not always be appropriate.

Function `ordisurf` fits surfaces of environmental variables to ordinations. It uses generalized additive models in function `gam` of package `mgcv`. Function `gam` uses thinplate splines in two dimensions, and automatically selects the degree of smoothing by generalized cross-validation. If the response really is linear and vectors are appropriate, the fitted surface is a plane whose gradient is parallel to the arrow, and the fitted contours are equally spaced parallel lines perpendicular to the arrow.

In the following example I introduce two new R features:

- Function `envfit` can be called with formula interface. Formula has a special character tilde (\(\sim\)), and the left-hand side gives the ordination results, and the right-hand side lists the environmental variables. In addition, we must define the name of the data containing the fitted variables.

- The variables in data frames are not visible to R session unless the data frame is `attach`ed to the session. We may not want to make all variables visible to the session, because there may be synonymous names, and we may use wrong variables with the same name in some analyses. We can use function `with` which makes the given data frame visible only to the following command.

Now we are ready for the example. We make vector fitting for selected variables and add fitted surfaces in the same plot.

```r
> ef <- envfit(vare.mds ~ Al + Ca, varechem)
> plot(vare.mds, display = "sites")
> plot(ef)
> tmp <- with(varechem, ordisurf(vare.mds, Al, add = TRUE))
> with(varechem, ordisurf(vare.mds, Ca, add = TRUE, col = "green4"))
```

Function `ordisurf` returns the result of fitted `gam`. If we save that result, like we did in the first fit with `Al`, we can use it for further analyses, such as statistical testing and prediction of new values. For instance, `fitted(ef)` will give the actual fitted values for sites.

3.3 Factors

Class centroids are a natural choice for factor variables, and \(R^2\) can be used as a goodness-of-fit statistic. The “significance” can be tested with permutations just like in vector fitting. Variables can be defined as factors in R, and they will be treated accordingly without any special tricks.

As an example, we shall inspect dune meadow data which has several class variables. Function `envfit` also works with factors:

```r
> data(dune)
> data(dune.env)
> dune.ca <- cca(dune)
```
> ef <- envfit(dune.ca, dune.env, permutations = 999)
> ef

***VECTORS

CA1  CA2  r2  Pr(r)
A1   0.9980 0.0606 0.31  0.04 *
---
Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1
Permutation: free
Number of permutations: 999

***FACTORS:

Centroids:

<table>
<thead>
<tr>
<th></th>
<th>CA1</th>
<th>CA2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture1</td>
<td>-0.75</td>
<td>-0.14</td>
</tr>
<tr>
<td>Moisture2</td>
<td>-0.47</td>
<td>-0.22</td>
</tr>
<tr>
<td>Moisture4</td>
<td>0.18</td>
<td>-0.73</td>
</tr>
<tr>
<td>Moisture5</td>
<td>1.11</td>
<td>0.57</td>
</tr>
<tr>
<td>ManagementBF</td>
<td>-0.73</td>
<td>-0.14</td>
</tr>
<tr>
<td>ManagementHF</td>
<td>-0.39</td>
<td>-0.30</td>
</tr>
<tr>
<td>ManagementNM</td>
<td>0.65</td>
<td>1.44</td>
</tr>
<tr>
<td>ManagementSF</td>
<td>0.34</td>
<td>-0.68</td>
</tr>
<tr>
<td>UseHayfield</td>
<td>-0.29</td>
<td>0.65</td>
</tr>
<tr>
<td>UseHaypastu</td>
<td>-0.07</td>
<td>-0.56</td>
</tr>
<tr>
<td>UsePasture</td>
<td>0.52</td>
<td>0.05</td>
</tr>
<tr>
<td>Manure0</td>
<td>0.65</td>
<td>1.44</td>
</tr>
<tr>
<td>Manure1</td>
<td>-0.46</td>
<td>-0.17</td>
</tr>
<tr>
<td>Manure2</td>
<td>-0.59</td>
<td>-0.36</td>
</tr>
<tr>
<td>Manure3</td>
<td>0.52</td>
<td>-0.32</td>
</tr>
<tr>
<td>Manure4</td>
<td>-0.21</td>
<td>-0.88</td>
</tr>
</tbody>
</table>

Goodness of fit:

<table>
<thead>
<tr>
<th></th>
<th>r2</th>
<th>Pr(r)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture</td>
<td>0.41</td>
<td>0.008</td>
</tr>
<tr>
<td>Management</td>
<td>0.44</td>
<td>0.001</td>
</tr>
<tr>
<td>Use</td>
<td>0.18</td>
<td>0.078</td>
</tr>
<tr>
<td>Manure</td>
<td>0.46</td>
<td>0.006</td>
</tr>
</tbody>
</table>
---
Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1
Permutation: free
Number of permutations: 999

> plot(dune.ca, display = "sites")
> plot(ef)

The names of factor centroids are formed by combining the name of the factor and the name of the level. Now the axes show the centroids for the level, and the $R^2$ values are for the whole factor, just like the significance test. The plot looks congested, and we may use tricks of §2.6 (p. 12) to make cleaner plots, but obviously not all factors are necessary in interpretation.

Package **vegan** has several functions for graphical display of factors. Function **ordihull** draws an enclosing convex hull for the items in a
class, `ordispider` combines items to their (weighted) class centroid, and `ordiellipse` draws ellipses for class standard deviations, standard errors or confidence areas. The example displays all these for `Management` type in the previous ordination and automatically labels the groups in `ordispider` command:

```r
> plot(dune.ca, display = "sites", type = "p")
> with(dune.env, ordiellipse(dune.ca, Management, kind = "se", conf = 0.95))
> with(dune.env, ordispider(dune.ca, Management, col = "blue", label = TRUE))
> with(dune.env, ordihull(dune.ca, Management, col = "blue", lty = 2))
```

Correspondence analysis is a weighted ordination method, and `vegan` functions `envfit` and `ordisurf` will do weighted fitting, unless the user specifies equal weights.

## 4 Constrained ordination

In unconstrained ordination we first find the major compositional variation, and then relate this variation to observed environmental variation. In constrained ordination we do not want to display all or even most of the compositional variation, but only the variation that can be explained by the used environmental variables, or constraints. Constrained ordination is often known as “canonical” ordination, but this name is misleading: there is nothing particularly canonical in these methods (see your favorite Dictionary for the term). The name was taken into use, because there is one special statistical method, canonical correlations, but these indeed are canonical: they are correlations between two matrices regarded to be symmetrically dependent on each other. The constrained ordination is non-symmetric: we have “independent” variables or constraints and we have “dependent” variables or the community. Constrained ordination rather is related to multivariate linear models.

The `vegan` package has three constrained ordination methods which all are constrained versions of basic ordination methods:

- Constrained analysis of proximities (CAP) in function `capscale` is related to metric scaling (`cmdscale`). It can handle any dissimilarity measures and performs a linear mapping.
- Redundancy analysis (RDA) in function `rda` is related to principal components analysis. It is based on Euclidean distances and performs linear mapping.
- Constrained correspondence analysis (CCA) in function `cca` is related to correspondence analysis. It is based on Chi-squared distances and performs weighted linear mapping.

We have already used functions `rda` and `cca` for unconstrained ordination: they will perform the basic unconstrained method as a special case if constraints are not used. All these three `vegan` functions are very similar. The following examples mainly use `cca`, but other methods can be used similarly. Actually, the results are similarly structured, and they inherit properties from each
other. For historical reasons, \texttt{cca} is the basic method, and \texttt{rda} inherits properties from it. Function \texttt{capscale} inherits directly from \texttt{rda}, and through this from \texttt{cca}. Many functions, are common with all these methods, and there are specific functions only if the method deviates from its ancestor. In \texttt{vegan} version 2.3-0 the following class functions are defined for these methods:

- \texttt{cca}: \texttt{add1}, \texttt{alias}, \texttt{anova}, \texttt{as.mlm}, \texttt{biplot}, \texttt{bstick}, \texttt{calibrate}, \texttt{coef}, \texttt{deviance}, \texttt{drop1}, \texttt{eigenvals}, \texttt{extractAIC}, \texttt{fitted}, \texttt{goodness}, \texttt{model.frame}, \texttt{model.matrix}, \texttt{nobs}, \texttt{permtest}, \texttt{plot}, \texttt{points}, \texttt{predict}, \texttt{print}, \texttt{residuals}, \texttt{RsquareAdj}, \texttt{scores}, \texttt{screeplot}, \texttt{simulate}, \texttt{stressplot}, \texttt{summary}, \texttt{text}, \texttt{tolerance}, \texttt{weights}
- \texttt{rda}: \texttt{as.mlm}, \texttt{biplot}, \texttt{coef}, \texttt{deviance}, \texttt{fitted}, \texttt{predict}, \texttt{RsquareAdj}, \texttt{scores}, \texttt{simulate}, \texttt{stressplot}, \texttt{weights}
- \texttt{capscale}: \texttt{fitted}, \texttt{print}, \texttt{RsquareAdj}, \texttt{simulate}, \texttt{stressplot}.

