Training in ade4 in R - Module I: Basic methods

Principal component analysis

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Data

We will analyze the doubs data set (see ?doubs)

```
library(ade4)
library(adegraphics)
data(doubs)
names(doubs)

## [1] "env"  "fish"  "xy"  "species"

names(doubs$env)

## [1] "dfs" "alt" "slo" "flo" "pH"  "har" "pho" "nit" "amm" "oxy" "bdo"
```

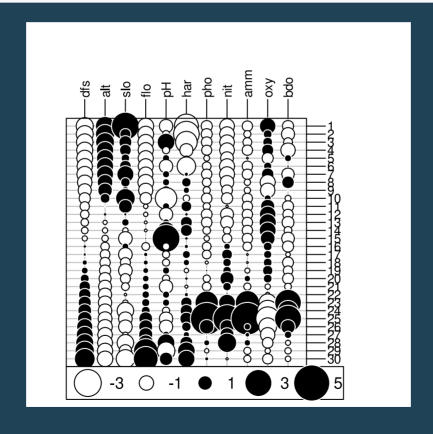
Some general information

```
env <- doubs$env
apply(env, 2, range)</pre>
```

```
## dfs alt slo flo pH har pho nit amm oxy bdo
## [1,] 3 172 1.099 84 77 40 1 15 0 41 13
## [2,] 4530 934 6.176 6900 86 110 422 620 180 124 167
```

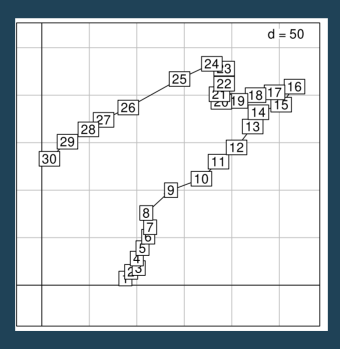
Display the data

```
table.value(scale(env), symbol = "circle")
```



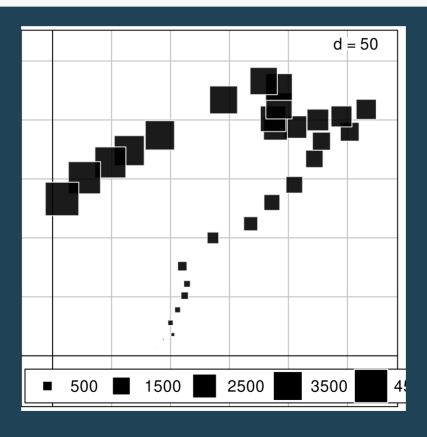
Spatial information

```
xy <- doubs$xy
sl1 <- s.label(xy, ppoints.cex = 0, plot = FALSE)
st1 <- s.traject(xy, ppoints.cex = 0, plabels.cex = 0,
    plot = FALSE)
s1 <- superpose(st1, sl1, plot = TRUE)</pre>
```



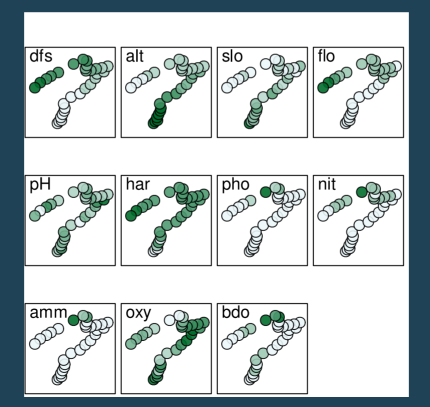
Map one variable

s.value(xy, doubs\$env[, 1])



Map the different variables

```
mypal <- colorRampPalette(c("#EDF8FB", "#006D2C"))
s.value(xy, doubs$env, pgrid.draw = FALSE, porigin.draw = FALSE,
    plegend.drawKey = FALSE, psub.cex = 2, method = "color",
    symbol = "circle", ppalette.quanti = mypal, ppoints.cex = 0.5)</pre>
```



Perform the PCA

Must we scale or not the data?

```
pca1 <- dudi.pca(env, scale = TRUE, scannf = FALSE,
    nf = 2)</pre>
```

How many axes?

Choosing the number of dimensions that can be interpreted is an open question. Several methods have been proposed in the literature. Using your eyes to detect breaks in eigenvalue decrease is probably the best method.

However, ade4 proposes one method developed for the case of PCA on scaled data. You can try it:

testdim(pca1)\$nb.cor

[1] 2

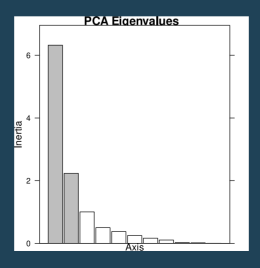
Display eigenvalues

- Make a barplot of eigenvalues
- Compute the percentage of variation explained by each dimension
- Compare to the results obtained by the function summary

```
round(pca1$eig/sum(pca1$eig) * 100, 2)

## [1] 57.47 20.29 9.13 4.55 3.41 2.25 1.51 0.97 0.21 0.16 0.04

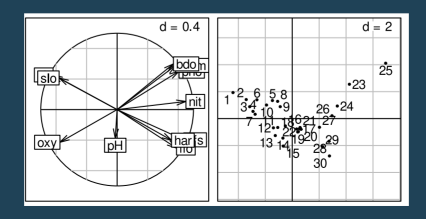
screeplot(pca1, main = "PCA Eigenvalues")
```



Graphical representation of PCA results

- Plot the variable scores with a correlation circle
- Plot the site scores on a separate plot
- Interpret

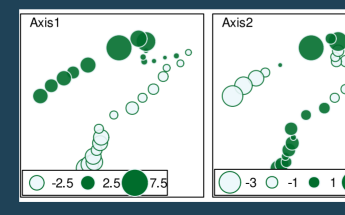
```
sc1 <- s.corcircle(pca1$co, plot = FALSE)
sl1 <- s.label(pca1$li, plabels.optim = TRUE, ppoints.cex = 0.5,
    plot = FALSE)
ADEgS(list(sc1, sl1))</pre>
```



PCA scores on the geographical map

- Draw maps of PCA scores on the first two axes
- Interpret the maps to describe the environmental structure of the river

```
s.value(xy, pca1$li[, 1:2], pgrid.draw = FALSE, porigin.draw = FALSE,
  method = "size", symbol = "circle", col = mypal(2),
  ppoints.cex = 1)
```



A look to variables

- Compute the correlations between original variables and PCA axes
- In which element of PCA outputs are stored these values

cor(pca1\$l1, doubs\$env)

```
dfs
                         alt
                                    slo
                                                flo
                                                                       har
##
                                                             Hq
        0.8733503 -0.8390535 -0.7602047 0.7777617 -0.02457895
                                                                 0.7122973 0.8
  RS2 -0.3963445
                   0.4537709 0.4145874 -0.5037445 -0.37560694 -0.4011204 0.5
##
             nit
                                            bdo
                       amm
                                  OXV
  RS1 0.9015491 0.7671086 -0.7489949 0.7375260
## RS2 0.1066847 0.5785809 -0.4280275 0.5985336
```

head(pca1\$co)

```
## Comp1 Comp2
## dfs 0.87335029 -0.3963445
## alt -0.83905354 0.4537709
## slo -0.76020468 0.4145874
## flo 0.77776175 -0.5037445
## pH -0.02457895 -0.3756069
## har 0.71229727 -0.4011204
```

A look to variables

The generic function score provides an optimal representation of the maximized criteria

score(pca1)

