# Multivariate analysis of genetic data: investigating spatial structures

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September 29, 2015

#### Abstract

This practical provides an introduction to the analysis of spatial genetic structures using R. Using an empirical dataset of microsatellites sampled from wild goats in the Bauges mountains (France), we illustrate univariate and multivariate tests of spatial structures, and then introduce the spatial Principal Component Analysis (sPCA) for uncovering spatial genetic patterns. For a more complete overview of spatial genetics methods, see the basics and sPCA tutorials for *adegenet*, which you can open using adegenetTutorial("basics") and adegenetTutorial("spca") (with an internet connection) or from the adegenet website.

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The chamois (*Rupicapra rupicapra*) is a conserved species in France. The Bauges mountains is a protected area in which the species has been recently studied. One of the most important questions for conservation purposes relates to whether individuals from this area form a single reproductive unit, or whether they are structured into sub-groups, and if so, what causes are likely to induce this structuring.

While field observations are very scarce and do not allow to answer this question, genetic data can be used to tackle the issue, as departure from panmixia should result in genetic structuring. The dataset *rupica* contains 335 georeferenced genotypes of Chamois from the Bauges mountains for 9 microsatellite markers, which we propose to analyse in this exercise.

#### 1 An overview of the data

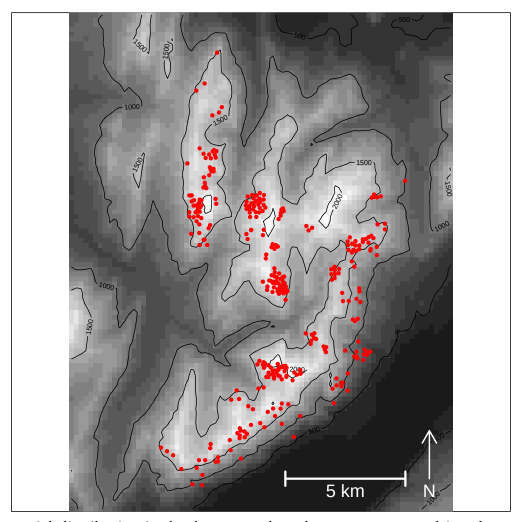
We first load some required packages and the data:

```
library(spdep)
library(adehabitat)
```

```
library(ade4)
library(adegenet)
##
##
      /// adegenet 2.0.1 is loaded /////////
##
##
      > overview:
                   '?adegenet'
      > tutorials/doc/questions:
                                   'adegenetWeb()'
##
      > bug reports/feature resquests: adegenetIssues()
##
data(rupica)
rupica
## /// GENIND OBJECT ///////
##
##
    // 335 individuals; 9 loci; 55 alleles; size: 217.2 Kb
##
    // Basic content
##
##
      Otab: 335 x 55 matrix of allele counts
      @loc.n.all: number of alleles per locus (range: 4-10)
##
      @loc.fac: locus factor for the 55 columns of @tab
##
      @all.names: list of allele names for each locus
##
      Oploidy: ploidy of each individual (range: 2-2)
##
      @type: codom
##
      @call: NULL
##
##
##
    // Optional content
      Oother: a list containing: xy mnt showBauges
##
```

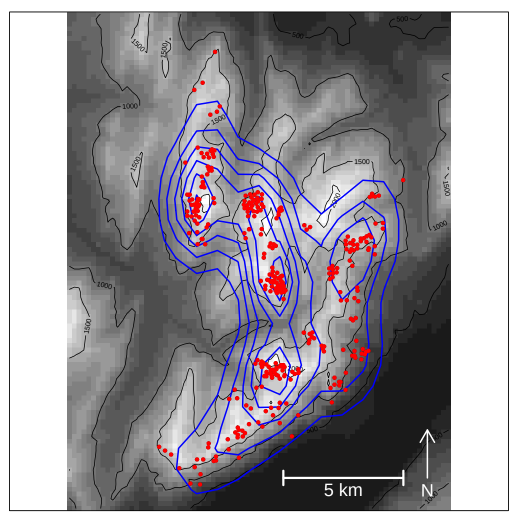
rupica is a typical genind object, which is the class of objects storing genotypes (as opposed to population data) in *adegenet*. rupica also contains topographic information about the sampled area, which can be displayed by calling rupica\$other\$showBauges. For instance, the spatial distribution of the sampling can be displayed as follows:

```
other(rupica)$showBauges()
points(other(rupica)$xy, col="red",pch=20)
```