Many of these methods are internal functions that users rarely need.

### 4.1 Model specification

The recommended way of defining a constrained model is to use model formula. Formula has a special character `∼`, and on its left-hand side gives the name of the community data, and right-hand gives the equation for constraints. In addition, you should give the name of the data set where to find the constraints. This fits a \texttt{cca} for \texttt{varespec} constrained by soil Al, K and P:

```r
> vare.cca <- cca(varespec ~ Al + P + K, varechem)
> vare.cca
```

```text
Call: cca(formula = varespec ~ Al + P + K, data = varechem)

                 Inertia Proportion Rank
Total             2.083      1.000
Constrained        0.644      0.309      3
Unconstrained      1.439      0.691     20

Inertia is mean squared contingency coefficient

Eigenvalues for constrained axes:
CCA1  CCA2  CCA3
0.362  0.170  0.113

Eigenvalues for unconstrained axes:
CA1  CA2  CA3  CA4  CA5  CA6  CA7  CA8
0.350  0.220  0.185  0.155  0.135  0.100  0.077  0.054
(Showed only 8 of all 20 unconstrained eigenvalues)
```

The output is similar as in unconstrained ordination. Now the total inertia is decomposed into constrained and unconstrained components. There were three constraints, and the rank of constrained component is three. The rank of unconstrained component is 20, when it used to be 23 in the previous analysis. The rank is the same as the number of axes: you have 3 constrained axes and 20 unconstrained axes. In some
It is very common to calculate the proportion of constrained inertia from the total inertia. However, total inertia does not have a clear meaning in CCA, and the meaning of this proportion is just as obscure. In RDA this would be the proportion of variance (or correlation). This may have a clearer meaning, but even in this case most of the total inertia may be random noise. It may be better to concentrate on results instead of these proportions.

Basic plotting works just like earlier:

```r
> plot(vare.cca)
```

have similar interpretation as fitted vectors: the arrow points to the direction of the gradient, and its length indicates the strength of the variable in this dimensionality of solution. The vectors will be of unit length in full rank solution, but they are projected to the plane used in the plot.

The vegan3d provides a primitive 3D plotting function ordiplot3d (which needs user interaction for final graphs) that shows all arrows in full length:

```r
> library(vegan3d)
> ordiplot3d(vare.cca, type = "h")
```

With function ordigr1 you can also inspect 3D dynamic plots that can be spinned or zoomed into with your mouse.

The formula interface works with factor variables as well:

```r
> dune.cca <- cca(dune ~ Management, dune.env)
> plot(dune.cca)
```

```r
Call: cca(formula = dune ~ Management, data = dune.env)

Inertia Proportion Rank
Total 2.115 1.000
Constrained 0.604 0.285 3
Unconstrained 1.511 0.715 16
```

Inertia is mean squared contingency coefficient

Eigenvalues for constrained axes:

CCA1  CCA2  CCA3
0.319  0.182  0.103

Eigenvalues for unconstrained axes:

CA1  CA2  CA3  CA4  CA5  CA6  CA7  CA8  CA9  CA10
0.447  0.203  0.163  0.135  0.129  0.095  0.079  0.065  0.050  0.043
CA11  CA12  CA13  CA14  CA15  CA16
0.039  0.024  0.018  0.009  0.008  0.004

Factor variable Management had four levels (BF, HF, NM, SF). Internally R expressed these four levels as three contrasts (sometimes called “dummy variables”). The applied contrasts look like this:
We do not need but three variables to express four levels: if there is number one in a column, the observation belongs to that level, and if there is a whole line of zeros, the observation must belong to the omitted level, or the first. The basic `plot` function displays class centroids instead of vectors for factors.

In addition to these ordinary factors, R also knows ordered factors. Variable `Moisture` in `dune.env` is defined as an ordered four-level factor. In this case the contrasts look different:

<table>
<thead>
<tr>
<th>Moisture.L</th>
<th>Moisture.Q</th>
<th>Moisture.C</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>-0.6708</td>
<td>-0.2236</td>
</tr>
<tr>
<td>2</td>
<td>-0.2236</td>
<td>0.6708</td>
</tr>
<tr>
<td>4</td>
<td>0.2236</td>
<td>-0.6708</td>
</tr>
<tr>
<td>5</td>
<td>0.6708</td>
<td>0.2236</td>
</tr>
</tbody>
</table>

R uses polynomial contrasts: the linear term L is equal to treating Moisture as a continuous variable, and the quadratic Q and cubic C terms show nonlinear features. There were four distinct levels, and the number of contrasts is one less, just like with ordinary contrasts. The ordination configuration, eigenvalues or rank do not change if the factor is unordered or ordered, but the presentation of the factor in the results may change:

```r
> vare.cca <- cca(dune ~ Moisture, dune.env)
> plot(vare.cca)
```

Now `plot` shows both the centroids of factor levels and the contrasts. If we could change the ordered factor to a continuous vector, only the linear effect arrow would be important. If the response to the variable is nonlinear, the quadratic (and cubic) arrows would be long as well.

I have explained only the simplest usage of the formula interface. The formula is very powerful in model specification: you can transform your contrasts within the formula, you can define interactions, you can use polynomial contrasts etc. However, models with interactions or polynomials may be difficult to interpret.

### 4.2 Permutation tests

The significance of all terms together can be assessed using permutation tests: the dat are permuted randomly and the model is refitted. When constrained inertia in permutations is nearly always lower than observed constrained inertia, we say that constraints are significant.

The easiest way of running permutation tests is to use the mock `anova` function in `vegan`:

```r
> anova(vare.cca)
```

Permutation test for cca under reduced model
Permutation: free
Number of permutations: 999
4.2 Permutation tests

Model: cca(formula = dune ~ Moisture, data = dune.env)

<table>
<thead>
<tr>
<th>Df</th>
<th>ChiSquare</th>
<th>F</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model</td>
<td>3</td>
<td>0.628</td>
<td>2.25</td>
</tr>
<tr>
<td>Residual</td>
<td>16</td>
<td>1.487</td>
<td></td>
</tr>
</tbody>
</table>

---

Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

The Model refers to the constrained component, and Residual to the unconstrained component of the ordination, ChiSquare is the corresponding inertia, and Df the corresponding rank. The test statistic F, or more correctly “pseudo-F” is defined as their ratio. You should not pay any attention to its numeric values or to the numbers of degrees of freedom, since this “pseudo-F” has nothing to do with the real F, and the only way to assess its “significance” is permutation. In simple models like the one studied here we could directly use inertia in testing, but the “pseudo-F” is needed in more complicated model including “partialed” terms.

The number of permutations was not specified in the mock anova function. The function tries to be lazy: it continues permutations only as long as it is uncertain whether the final P-value will be below or above the critical value (usually P = 0.05). If the observed inertia is never reached in permutations, the function may stop after 200 permutations, and if it is very often exceeded, it may stop after 100 permutations. When we are close to the critical level, the permutations may continue to thousands. In this way the calculations are fast when this is possible, but they are continued longer in uncertain cases. If you want to have a fixed number of iterations, you must specify that in anova call or directly use the underlying function permute.test.cca

In addition to the overall test for all constraints together, we can also analyse single terms or axes by setting argument by. The following command analyses all terms separately in a sequential (“Type I”) test:

```r
mod <- cca(varespec ~ Al + P + K, varechem)
> anova(mod, by = "term", step=200)
```

Permutation test for cca under reduced model
Terms added sequentially (first to last)
Permutation: free
Number of permutations: 999

Model: cca(formula = varespec ~ Al + P + K, data = varechem)

<table>
<thead>
<tr>
<th>Df</th>
<th>ChiSquare</th>
<th>F</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>1</td>
<td>0.298</td>
<td>4.14</td>
</tr>
<tr>
<td>P</td>
<td>1</td>
<td>0.190</td>
<td>2.64</td>
</tr>
<tr>
<td>K</td>
<td>1</td>
<td>0.156</td>
<td>2.17</td>
</tr>
<tr>
<td>Residual</td>
<td>20</td>
<td>1.439</td>
<td></td>
</tr>
</tbody>
</table>

---

Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

All terms are compared against the same residuals, and there is no heuristic for the number permutations. The test is sequential, and the order of terms will influence the results, unless the terms are uncorrelated. In this case the same number of permutations will be used for all terms. The

\[ F = \frac{0.628/3}{1.487/16} = 2.254 \]
sum of test statistics (ChiSquare) for terms is the same as the Model test statistic in the overall test.

"Type III" tests analyse the marginal effects when each term is eliminated from the model containing all other terms:

```r
> anova(mod, by = "margin", perm=500)
```

Permutation test for cca under reduced model
Marginal effects of terms
Permutation: free
Number of permutations: 999

<table>
<thead>
<tr>
<th>Df</th>
<th>ChiSquare</th>
<th>F</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>1</td>
<td>0.312</td>
<td>4.33</td>
</tr>
<tr>
<td>P</td>
<td>1</td>
<td>0.168</td>
<td>2.34</td>
</tr>
<tr>
<td>K</td>
<td>1</td>
<td>0.156</td>
<td>2.17</td>
</tr>
<tr>
<td>Residual</td>
<td>20</td>
<td>1.439</td>
<td></td>
</tr>
</tbody>
</table>

---
Signif. codes:  0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

The marginal effects are independent of the order of the terms, but correlated terms will get higher ("worse") P-values. Now the sum of test statistics is not equal to the Model test statistic in the overall test, unless the terms are uncorrelated.

We can also ask for a test of individual axes:

```r
> anova(mod, by="axis", perm=1000)
```

Permutation test for cca under reduced model
Marginal tests for axes
Permutation: free
Number of permutations: 999

<table>
<thead>
<tr>
<th>Df</th>
<th>ChiSquare</th>
<th>F</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCA1</td>
<td>1</td>
<td>0.362</td>
<td>5.02</td>
</tr>
<tr>
<td>CCA2</td>
<td>1</td>
<td>0.170</td>
<td>2.36</td>
</tr>
<tr>
<td>CCA3</td>
<td>1</td>
<td>0.113</td>
<td>1.57</td>
</tr>
<tr>
<td>Residual</td>
<td>20</td>
<td>1.439</td>
<td></td>
</tr>
</tbody>
</table>

---
Signif. codes:  0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

4.3 Model building

It is very popular to perform constrained ordination using all available constraints simultaneously. Increasing the number of constraints actually means relaxing constraints: the ordination becomes more similar to the unconstrained one. When the rank of unconstrained component reduces towards zero, there are absolutely no constraints. However, the relaxation of constraints often happens much earlier in first ordination axes. If we do not have strict constraints, it may be better to use unconstrained ordination with vector fitting (or surface fitting), which allows detection of compositional variation for which we have not observed environmental
4.3 Model building

In constrained ordination it is best to reduce the number of constraints to just a few, say three to five.

I do not want to encourage using all possible environmental variables together as constraints. However, there still is a shortcut for that purpose in formula interface:

```r
> mod1 <- cca(varespec ~ ., varechem)
> mod1
```

```
Call: cca(formula = varespec ~ N + P + K + Ca + Mg + S + Al + Fe + Mn + Zn + Mo + Baresoil + Humdepth + pH, data = varechem)

Inertia Proportion Rank
Total 2.083 1.000
Constrained 1.441 0.692 14
Unconstrained 0.642 0.308 9
Inertia is mean squared contingency coefficient

Eigenvalues for constrained axes:
CCA1 CCA2 CCA3 CCA4 CCA5 CCA6 CCA7 CCA8 CCA9 CCA10
0.439 0.292 0.163 0.142 0.118 0.089 0.070 0.058 0.031 0.013
CCA11 CCA12 CCA13 CCA14
0.008 0.007 0.006 0.005

Eigenvalues for unconstrained axes:
CA1 CA2 CA3 CA4 CA5 CA6 CA7 CA8 CA9
0.1978 0.1419 0.1012 0.0708 0.0533 0.0333 0.0189 0.0151 0.0095

This result probably is very similar to unconstrained ordination:

```r
> plot(procrustes(cca(varespec), mod1))
```

For heuristic purposes we should reduce the number of constraints to find important environmental variables. In principle, constrained ordination only should be used with designed a priori constraints. All kind of automatic tools of model selection are dangerous: There may be several alternative models which are nearly equally good; Small changes in data can cause large changes in selected models; There may be no route to the best model with the adapted strategy; The model building has a history: one different step in the beginning may lead into wildly different final models; Significance tests are biased, because the model is selected for the best test performance.

After all these warnings, I show how vegan can be used to automatically select constraints into model using standard R function `step`. The `step` uses Akaike’s information criterion (AIC) as the selection criterion. AIC is a penalized goodness-of-fit measure: the goodness-of-fit is basically derived from the residual (unconstrained) inertia penalized by the rank of the constraints. In principle AIC is based on log-Likelihood that ordination does not have. However, a `deviance` function changes the unconstrained inertia to Chi-squared in `cca` or sum of squares in `rda` and `capscale`. This deviance is treated like sum of squares in Gaussian models. If we have only continuous (or 1 d.f.) terms, this is the same as selecting variables by their contributions to constrained eigenvalues (inertia). With factors the situation is more tricky, because factors must be
penalized by their degrees of freedom, and there is no way of knowing the magnitude of penalty. The `step` function may still be useful in helping to gain insight into the data, but it should not be trusted blindly (or at all), but only regarded as an aid in model building.

After this longish introduction the example: using `step` is much simpler than explaining how it works. We need to give the model we start with, and the scope of possible models inspected. For this we need another formula trick: formula with only 1 as the constraint defines an unconstrained model. We must define it like this so that we can add new terms to initially unconstrained model. The AIC used in model building is not based on a firm theory, and therefore we also ask for permutation tests at each step. In ideal case, all included terms should be significant and all excluded terms insignificant in the final model. The `scope` must be given as a list a formula, but we can extract this from fitted models using function `formula`. The following example begins with an unconstrained model `mod0` and steps towards the previously fitted maximum model `mod1`:

```r
> mod0 <- cca(varespec ~ 1, varechem)
> mod <- step(mod0, scope = formula(mod1), test = "perm")
```

Start: AIC=130.31

```r
varespec ~ 1

<table>
<thead>
<tr>
<th></th>
<th>Df</th>
<th>AIC</th>
<th>F</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>+ Al</td>
<td>1</td>
<td>128.61</td>
<td>3.67</td>
<td>0.005 **</td>
</tr>
<tr>
<td>+ Mn</td>
<td>1</td>
<td>128.95</td>
<td>3.31</td>
<td>0.005 **</td>
</tr>
<tr>
<td>+ Humdepth</td>
<td>1</td>
<td>129.24</td>
<td>3.00</td>
<td>0.005 **</td>
</tr>
<tr>
<td>+ Baresoil</td>
<td>1</td>
<td>129.77</td>
<td>2.46</td>
<td>0.020 *</td>
</tr>
<tr>
<td>+ Fe</td>
<td>1</td>
<td>129.79</td>
<td>2.44</td>
<td>0.020 *</td>
</tr>
<tr>
<td>+ P</td>
<td>1</td>
<td>130.03</td>
<td>2.19</td>
<td>0.030 *</td>
</tr>
<tr>
<td>+ Zn</td>
<td>1</td>
<td>130.30</td>
<td>1.92</td>
<td>0.060 .</td>
</tr>
<tr>
<td>&lt;none&gt;</td>
<td></td>
<td>130.31</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ Mg</td>
<td>1</td>
<td>130.35</td>
<td>1.87</td>
<td>0.075 .</td>
</tr>
<tr>
<td>+ K</td>
<td>1</td>
<td>130.37</td>
<td>1.86</td>
<td>0.050 *</td>
</tr>
<tr>
<td>+ Ca</td>
<td>1</td>
<td>130.43</td>
<td>1.79</td>
<td>0.060 .</td>
</tr>
<tr>
<td>+ pH</td>
<td>1</td>
<td>130.57</td>
<td>1.66</td>
<td>0.115</td>
</tr>
<tr>
<td>+ S</td>
<td>1</td>
<td>130.72</td>
<td>1.51</td>
<td>0.120</td>
</tr>
<tr>
<td>+ N</td>
<td>1</td>
<td>130.77</td>
<td>1.46</td>
<td>0.170</td>
</tr>
<tr>
<td>+ Mo</td>
<td>1</td>
<td>131.19</td>
<td>1.06</td>
<td>0.415</td>
</tr>
</tbody>
</table>
```

---

Signif. codes:  0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

Step: AIC=128.61

```r
varespec ~ Al

<table>
<thead>
<tr>
<th></th>
<th>Df</th>
<th>AIC</th>
<th>F</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>+ P</td>
<td>1</td>
<td>127.91</td>
<td>2.50</td>
<td>0.005 **</td>
</tr>
<tr>
<td>+ K</td>
<td>1</td>
<td>128.09</td>
<td>2.32</td>
<td>0.010 **</td>
</tr>
<tr>
<td>+ Zn</td>
<td>1</td>
<td>128.26</td>
<td>2.15</td>
<td>0.015 *</td>
</tr>
<tr>
<td>+ Mn</td>
<td>1</td>
<td>128.44</td>
<td>1.98</td>
<td>0.030 *</td>
</tr>
<tr>
<td>&lt;none&gt;</td>
<td></td>
<td>128.53</td>
<td>1.89</td>
<td>0.025 *</td>
</tr>
</tbody>
</table>
```

Step: AIC=128.61