This spatial distribution is clearly not random, but seems arranged into loose clusters. However, superimposed samples can bias our visual assessment of the spatial clustering. Use a two-dimensional kernel density estimation (function s.kde2d) to overcome this possible issue.

```
other(rupica)$showBauges()
s.kde2d(other(rupica)$xy,add.plot=TRUE)
points(other(rupica)$xy, col="red",pch=20)
```



Is geographical clustering strong enough to assign safely each individual to a group? Accordingly, shall we analyse these data at individual or group level?

## 2 Summarising the genetic diversity

As a prior clustering of genotypes is not known, we cannot employ usual  $F_{ST}$ -based approaches to detect genetic structuring. However, genetic structure could still result in a deficit of heterozygosity. Use the summary of genind objects to compare expected and observed heterozygosity:

```
rupica.smry <- summary(rupica)

##

## # Total number of genotypes: 335

##

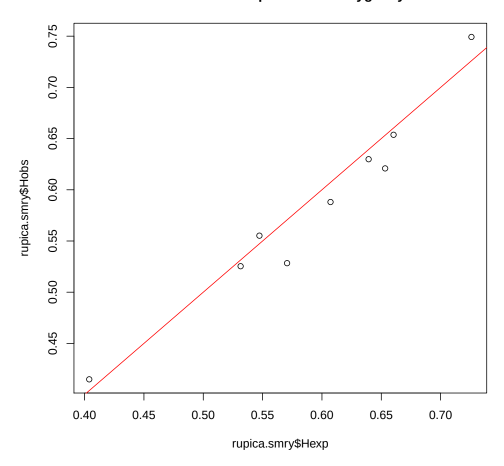
## # Population sample sizes:

## P1

## 335</pre>
```

```
##
## # Number of alleles per locus:
## Bm1818 Bm203 Eth225
                            Hel1 Bm4505 Bmc1009 Ilst030 Maf70
                                                                     Sr
                       7
                               6
                                     5
                                              5
##
               10
                                                                      5
##
## # Number of alleles per population:
## P1
## 55
##
## # Percentage of missing data:
## [1] 0
##
## # Observed heterozygosity:
              Bm203 Eth225
     Bm1818
                                    Hel1 Bm4505
                                                    Bmc1009
## 0.5880597 0.6208955 0.5253731 0.7492537 0.6537313 0.5283582 0.6298507
##
      Maf70
                   Sr
## 0.5552239 0.4149254
##
## # Expected heterozygosity:
               Bm203
    Bm1818
                        Eth225
                                    Hel1
                                            Bm4505
                                                    Bmc1009
                                                              Ilst030
## 0.6072158 0.6532517 0.5314591 0.7259626 0.6603405 0.5706082 0.6393235
##
      Maf70
                   Sr
## 0.5473112 0.4039120
plot(rupica.smry$Hexp, rupica.smry$Hobs, main="Observed vs expected heterozygosity")
abline(0,1,col="red")
```