```r
varespec ~ Al
```

25
4.3 Model building

### CONstrained ORDINATION

| + N         | 1 | 128.85 | 1.5900 | 0.065 |
| + Baresoil  | 1 | 128.88 | 1.5670 | 0.090 |
| + Ca        | 1 | 129.04 | 1.4180 | 0.190 |
| + Humdepth  | 1 | 129.08 | 1.3814 | 0.205 |
| + Mo        | 1 | 129.50 | 0.9884 | 0.435 |
| + pH        | 1 | 129.63 | 0.8753 | 0.575 |
| + Fe        | 1 | 130.02 | 0.5222 | 0.875 |
| - Al        | 1 | 130.31 | 3.6749 | 0.010 ** |

Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

Step: AIC=127.91

\[ \text{varespec} \sim \text{Al} + \text{P} \]

<table>
<thead>
<tr>
<th>Df</th>
<th>AIC</th>
<th>F</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>127.91</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ K</td>
<td>127.44</td>
<td>2.1688</td>
<td>0.040 *</td>
</tr>
<tr>
<td>&lt;none&gt;</td>
<td>127.91</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ Baresoil</td>
<td>127.99</td>
<td>1.6606</td>
<td>0.070</td>
</tr>
<tr>
<td>+ N</td>
<td>128.11</td>
<td>1.5543</td>
<td>0.135</td>
</tr>
<tr>
<td>+ S</td>
<td>128.36</td>
<td>1.3351</td>
<td>0.255</td>
</tr>
<tr>
<td>+ Mn</td>
<td>128.44</td>
<td>1.2641</td>
<td>0.280</td>
</tr>
<tr>
<td>+ Zn</td>
<td>128.51</td>
<td>1.2002</td>
<td>0.300</td>
</tr>
<tr>
<td>+ Humdepth</td>
<td>128.56</td>
<td>1.1536</td>
<td>0.320</td>
</tr>
<tr>
<td>- P</td>
<td>128.61</td>
<td>2.5001</td>
<td>0.020 *</td>
</tr>
<tr>
<td>+ Mo</td>
<td>128.75</td>
<td>0.9837</td>
<td>0.435</td>
</tr>
<tr>
<td>+ Mg</td>
<td>128.79</td>
<td>0.9555</td>
<td>0.430</td>
</tr>
<tr>
<td>+ pH</td>
<td>128.82</td>
<td>0.9247</td>
<td>0.555</td>
</tr>
<tr>
<td>+ Fe</td>
<td>129.28</td>
<td>0.5253</td>
<td>0.875</td>
</tr>
<tr>
<td>+ Ca</td>
<td>129.36</td>
<td>0.4648</td>
<td>0.895</td>
</tr>
<tr>
<td>- Al</td>
<td>130.03</td>
<td>3.9401</td>
<td>0.005 **</td>
</tr>
</tbody>
</table>

Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

Step: AIC=127.44

\[ \text{varespec} \sim \text{Al} + \text{P} + \text{K} \]

<table>
<thead>
<tr>
<th>Df</th>
<th>AIC</th>
<th>F</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>127.44</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ N</td>
<td>127.59</td>
<td>1.5148</td>
<td>0.115</td>
</tr>
<tr>
<td>+ Baresoil</td>
<td>127.67</td>
<td>1.4544</td>
<td>0.120</td>
</tr>
<tr>
<td>+ Zn</td>
<td>127.84</td>
<td>1.3067</td>
<td>0.290</td>
</tr>
<tr>
<td>+ S</td>
<td>127.89</td>
<td>1.2604</td>
<td>0.230</td>
</tr>
<tr>
<td>- K</td>
<td>127.91</td>
<td>2.1688</td>
<td>0.035 *</td>
</tr>
<tr>
<td>+ Mo</td>
<td>127.92</td>
<td>1.2350</td>
<td>0.280</td>
</tr>
<tr>
<td>- P</td>
<td>128.09</td>
<td>2.3362</td>
<td>0.025 *</td>
</tr>
<tr>
<td>+ Mg</td>
<td>128.17</td>
<td>1.0300</td>
<td>0.400</td>
</tr>
<tr>
<td>+ Mn</td>
<td>128.34</td>
<td>0.8879</td>
<td>0.545</td>
</tr>
<tr>
<td>+ Humdepth</td>
<td>128.44</td>
<td>0.8056</td>
<td>0.570</td>
</tr>
<tr>
<td>+ Fe</td>
<td>128.79</td>
<td>0.5215</td>
<td>0.920</td>
</tr>
<tr>
<td>+ pH</td>
<td>128.81</td>
<td>0.5067</td>
<td>0.855</td>
</tr>
<tr>
<td>+ Ca</td>
<td>128.89</td>
<td>0.4358</td>
<td>0.910</td>
</tr>
<tr>
<td>- Al</td>
<td>130.14</td>
<td>4.3340</td>
<td>0.005 **</td>
</tr>
</tbody>
</table>

Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1
> mod
Call: cca(formula = varespec ~ Al + P + K, data = varechem)

<table>
<thead>
<tr>
<th>Inertia Proportion Rank</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
</tr>
<tr>
<td>2.083</td>
</tr>
<tr>
<td>Constrained</td>
</tr>
<tr>
<td>0.644</td>
</tr>
<tr>
<td>Unconstrained</td>
</tr>
<tr>
<td>1.439</td>
</tr>
</tbody>
</table>

Inertia is mean squared contingency coefficient

Eigenvalues for constrained axes:
CCA1 CCA2 CCA3
0.362 0.170 0.113

Eigenvalues for unconstrained axes:
CA1 CA2 CA3 CA4 CA5 CA6 CA7 CA8
0.350 0.220 0.185 0.155 0.135 0.100 0.077 0.054
(Showed only 8 of all 20 unconstrained eigenvalues)

We ended up with the same familiar model we have been using all the time (and now you know the reason why this model was used in the first place). The AIC was based on deviance, and penalty for each added parameter was 2 per degree of freedom. At every step the AIC was evaluated for all possible additions (+) and removals (-), and the variables are listed in the order of AIC. The stepping stops when <none> or the current model is at the top.

Model building with step is fragile, and the strategy of model building can change the final model. If we start with the largest model (mod1), the final model will be different:

> modb <- step(mod1, scope = list(lower = formula(mod0), upper = formula(mod1)), trace = 0)
> modb
Call: cca(formula = varespec ~ P + K + Mg + S + Mn + Mo + Baresoil + Humdepth, data = varechem)

<table>
<thead>
<tr>
<th>Inertia Proportion Rank</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
</tr>
<tr>
<td>2.083</td>
</tr>
<tr>
<td>Constrained</td>
</tr>
<tr>
<td>1.117</td>
</tr>
<tr>
<td>Unconstrained</td>
</tr>
<tr>
<td>0.967</td>
</tr>
</tbody>
</table>

Inertia is mean squared contingency coefficient

Eigenvalues for constrained axes:
CCA1 CCA2 CCA3 CCA4 CCA5 CCA6 CCA7 CCA8
0.401 0.249 0.149 0.127 0.088 0.066 0.025 0.013

Eigenvalues for unconstrained axes:
CA1 CA2 CA3 CA4 CA5 CA6 CA7 CA8 CA9
0.2582 0.1881 0.1193 0.1020 0.0879 0.069 0.0446 0.0278 0.0269
CA10 CA11 CA12 CA13 CA14 CA15
0.0165 0.0136 0.0082 0.0066 0.0037 0.0024

The AIC of this model is 127.89 which is higher (worse) than reached in forward selection (127.44). We supressed tracing to save some pages of output, but step adds its history in the result:
4.4 Linear combinations and weighted averages

Variable Al was the first to be selected in the model in forward selection, but it was the second to be removed in backward elimination. Variable Al is strongly correlated with many other explanatory variables. This is obvious when looking at the variance inflation factors (vif) in the full model mod1:

```
> vif.cca(mod1)
   N     P     K     Ca     Mg     S     Al
Mn 9.128 5.380 7.740 4.320 2.254 6.013
Zn 9.128 5.380 7.740 4.320 2.254 6.013
Mo 9.128 5.380 7.740 4.320 2.254 6.013
Baresoil 9.128 5.380 7.740 4.320 2.254 6.013
Hundepth 9.128 5.380 7.740 4.320 2.254 6.013
pH 9.128 5.380 7.740 4.320 2.254 6.013
```

A common rule of thumb is that $vif > 10$ indicates that a variable is strongly dependent on others and does not have independent information. On the other hand, it may not be the variable that should be removed, but alternatively some other variables may be removed. The $vifs$ were all modest in model found by forward selection, including Al:

```
> vif.cca(mod)
   Al   P   K
Fe 1.012 2.365 2.379
```

4.4 Linear combinations and weighted averages

There are two kind of site scores in constrained ordinations:

1. Linear combination scores $lc$ which are linear combinations of constraining variables.

2. Weighted averages scores $wa$ which are weighted averages of species scores.

These two scores are as similar as possible, and their (weighted) correlation is called the species–environment correlation:

```
> spenvcor(mod)

CCA1  CCA2  CCA3
0.8555 0.8133 0.8793
```

Correlation coefficient is very sensitive to single extreme values, like seems to happen in the example above where axis 3 has the “best” correlation simply because it has some extreme points, and eigenvalue is a more appropriate measure of similarity between $lc$ and $wa$ scores.
The opinions are divided on using LC or WA as primary results in ordination graphics. The vegan package prefers WA scores, whereas the major commercial programme for CCA prefers LC scores. The vegan package comes with a separate document (“vegan FAQ”) which studies the issue in more detail, but I will briefly discuss the subject here also, and show how you can circumvent my decisions.

The practical reason to prefer WA scores is that they are more robust against random error in environmental variables. All ecological observations have random error, and therefore it is better to use scores that are resistant to this variation. Another point is that I see LC scores as constraints: the scores are dependent only on environmental variables, and community composition does not influence them. The WA scores are based on community composition, but so that they are as similar as possible to the constraints. This duality is particularly clear when using a single factor variable as constraint: the LC scores are constant within each level of the factor and fall in the same point. The WA scores show how well we can predict the factor level from community composition.

The vegan package has a graphical function ordispider which (among other alternatives) will combine WA scores to the corresponding LC score. With a single factor constraint:

```R
> dune.cca <- cca(dune ~ Management, dune.env)
> plot(dune.cca, display = c("lc", "wa"), type = "p")
> ordispider(dune.cca, col="blue")
```

The interpretation is similar as in discriminant analysis: LC scores give the predicted class centroids, and WA scores give the predicted values. For distinct classes, there is no overlap among groups. In general, the length of ordispider segments is a visual image of species–environment correlation.

### 4.5 Biplot arrows and environmental calibration

Biplot arrows are an essential part of constrained ordination plots. The arrows are based on (weighted) correlation of LC scores and environmental variables. They are scaled to unit length in the constrained ordination of full rank. When these arrows are projected onto 2D ordination plot, they look shorter if they go off the plane.

In vegan the biplot arrows are always scaled similarly irrespective of scaling of sites or species. With default `scaling = 2`, the biplot arrows have optimal relation to sites, but with `scaling = 1` they rather are related to species.

The standard interpretation of biplot arrows is that a site should be perpendicularly projected onto the arrow to predict the value of the variable. The arrow starts from the (weighted) mean of the environmental variable, and the values increase towards the arrow head, and decrease to the opposite direction. Then we still should figure out the unit of change. Function `calibrate.cca` performs this automatically in vegan. Let us inspect the result of the `step` function with three constraints:

```R
> pred <- calibrate(mod)
> head(pred)
```
4.6 Conditioned or partial models

The effect of some environmental variables can be removed from the ordination before constraining with other variables. The analysis is said to be conditioned on variables, or in other words, it is partial after removing variation caused by some variables. These conditioning variables typically are “random” or background variables, and their effect is removed from the analysis based on “fixed” or interesting variables.

In vegan, the formula for constrained ordination can contain a Condition which specifies the variable or variables whose effect is removed from the analysis before constraining with other variables. As an example, let us inspect what would be the effect of designed Management after removing the natural variation caused by Moisture:

```r
> dune.cca <- cca(dune ~ Management + Condition(Moisture), dune.env)
> plot(dune.cca)
> dune.cca
```

Actually, this is not based on biplot arrows, but on regression coefficients used internally in constrained ordination. Biplot arrows should only be seen as a visual approximation. The fitting is done in full constrained rank as default and for all constraints simultaneously. The example draws a residual plot of predictions:

```r
> with(varechem, plot(Al, pred[,"Al"] - Al, ylab="Prediction Error"))
> abline(h=0, col="grey")

The vegan package provides function ordisurf which is based on gam in the mgcv package, and can automatically detect the degree of smoothness needed, and can be used to check the linearity hypothesis of the biplot method. Function performs weighted fitting, and the model should be consistent with the one used in arrow fitting. Aluminium was the most important of three constraints in our example. Now we should fit the model to the LC scores, just like the arrows:

```r
> ef <- with(varechem, ordisurf(mod, Al, display = "lc", add = TRUE))
```

The results are not like we expected: we get curves instead of parallel lines perpendicular to the Al arrow. It seems that we cannot use linear projection in this case. Linear projection actually works, but only in the full constrained rank, or in three dimensions. When we project the multidimensional solution onto a plane, we get the distortion observed. Projections become unrealistic as soon as we have more than two constrained axes — but sometimes they may work quite well. In this case, P would display a linear response surface, although it was less important than Al in model building.

4.6 Conditioned or partial models

The effect of some environmental variables can be removed from the ordination before constraining with other variables. The analysis is said to be conditioned on variables, or in other words, it is partial after removing variation caused by some variables. These conditioning variables typically are “random” or background variables, and their effect is removed from the analysis based on “fixed” or interesting variables.

In vegan, the formula for constrained ordination can contain a Condition which specifies the variable or variables whose effect is removed from the analysis before constraining with other variables. As an example, let us inspect what would be the effect of designed Management after removing the natural variation caused by Moisture:

```r
> dune.cca <- cca(dune ~ Management + Condition(Moisture), dune.env)
> plot(dune.cca)
> dune.cca
```
4 CONSTRAINED ORDINATION

4.6 Conditioned or partial models

Call: cca(formula = dune ~ Management + Condition(Moisture), data = dune.env)

<table>
<thead>
<tr>
<th>Inertia</th>
<th>Proportion</th>
<th>Rank</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
<td>2.115</td>
<td>1.000</td>
</tr>
<tr>
<td>Conditional</td>
<td>0.628</td>
<td>0.297</td>
</tr>
<tr>
<td>Constrained</td>
<td>0.374</td>
<td>0.177</td>
</tr>
<tr>
<td>Unconstrained</td>
<td>1.113</td>
<td>0.526</td>
</tr>
</tbody>
</table>

Inertia is mean squared contingency coefficient

Eigenvalues for constrained axes:
CCA1  CCA2  CCA3
0.2278  0.0849  0.0614

Eigenvalues for unconstrained axes:
CA1  CA2  CA3  CA4  CA5  CA6  CA7  CA8  CA9  CA10  CA11  CA12  CA13
0.350  0.152  0.125  0.110  0.092  0.077  0.059  0.048  0.037  0.022  0.021  0.011  0.008

Now the total inertia is decomposed into three components: inertia explained by conditions, inertia explained by constraints and the remaining unconstrained inertia. We previously fitted a model with Management as the only constraint, and in that case constrained inertia was clearly higher than now. It seems that different Management was practised in different natural conditions, and the variation we previously attributed to Management may be due to Moisture.

We can perform permutation tests for Management in conditioned model, and Management alone:

> anova(dune.cca, perm.max = 2000)

Permutation test for cca under reduced model
Permutation: free
Number of permutations: 999

Model: cca(formula = dune ~ Management + Condition(Moisture), data = dune.env)
   Df  ChiSquare   F Pr(>F)
Model  3 0.374 1.46 0.034 *
Residual 13 1.113

---
Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

> anova(cca(dune ~ Management, dune.env))

Permutation test for cca under reduced model
Permutation: free
Number of permutations: 999

Model: cca(formula = dune ~ Management, data = dune.env)
   Df ChiSquare   F Pr(>F)
Model  3 0.604 2.13 0.001 ***
Residual 16 1.511

---
Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1
Inspected alone, Management seemed to be very significant, but the situation is much less clear after removing the variation due to Moisture.

The anova function (like any permutation test in vegan) can be restricted so that permutation are made only within strata or within a level of a factor variable:

```r
> with(dune.env, anova(dune.cca, strata = Moisture))
```

Permutation test for cca under reduced model
Blocks: strata
Permutation: free
Number of permutations: 999

<table>
<thead>
<tr>
<th>Df</th>
<th>ChiSquare</th>
<th>F</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model</td>
<td>3</td>
<td>0.374</td>
<td>1.46</td>
</tr>
<tr>
<td>Residual</td>
<td>13</td>
<td>1.113</td>
<td>---</td>
</tr>
</tbody>
</table>

Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

Conditioned or partial models are sometimes used for decomposition of inertia into various components attributed to different sets of environmental variables. In some cases this gives meaningful results, but the groups of environmental variables should be non-linearly independent for unbiased decomposition. If the groups of environmental variables have polynomial dependencies, some of the components of inertia may even become negative (that should be impossible). That kind of higher-order dependencies are almost certain to appear with high number of variables and high number of groups. However, varpart performs decomposition of rda models among two to four components.

5 Dissimilarities and environment

We already discussed environmental interpretation of ordination and environmentally constrained ordination. These both reduce the variation into an ordination space, and mainly inspect the first dimensions. Sometimes we may wish to analyse vegetation–environment relationships without ordination, or in full space. Typically these methods use the dissimilarity matrix in analysis. The recommended method in vegan is adonis which implements a multivariate analysis of variances using distance matrices. Function adonis can handle both continuous and factor predictors. Other methods in vegan include multiresponse permutation procedure (mrpp) and analysis of similarities (anosim). Both of these handle only class predictors, and they are less robust than adonis.

5.1 adonis: Multivariate ANOVA based on dissimilarities

Function adonis partitions dissimilarities for the sources of variation, and uses permutation tests to inspect the significances of those partitions. With Euclidean distances the results are similar as in rda and its anova permutation tests, but adonis can handle any dissimilarity objects.
The example uses `adonis` to study beta diversity between Management classes in the dune meadow data. We define beta diversity as the slope of species-area curve, or the exponent \( z \) of the Arrhenius model where the number of species \( S \) is dependent on the size \( X \) of the study area. For pairwise comparison of sites the slope \( z \) can be found from the number of species shared between two sites \( (a) \) and the number of species unique to each sites \( (b \text{ and } c) \). It is commonly regarded that \( z \approx 0.3 \) implies random sampling variability, and only higher values mean real systematic differences. The Arrhenius \( z \) can be directly found with function `betadiver` that also provided many other indices of pairwise beta diversity.

\[
S = kX^z \\
z = \left[ \log(2) - \log(2a + b + c) + \log(a + b + c) \right] / \log(2)
\]

> betad <- betadiver(dune, "z")

Function `adonis` can use formula interface, and the dependent data can be either dissimilarities or data frame, and in the latter case `adonis` uses `vegdist` to find the dissimilarities.

> adonis(betad ~ Management, dune.env, perm=200)

Call:
adonis(formula = betad ~ Management, data = dune.env, permutations = 200)

Permutation: free
Number of permutations: 200

Terms added sequentially (first to last)

<table>
<thead>
<tr>
<th>Df</th>
<th>SumsOfSqs</th>
<th>MeanSqs</th>
<th>F.Model</th>
<th>R2</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Management 3</td>
<td>1.24</td>
<td>0.412</td>
<td>2.36</td>
<td>0.307</td>
<td>0.01 **</td>
</tr>
<tr>
<td>Residuals 16</td>
<td>2.79</td>
<td>0.174</td>
<td>0.693</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total 19</td>
<td>4.03</td>
<td>1.000</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

The models can be more complicated, and sequential test of permutational ANOVA is performed if there are several parameters:

> adonis(betad ~ A1*Management, dune.env, perm = 200)

Call:
adonis(formula = betad ~ A1 * Management, data = dune.env, permutations = 200)

Permutation: free
Number of permutations: 200

Terms added sequentially (first to last)

<table>
<thead>
<tr>
<th>Df</th>
<th>SumsOfSqs</th>
<th>MeanSqs</th>
<th>F.Model</th>
<th>R2</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1 1</td>
<td>0.65</td>
<td>0.655</td>
<td>4.13</td>
<td>0.005 **</td>
<td></td>
</tr>
<tr>
<td>Management 3</td>
<td>1.00</td>
<td>0.334</td>
<td>2.11</td>
<td>0.055 .</td>
<td></td>
</tr>
<tr>
<td>A1:Management 3</td>
<td>0.47</td>
<td>0.156</td>
<td>0.99</td>
<td>0.117 0.458</td>
<td></td>
</tr>
<tr>
<td>Residuals 12</td>
<td>1.90</td>
<td>0.158</td>
<td>0.472</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total 19</td>
<td>4.03</td>
<td>1.000</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1
5.2 Homogeneity of groups and beta diversity

Function `adonis` studied the differences in the group means, but function `betadisper` studies the differences in group homogeneities. Function `adonis` was analogous to multivariate analysis of variance, and `betadisper` is analogous to Levene’s test of the equality of variances.

The example continues the analysis of the previous section and inspects the beta diversity. The function can only use one factor as an independent variable, and it does not know the formula interface, so that we need to `attach` the data frame or use `with` to make the factor visible to the function:

```r
> mod <- with(dune.env, betadisper(betad, Management))
> mod
```

```
Call: betadisper(d = betad, group = Management)

No. of Positive Eigenvalues: 12
No. of Negative Eigenvalues: 7

Average distance to median:
  BF  HF  NM  SF
0.308 0.251 0.441 0.363

Eigenvalues for PCoA axes:
  PCoA1  PCoA2  PCoA3  PCoA4  PCoA5  PCoA6  PCoA7  PCoA8
1.655  0.887  0.533  0.374  0.287  0.224  0.161  0.081

The function has `plot` and `boxplot` methods for graphical display.
```

```r
> plot(mod)
> boxplot(mod)
```

The significance of the fitted model can be analysed either using standard parametric `anova` or permutation tests (`permutest`):

```r
> anova(mod)
> permutest(mod)
```

```
Analysis of Variance Table

Response: Distances
  Df  Sum Sq  Mean Sq   F value  Pr(>F)
Groups   3 0.1048  0.03494 1.260000 0.3241
Residuals 16 0.4428  0.02770

Permutation test for homogeneity of multivariate dispersions
  Permutation: free
  Number of permutations: 999

Response: Distances
  Df  Sum Sq  Mean Sq F  N.Perm Pr(>F)
Groups   3 0.1048  0.03494 1.26  999    0.315
Residuals 16 0.4428  0.02770
```

34
Moreover, it is possible to analyse pairwise differences between groups using parametric Tukey’s HSD test:

\[
> \text{TukeyHSD(mod)}
\]

Tukey multiple comparisons of means
95% family-wise confidence level

Fit: \(\text{aov(formula = distances ~ group, data = df)}\)

\[
\begin{array}{cccc}
\text{group} & \text{diff} & \text{lwr} & \text{upr} & \text{p adj} \\
\text{HF-BF} & -0.05682 & -0.40452 & 0.2909 & 0.9651 \\
\text{NM-BF} & 0.13256 & -0.20409 & 0.4692 & 0.6791 \\
\text{SF-BF} & 0.05547 & -0.28119 & 0.3921 & 0.9643 \\
\text{NM-HF} & 0.18938 & -0.09891 & 0.4777 & 0.2752 \\
\text{SF-HF} & 0.11229 & -0.17600 & 0.4006 & 0.6862 \\
\text{SF-NM} & -0.07709 & -0.35197 & 0.1978 & 0.8523 \\
\end{array}
\]

### 5.3 Mantel test

Mantel test compares two sets of dissimilarities. Basically, it is the correlation between dissimilarity entries. As there are \(N(N-1)/2\) dissimilarities among \(N\) objects, normal significance tests are not applicable. Mantel developed asymptotic test statistics, but \textit{vegan} function \texttt{mantel} uses permutation tests.

In this example we study how well the lichen pastures (\texttt{varespec}) correspond to the environment. We have already used vector fitting after ordination. However, the ordination and environment may be non-linearly related, and we try now with function \texttt{mantel}. We first perform a \texttt{PCA} of environmental variables, and then compute dissimilarities for first principal components. We use standard R function \texttt{prcomp}, but \texttt{princomp} or \texttt{rda} will work as well. Function \texttt{scores} in \textit{vegan} will work with all these methods. The following uses the same standardizations for community dissimilarities as previously used in \texttt{metaMDS}.

\[
> \text{pc <- prcomp(varechem, scale = TRUE)}
\]
\[
> \text{pc$\text{scores}(pc, display = \"sites\", choices = 1:4)}
\]
\[
> \text{edis <- vegdist(pc, method = \"euclid\")}
\]
\[
> \text{vare.dis <- vegdist(wisconsin(sqrt(varespec))))}
\]
\[
> \text{mantel(vare.dis, edis)}
\]

Mantel statistic based on Pearson’s product-moment correlation

Call:
\[
\text{mantel(xdis = vare.dis, ydis = edis)}
\]

Mantel statistic \(r:\ 0.381\)

Significance: 0.001

Upper quantiles of permutations (null model):
90% 95% 97.5% 99%
0.147 0.186 0.218 0.250

Permutation: free
Number of permutations: 999
5.4 Protest: Procrustes test

We could use a selection of environmental variables in PCA, or we could use standardized environmental variables directly without PCA — tastes vary. Function bioenv gives an intriguing alternative for selecting optimal subsets for comparing ordination and environment. There also is a partial Mantel test where we can remove the influence of third set dissimilarities from the analysis, but its results often are difficult to interpret.

Function mantel does not have diagnostic plot functions, but you can directly plot two dissimilarity matrices against each other:

```r
> plot(vare.dis, edis)
```

Everything is O.K. if the relationship is more or less monotonous, or even linear and positive. In spatial models we even may observe a hump which indicates spatial aggregation.