#### Observed vs expected heterozygosity



The red line indicate identity between both quantities. What can we say about heterozygosity in this population? How can this be tested? The result below can be reproduced using a standard testing procedure:

```
t.test(rupica.smry$Hexp, rupica.smry$Hobs,paired=TRUE,var.equal=TRUE)

##

## Paired t-test

##

## data: rupica.smry$Hexp and rupica.smry$Hobs

## t = 1.1761, df = 8, p-value = 0.2734

## alternative hypothesis: true difference in means is not equal to 0

## 95 percent confidence interval:

## -0.007869215 0.024249885

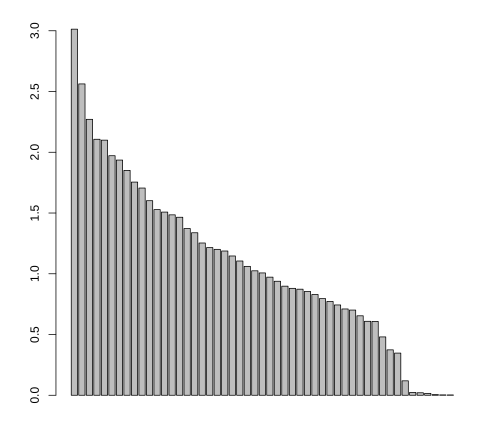
## sample estimates:

## mean of the differences

## 0.008190335
```

We can seek a global picture of the genetic diversity among genotypes using a Principal Component Analysis (PCA, [4, 2], dudi.pca in ade4 package). The analysis is performed on a table of allele frequencies, obtained by tab. The function dudi.pca displays a barplot of eigenvalues and asks for a number of retained principal components:

```
rupica.X <- tab(rupica, freq=TRUE, NA.method="mean")
rupica.pca1 <- dudi.pca(rupica.X, cent=TRUE, scannf=FALSE, nf=2)
barplot(rupica.pca1$eig)</pre>
```



The output produced by dudi.pca is a dudi object. A dudi object contains various information; in the case of PCA, principal axes (loadings), principal components (synthetic variable), and eigenvalues are respectively stored in \$c1, \$li, and \$eig slots. Here is the content of the PCA:

```
rupica.pca1
## Duality diagramm
## class: pca dudi
```

```
## $call: dudi.pca(df = rupica.X, center = TRUE, scannf = FALSE, nf = 2)
##
## $nf: 2 axis-components saved
## $rank: 51
## eigen values: 3.013 2.563 2.271 2.107 2.1 ...
     vector length mode
                            content
## 1 $cw
            55
                   numeric column weights
## 2 $1w
                   numeric row weights
            335
                   numeric eigen values
## 3 $eig
            51
##
##
     data.frame nrow ncol content
## 1 $tab
                335
                     55
                           modified array
## 2 $li
                335
                      2
                           row coordinates
## 3 $11
                335
                     2
                          row normed scores
## 4 $co
                      2
                           column coordinates
                55
## 5 $c1
                      2
                55
                           column normed scores
## other elements: cent norm
```

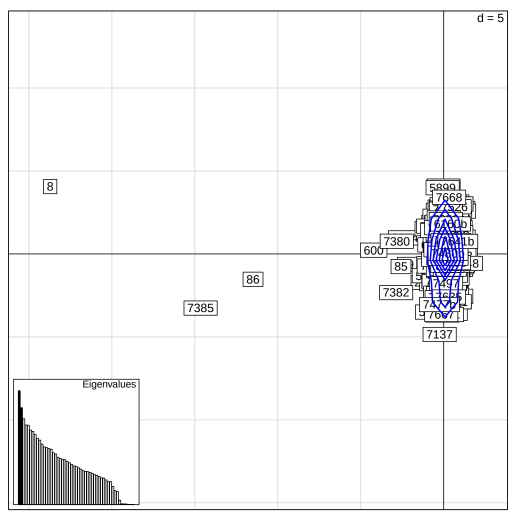
In general, eigenvalues represent the amount of genetic diversity — as measured by the multivariate method being used — represented by each principal component (PC). Verify that here, each eigenvalue is the variance of the corresponding PC.

```
head(rupica.pca1$eig)
## [1] 3.013193 2.562510 2.271093 2.107049 2.099738 1.971703
apply(rupica.pca1$li,2,var)*334/335
## Axis1 Axis2
## 3.013193 2.562510
```

An abrupt decrease in eigenvalues is likely to indicate the boundary between true patterns and non-interpretable structures. In this case, how many PCs would you interprete?

Use s.label to display to two first components of the analysis. Then, use a kernel density (s.kde2d) for a better assessment of the distribution of the genotypes onto the principal axes:

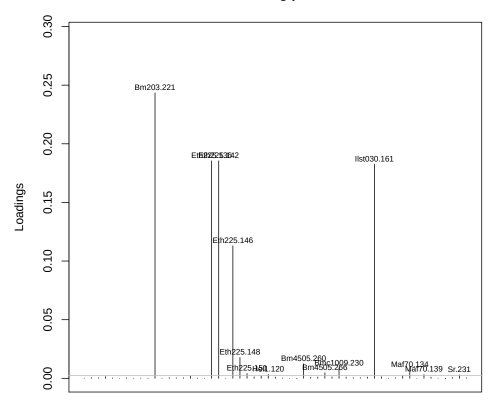
```
s.label(rupica.pca1$li)
s.kde2d(rupica.pca1$li, add.p=TRUE, cpoint=0)
add.scatter.eig(rupica.pca1$eig,2,1,2)
```



What can we say about the genetic diversity among these genotypes as inferred by PCA? The function loadingplot allows to visualize the contribution of each allele, expressed as squared loadings, for a given principal component. Using this function, reproduce this figure:

loadingplot(rupica.pca1\$c1^2)

#### **Loading plot**



#### Variables

What do we observe? We can get back to the genotypes for the concerned markers (e.g., Bm203) to check whether the highlighted genotypes are uncommon. tab extracts the table of allele frequencies from a genind object:

```
X <- tab(rupica)
class(X)

## [1] "matrix"

dim(X)

## [1] 335 55

bm203.221 <- X[,"Bm203.221"]
table(bm203.221)

## bm203.221

## 0 1
## 331 4</pre>
```

Only 4 genotypes possess one copy of the allele 221 of marker bm203 (the second result corresponds to a replaced missing data). Which individuals are they?

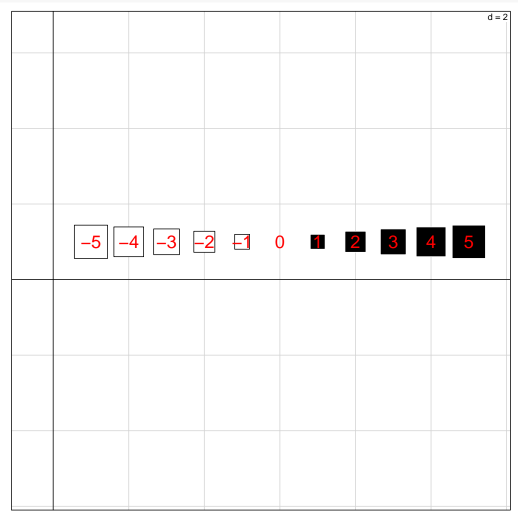
```
rownames(X)[bm203.221>0]
## [1] "8" "86" "600" "7385"
```

Conclusion?

## 3 Mapping and testing PCA results

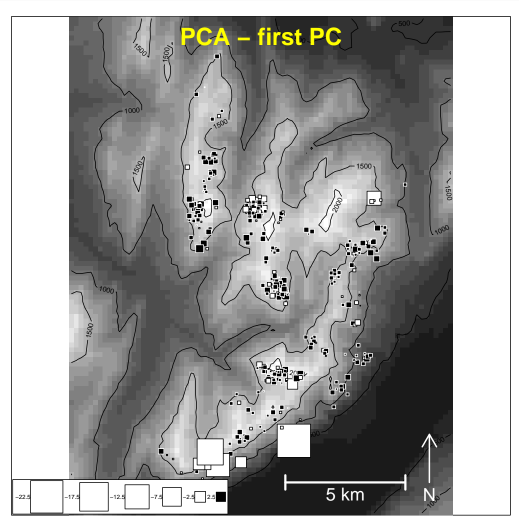
A frequent practice in spatial genetics is mapping the first principal components (PCs) onto the geographic space. The function **s.value** is well-suited to do so, using black and white squares of variable size for positive and negative values. To give a legend for this type of representation:

```
s.value(cbind(1:11,rep(1,11)), -5:5, cleg=0)
text(1:11,rep(1,11), -5:5, col="red",cex=1.5)
```