5.4 Protest: Procrustes test

Procrustes test or protest compares two ordinations using symmetric Procrustes analysis. It is an alternative to Mantel tests, but uses reduced space instead of complete dissimilarity matrices. We can repeat the previous analysis, but now with the solution of metaMDS and two first principal components of the environmental analysis:

```r
> pc <- scores(pc, choices = 1:2)
> pro <- protest(vare.mds, pc)
> plot(pro)
> pro
```

Call:
protest(X = vare.mds, Y = pc)

Procrustes Sum of Squares (m2 squared): 0.533
Correlation in a symmetric Procrustes rotation: 0.683
Significance: 0.001

Permutation: free
Number of permutations: 999

The significance is assessed by permutation tests. The statistic is now Procrustes correlation \( r \) derived from the symmetric Procrustes residual \( m^2 \). The correlation is clearly higher than the Mantel correlation for the corresponding dissimilarities. In lower number of dimensions we remove noise from the data which may explain higher correlations (as well as different methods of calculating the correlation). However, both methods are about as significant. Significance often is not a significant concept: even small deviations from randomness may be highly significant in large data sets. Function protest provides graphical presentations (“Procrustes superimposition plot”) which may be more useful in evaluating the congruence between configurations.

PROTEST (as ordinary Procrustes analysis) is often used in assessing similarities between different community ordinations. This is known as analysis of congruence.
6 Classification

The vegan mainly is a package for ordination and diversity analysis, and there is only a scanty support to classification. There are several other R packages with more extensive classification functions. Among community ecological packages, labdsv package by Dave Roberts is particularly strong in classification functions.

This chapter describes performing simple classification tasks in community ecology that are sufficient to many community ecologists.

6.1 Cluster analysis

Hierarchic clustering can be performed using standard R function hclust. In addition, there are several other clustering packages, some of which may be compatible with hclust. Function hclust needs a dissimilarities as input.

Function hclust provides several alternative clustering strategies. In community ecology, most popular are single linkage a.k.a. nearest neighbour, complete linkage a.k.a. furthest neighbour, and various brands of average linkage methods. These are best illustrated with examples:

```r
> dis <- vegdist(dune)
> clus <- hclust(dis, "single")
> plot(clus)
```

Some people prefer single linkage, because it is conceptually related to minimum spanning tree which nicely can be represented in ordinations, and it is able to find discontinuities in the data. However, single linkage is prone to chain data so that single sites are joined to large clusters.

```r
> cluc <- hclust(dis, "complete")
> plot(cluc)
```

Some people prefer complete linkage because it makes compact clusters. However, this is in part an artefact of the method: the clusters are not allowed to grow, because the complete linkage criterion would be violated.

```r
> clua <- hclust(dis, "average")
> plot(clua)
```

Some people (I included) prefer average linkage clustering, because it seems to be a compromise between the previous two extremes, and more neutral in grouping. There are several alternative methods loosely connected to "average linkage" family. Ward’s method seems to be popular in publications. It approaches complete linkage in its attempt to minimize variances in agglomeration. The default "average" method is the one often known as UPGMA which was popular in old-time genetics.

All these clustering methods are agglomerative. They start with combining two most similar sites to each other. Then they proceed by combining points to points or to groups, or groups to groups. The fusion criteria vary. The vertical axis in all graphs shows the level of fusion. The numbers vary among methods, but all are based on the same dissimilarities with range:

```r
> range(dis)
```
The first fusion is between the same two most similar sites in all examples, and at the same minimum dissimilarity. In complete linkage the last fusion combines the two most dissimilar sites, and it is at the maximum dissimilarity. In single linkage the fusion level always is at the smallest gap between groups, and the reported levels are much lower than with complete linkage. Average linkage makes fusions between group centre points, and its fusion levels are between the previous two trees. The estimated dissimilarity between two points is the level where they are fused in a tree. Function \texttt{cophenetic} finds this estimated dissimilarity from a tree for every pair of points – the name of the function reflects the history of clustering in numerical taxonomy. Cophenetic correlation measures the similarity between original dissimilarities and dissimilarities estimated from the tree. For our three example methods:

\begin{verbatim}
> cor(dis, cophenetic(clus))
[1] 0.6602
> cor(dis, cophenetic(cluc))
[1] 0.6707
> cor(dis, cophenetic(clua))
[1] 0.8169
\end{verbatim}

Approximating dissimilarities is the same task that ordinations perform, and average linkage is the best performer.

6.2 Display and interpretation of classes

Cluster analysis performs a hierarchic clustering, and its results can be inspected at as many levels as there are points: the extremes are that every point is in its private cluster, or that all points belong to the same cluster. We commonly want to inspect clustering at a certain level, as a non-hierarchic system of certain number of clusters. The flattening of the clustering happens by cutting the tree at some fusion level so that we get a desired number of clusters.

Base R provides function \texttt{rect.hclust} to visualize the cutting, and function \texttt{cutree} to make a classification vector with certain number of classes:

\begin{verbatim}
> plot(cluc)
> rect.hclust(cluc, 3)
> grp <- cutree(cluc, 3)
\end{verbatim}

The classification vector can be used as any other factor variable. A natural way of inspecting the goodness of community classification is to see how well it predicts external environmental variables that were not used in clustering. The only continuous variable in the Dune data is the thickness of the A1 horizon:

\begin{verbatim}
> boxplot(A1 ~ grp, data=dune.env, notch = TRUE)
\end{verbatim}
If we wish, we may use all normal statistical methods with factors, such as functions `lm` or `aov` for formal testing of “significance” of clusters. Classification can be compared against external factor variables as well. However, `vegan` does not provide any tools for this. It may be best to see the labdsv package and its tutorial for this purpose.

The clustering results can be displayed in ordination diagrams. All usual `vegan` functions for factors can be used: `ordihull`, `ordispider`, and `ordiellipse`. We shall see only the first as an example:

```r
> ord <- cca(dune)
> plot(ord, display = "sites")
> ordihull(ord, grp, lty = 2, col = "red")
```

It is said sometimes that overlaying classification in ordination can be used as a cross-check: if the clusters look distinct in the ordination diagram, (both) analyses probably were adequate. However, the classes can overlap and the analyses can still be good. It may be that you need three or more axes to display the multivariate class structure. In addition, ordination and classification may use different criteria. In our example, CA uses weighted Chi-squared criteria, and the clustering uses Bray–Curtis dissimilarities which may be quite different. Function `ordirgl` with its support function `orglspider` can be used to inspect classification using dynamic 3D graphics.

The `vegan` package has function `ordicluster` to overlay `hclust` tree in an ordination:

```r
> plot(ord, display="sites")
> ordicluster(ord, cluc, col="blue")
```

The function combines points and cluster midpoints similarly as in the original cluster dendrogram.

Single linkage clustering is the method most often used with ordination diagrams. Single linkage clustering is special among the clustering algorithms, because it always combines points to points: it is only the nearest point that is recognized and no information on its cluster membership is used. The dendrogram, however, hides this information: it only shows the fusions between clusters, but does not show which were the actual points that were joined. The tree connecting individual points is called a minimum spanning tree (MST). In graph theory, ‘tree’ is a connected graph with no loops, ‘spanning tree’ is tree that connects all points, and minimum spanning tree is the one where the total length of connecting segments is shortest. Function `spantree` in `vegan` find this tree, and it has a `lines` function to overlay the tree onto ordination:

```r
> mst <- spantree(dis, toolong = 1)
> plot(mst, ord=ord, pch=21, col = "red", bg = "yellow", type = "t")
```

In our dissimilarity index, distance = 1 means that there is nothing in common with two sample plots. Function `spantree` regards these maximum dissimilarities as missing data, and does not use them in building the tree. If all points cannot be connected because of these missing values, the result will consist of disconnected spanning trees. In graph theory this is known as a ‘forest’. `MST` is used sometimes to cross-check ordination: if the tree is linear, the ordination might be good. A curved tree may
indicate arc or horseshoe artefacts, and a messy tree a bad ordination, or a need of higher number of dimensions. However, the results often are difficult to interpret.

### 6.3 Classified community tables

The aim of classification often is to make a classified community table. For this purpose, both sites and species should be arranged so that the table looks structured. The original clustering may not be ideally structured, because the ordering of sites is not strictly defined in the cluster dendrogram. You can take any branch and rotate it around its base, and the clustering is the same. The tree drawing algorithms use heuristic rules to make the tree look aesthetically pleasing, but this ordering may not be the best one for a structured community table.