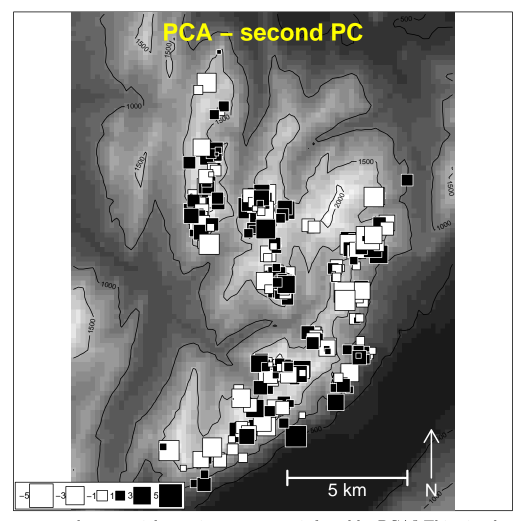


Apply this graphical representation to the first two PCs of the PCA:

```
showBauges <- other(rupica)$showBauges
showBauges()
s.value(other(rupica)$xy, rupica.pca1$li[,1], add.p=TRUE, cleg=0.5)
title("PCA - first PC",col.main="yellow" ,line=-2, cex.main=2)</pre>
```



```
showBauges()
s.value(other(rupica)$xy, rupica.pca1$li[,2], add.p=TRUE, csize=0.7)
title("PCA - second PC",col.main="yellow",line=-2, cex.main=2)
```



What can we say about spatial genetic structure as inferred by PCA? This visual assessment can be complemented by testing the spatial autocorrelation in the first PCs of PCA. This can be achieved using Moran's I test. Use the function moran.mc in the package spdep to perform these tests. You will need first to define the spatial connectivity between the sampled individuals. For these data, spatial connectivity is best defined as the overlap between home ranges of individuals. Home ranges will be modelled as disks with a radius of 1150m. Use chooseCN to create a connection network based on distance range ("neighbourhood by distance"). What threshold distance do you choose for individuals to be considered as neighbours?

The connection network should ressemble this:

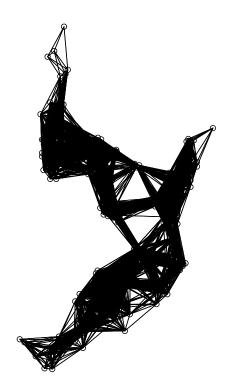
```
rupica.graph

## Characteristics of weights list object:
## Neighbour list object:
```

```
## Number of regions: 335
## Number of nonzero links: 18018
## Percentage nonzero weights: 16.05525
## Average number of links: 53.78507
##
## Weights style: W
## Weights constants summary:
## n nn S0 S1 S2
## W 335 112225 335 15.04311 1352.07

plot(rupica.graph, other(rupica)$xy)
title("rupica.graph")
```

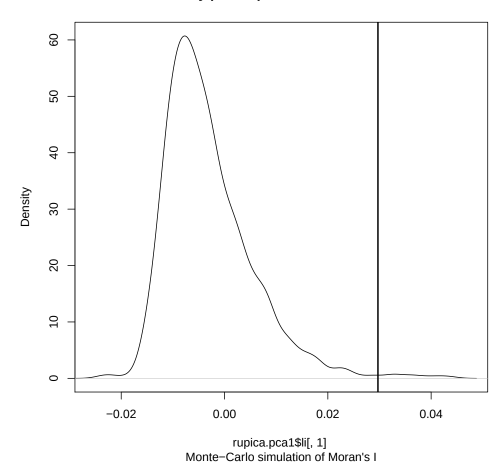
#### rupica.graph



Perform Moran's test for the first two PCs, and plot the results. The first test should be significant:

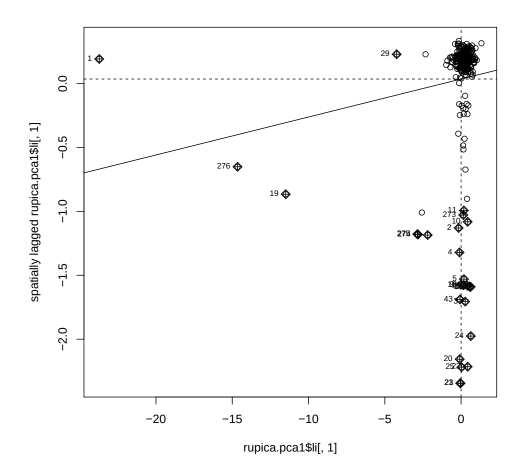
```
pc1.mctest <- moran.mc(rupica.pca1$li[,1], rupica.graph, 999)
plot(pc1.mctest)</pre>
```

#### Density plot of permutation outcomes



Compare this result to the mapping of the first PC of PCA. What is wrong? When a test gives unexpected results, it is worth looking into the data in more details. Moran's plot (moran.plot) plots the tested variable against its lagged vector. Use it on the first PC:

moran.plot(rupica.pca1\$li[,1], rupica.graph)

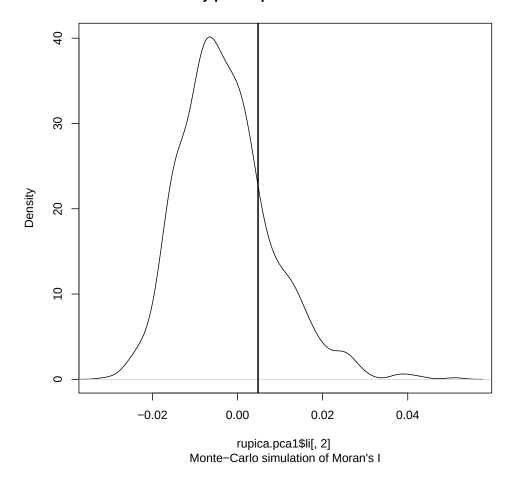


Actual positive autocorrelation corresponds to a positive correlation between a variable and its lag vector. Is it the case here? How can we explain that Moran's test was significant? Repeat these analyses for the second PC. What are your conclusions?

```
pc2.mctest <- moran.mc(rupica.pca1$li[,2], rupica.graph, 999)
pc2.mctest

##
## Monte-Carlo simulation of Moran's I
##
## data: rupica.pca1$li[, 2]
## weights: rupica.graph
## number of simulations + 1: 1000
##
## statistic = 0.0048162, observed rank = 797, p-value = 0.203
## alternative hypothesis: greater</pre>
plot(pc2.mctest)
```