Base R has a general tree class called `dendrogram` which is intended as a common base for any tree-like presentations. This class has a function to reorder a tree according to some external variable. The `hclust` result can be changed into `dendrogram` with function `as.dendrogram`, and this can be reordered with function `reorder`. The only continuous variable in the Dune data is the thickness of A1 horizon, and this could be used to arrange the tree. However, for a nicely structured community table we use another trick: CA is an ordination method that structures table optimally into a diagonal structure, and we can use its first axis to reorder the tree:

```r
> wa <- scores(ord, display = "sites", choices = 1)
> den <- as.dendrogram(clua)
> oden <- reorder(den, wa, mean)
```

The results really change, and it may take some effort to see that these two trees really are identical, except for the order of leaves.

```r
> op <- par(mfrow=c(2,1), mar=c(3,5,1,2)+.1)
> plot(den)
> plot(oden)
> par(op)
```

Function `vegemite` in vegan produces compact vegetation tables. It can take an argument `use` to arrange the sites (and species, if possible). This argument can be a vector used to arrange sites, or it can be an ordination result, or it can be an `hclust` result or a `dendrogram` object.

```r
> vegemite(dune, use = oden, zero = "-")
```

<table>
<thead>
<tr>
<th>Airaprae</th>
<th>23-1111111</th>
</tr>
</thead>
<tbody>
<tr>
<td>Empenigr</td>
<td>-2-11111111</td>
</tr>
<tr>
<td>Hyporadi</td>
<td>25-2222222</td>
</tr>
<tr>
<td>Anthodor</td>
<td>44-1111111</td>
</tr>
<tr>
<td>Trifprat</td>
<td>2222222222</td>
</tr>
<tr>
<td>Achimill</td>
<td>2-2222222222</td>
</tr>
<tr>
<td>Planlanc</td>
<td>2-2222222222</td>
</tr>
<tr>
<td>Rumeacet</td>
<td>2-2222222222</td>
</tr>
<tr>
<td>Bromhord</td>
<td>2-2222222222</td>
</tr>
<tr>
<td></td>
<td>79157602183498234506</td>
</tr>
<tr>
<td></td>
<td>Airaprae 23-1111111</td>
</tr>
<tr>
<td></td>
<td>Empenigr -2-11111111</td>
</tr>
<tr>
<td></td>
<td>Hyporadi 25-2222222</td>
</tr>
<tr>
<td></td>
<td>Anthodor 44-1111111</td>
</tr>
<tr>
<td></td>
<td>Trifprat 2222222222</td>
</tr>
<tr>
<td></td>
<td>Achimill 2-2222222222</td>
</tr>
<tr>
<td></td>
<td>Planlanc 2-2222222222</td>
</tr>
<tr>
<td></td>
<td>Rumeacet 2-2222222222</td>
</tr>
<tr>
<td></td>
<td>Bromhord 2-2222222222</td>
</tr>
</tbody>
</table>
Lolipere --72665726524------
Bellipere --2--23--222--------
Vicilath --------1-21----------
Elymrepe --44--4--446--------
Poaprat 1-424344435444-2----
Scorautu 26-333355223222222-
Trifrepe -2-2256321323261--
Poatriv --265447--655449-----2
Bracrutu -3-2262-4622224----444
Sagiproc -3--------2--52242----
Salirepe -3--------3----------5-
Cirscarv ---------------2--------
Juncbufo ----2-------4-43----
Alopgeni --------2--723855--4
Agrostol --------4834545457
Chenalbu ---------------1----
Juncartt -------------44--343
Comapalu ----------------22--
Ranuflam -------------2-22242
Eleopalu -------------4--4548
Callcusp ----------------4-33

<table>
<thead>
<tr>
<th>sites</th>
<th>species</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>30</td>
</tr>
</tbody>
</table>

The dendrogram had no information on species, but it uses weighted averages to arrange them similarly as sites. This may not be optimal for a clustering results, but if the clusters are reordered nicely, the results may be very satisfactory with a nicely structured community table.

The vegemite output is very compact (hence the name), and it uses only one column for sites. In this case this was automatic, since Dune meadow data uses class scales. Percent cover scale can be transformed to traditional class scales, such as Braun-Blanquet, Domin or Hult–Sernander–Du Rietz.

**Session Info**

- R Under development (unstable) (2015-06-09 r68498),
  x86_64-unknown-linux-gnu
- Locale: LC_CTYPE=en_IE.UTF-8, LC_NUMERIC=C, LC_TIME=en_IE.UTF-8, LC_COLLATE=en_IE.UTF-8, LC_MONETARY=en_IE.UTF-8, LC_MESSAGES=en_IE.UTF-8, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_IE.UTF-8, LC_IDENTIFICATION=C
- Base packages: base, datasets, graphics, grDevices, methods, stats, utils
- Other packages: lattice 0.20-31, MASS 7.3-40, permute 0.8-3, vegan 2.3-0, veganId 0.65-0
- Loaded via a namespace (and not attached): cluster 2.0.1, grid 3.3.0, Matrix 1.2-0, mgcv 1.8-6, nlme 3.1-120, parallel 3.3.0, rgl 0.95.1247, scatterplot3d 0.3-35, tools 3.3.0
Index

A
adonis, 32–34
AIC, 24, 25, 27
anosim, 32
anova, 21, 22, 32, 34
aov, 39
apply, 9
arc effect, 11, 12
as.dendrogram, 40
attach, 16, 34

B
beta diversity, 7, 33, 34
betadisper, 34
betadiver, 7, 33
bioenv, 36
biplot, 9

C
calibrate.cca, 29
capscale, 18, 19, 24
cca, 9, 12, 18, 19, 24
clustering, 37–39
cmdscale, 3, 18
constrained analysis of proximities, 18
constrained correspondence analysis, 18, 20
constrained ordination, 18, 19, 21, 23, 28
contrasts, 20, 21
cophenetic, 38
correspondence analysis, 8, 10–12, 18, 40
cutree, 38

daisy, 7
decorana, 11–13
decostand, 7
dendrogram, 40, 41
designdist, 7
detrended correspondence analysis, 11, 12
deviance, 24
dissimilarity
  Arrhenius, 33
  binomial, 6
  Bray-Curtis, 3, 5, 6
  Canberra, 6
  Chi-square, 8, 10, 18
  chord, 7
  Czekanowski, 6
  Euclidean, 5–8, 18, 32
  Gower, 6
  Hellinger, 7
  Horn-Morisita, 6
  Jaccard, 5, 6
  Kulczyński, 5
  Manhattan, 5
  metric properties, 6
  Morisita, 6
  Mountford, 6
  Raup-Crick, 6
  Ružička, 6
  semimetric, 6
  Sørensen, 6, 7
  Steinhaus, 3, 6
dist, 5, 7
distance, 7
downweight, 12
downweighting, 11, 12
dsvdis, 7

e
envfit, 15, 16, 18

F
factor fitting, 16
formula, 16, 19, 21, 24, 25

g
gam, 16, 30

H
half-change scaling, 5
hclust, 37, 39, 40

I
identify, 8, 13, 14
inertia, 9–11, 19–22, 24
isoMDS, 3–5, 8

L
LC scores, 28–30
Levene’s test, 34
lm, 39
local optimum, 4

M
make.cepnames, 13
mantel, 35, 36
Mantel test, 35, 36
partial, 36
metaMDS, 4–6, 8, 36
metric scaling, 3, 8, 18
minimum spanning tree, 39
mrpp, 32

N
non-metric multidimensional scaling, 3–8, 15

O
ordered factors, 21
ordi.cluster, 39
ordi.ellipse, 18, 39
ordihull, 17, 39
ordi.label, 13, 14
ordi.plot, 4, 12, 13
ordi.plot3d, 20
ordi.plot.label, 14
ordi.gl, 20, 39
ordi.spider, 18, 29, 39
ordi.surf, 16, 18, 30
ordi.tkplot, 14
ordi.tor, 13, 14
ordi.g.lspider, 39

P
package
analogue, 7
cluster, 7
labdsv, 7, 37, 39
MASS, 3
mgcv, 16, 30
partial ordination, 30
permutation tests, 21, 33–36
permute.test, 34
permute.test.cca, 22
points, 12, 14
prcomp, 8, 35
principal components analysis, 8, 9, 12, 35
principal coordinates analysis, 8
princomp, 8, 35
procrustes, 8
Procrustes analysis, 36
Procrustes rotation, 8
protest, 36

R
rankindex, 6
rda, 9, 18, 19, 24, 35
rect.hclust, 38
redundancy analysis, 18, 20
reorder, 40

S
scores, 4
Shepard, 3
spantree, 39
species space, 5
species–environment correlation, 28, 29
splines, 16
standardization
Hellinger, 7
norm, 7
Wisconsin, 4, 7
step, 24, 25, 27, 29
stress, 3
stressplot, 3
surface fitting, 16

T
Tcl/Tk, 14
text, 12, 14
transformation
square root, 4, 7
Tukey’s HSD, 35

V
var, 9
variance inflation factor, 28
varpart, 32
vector fitting, 14
vegdist, 3, 5–7, 33
vegemite, 40, 41

W
WA scores, 28, 29
weighted averages, 5, 11, 28
wisconsin, 7
with, 16, 34