#### **Density plot of permutation outcomes**

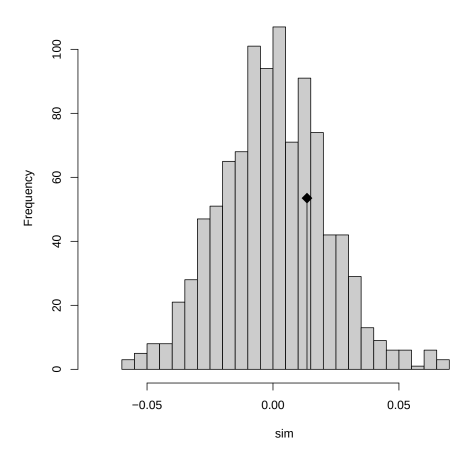


## 4 Multivariate tests of spatial structure

So far, we have only tested the existence of spatial structures in the first two principal components of a PCA of the data. Therefore, these tests only describe one fragment of the data, and do not encompass the whole diversity in the data. As a complement, we can use Mantel test (mantel.randtest) to test spatial structures in the whole data, by assessing the correlation between genetic distances and geographic distances. Pairwise Euclidean distances are computed using dist. Perform Mantel test, using the scaled genetic data you used before in PCA, and the geographic coordinates.

```
mtest <- mantel.randtest(dist(rupica.X), dist(other(rupica)$xy))
plot(mtest, nclass=30)</pre>
```

#### Histogram of sim



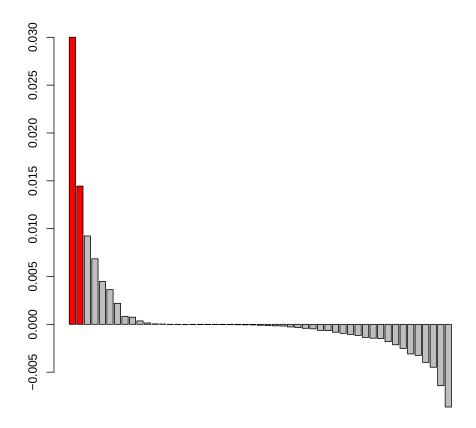
What is your conclusion? Shall we be looking for spatial structures? If so, how can we explain that PCA did not reveal them?

### 5 spatial Principal Component Analysis

The spatial Principal Component Analysis (sPCA, function spca [3]) has been especially developed to investigate hidden or non-obvious spatial genetic patterns. Like Moran's I test, sPCA first requires the spatial proximities between genotypes to be modeled. You will reuse the connection network defined previously using chooseCN, and pass it as the 'cn' argument of the function spca.

Read the documentation of spca, and apply the function to the dataset rupica. The function will display a barplot of eigenvalues:

```
rupica.spca1 <- spca(rupica, cn=rupica.graph,scannf=FALSE, nfposi=2,nfnega=0)
barplot(rupica.spca1$eig, col=rep(c("red","grey"), c(2,1000)) )</pre>
```



This figure illustrates the fundamental difference between PCA and sPCA. Like  $\mathtt{dudi.pca}$ , spca displays a barplot of eigenvalues, but unlike in PCA, eigenvalues of sPCA can also be negative. This is because the criterion optimized by the analysis can have positive and negative values, corresponding respectively to positive and negative autocorrelation. Positive spatial autocorrelation correspond to greater genetic similarity between geographically closer individuals. Conversely, negative spatial autocorrelation corresponds to greater dissimilarity between neighbours. The spatial autocorrelation of a variable is measured by Moran's I, and interpreted as follows:

- $I_0 = -1/(n-1)$ : no spatial autocorrelation (x is randomly distributed across space)
- $I > I_0$ : positive spatial autocorrelation
- $I < I_0$ : negative spatial autocorrelation

Principal components of PCA ensure that  $(\phi \text{ referring to one PC}) \ var(\phi)$  is maximum. By contrast, sPCA provides PC which decompose the quantity  $var(\phi)I(\phi)$ . In other words, PCA focuses on variability only, while sPCA is a compromise between variability  $(var(\phi))$  and spatial structure  $(I(\phi))$ .

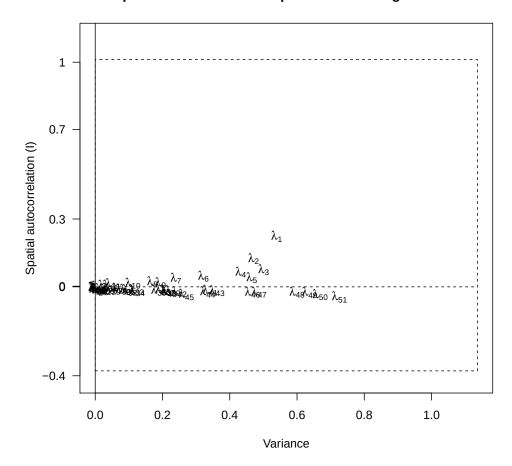
In this case, only the principal components associated with the two first positive eigenvalues (in red) shall be retained. The printing of spca objects is more explicit than dudi objects, but named with the same conventions:

```
rupica.spca1
   ##
   # spatial Principal Component Analysis #
   ##
## class: spca
## $call: spca(obj = rupica, cn = rupica.graph, scannf = FALSE, nfposi = 2,
##
      nfnega = 0)
## $nfposi: 2 axis-components saved
## $nfnega: 0 axis-components saved
## Positive eigenvalues: 0.03001 0.01445 0.00924 0.006851 0.004485 ...
## Negative eigenvalues: -0.008643 -0.006407 -0.004488 -0.003977 -0.003248 ...
##
    vector length mode
                         content
## 1 $eig
           51
                 numeric eigenvalues
##
##
    data.frame nrow ncol
## 1 $c1
               55
## 2 $li
               335
                   2.
## 3 $1s
                   2
               335
## 4 $as
##
    content
## 1 principal axes: scaled vectors of alleles loadings
## 2 principal components: coordinates of entities ('scores')
## 3 lag vector of principal components
## 4 pca axes onto spca axes
## $xy: matrix of spatial coordinates
## $lw: a list of spatial weights (class 'listw')
##
## other elements: NULL
```

Unlike usual multivariate analyses, eigenvalues of sPCA are composite: they measure both the genetic diversity (variance) and the spatial structure (spatial autocorrelation measured by Moran's I). This decomposition can also be used to choose which principal component to interprete. The function **screeplot** allows to display this information graphically:

```
screeplot(rupica.spca1)
```

#### Spatial and variance components of the eigenvalues



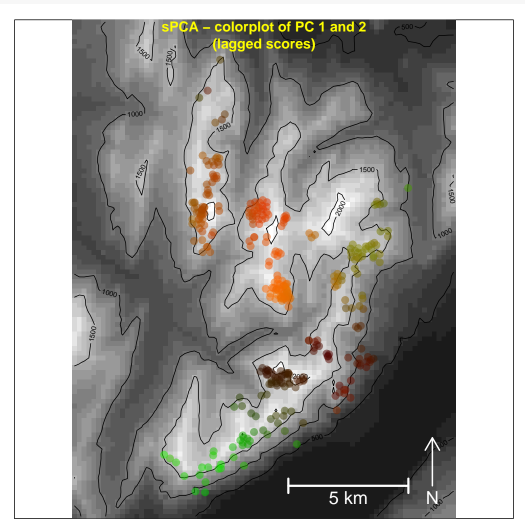
While  $\lambda_1$  indicates with no doubt a structure, the second eigenvalue,  $\lambda_2$  is less clearly distinct from the successive values. Thus, we shall keep in mind this uncertainty when interpreting the second principal component of the analysis.

Try visualising the sPCA results as you did before with the PCA results. To clarify the possible spatial patterns, you can map the lagged PC (\$ls) instead of the PC (\$li), which are a 'denoisified' version of the PCs.

First, map the first principal component of sPCA. How would you interprete this result? How does it compare to the first PC of PCA? What inference can we make about the way the landscape influences gene flow in this population of Chamois?

Do the same with the second PC of sPCA. Some field observations suggest that this pattern is not artefactual. How would you interprete this second structure?

To finish, you can try representing both structures at the same time using the color coding introduced by [1] (?colorplot). The final figure should ressemble this (although colors may change from one computer to another):



## 6 To go further

More spatial genetics methods and are presented in the  $basics\ tutorial$  as well as in the tutorial dedicated to sPCA, which you can access from the adegenet website:

```
http://adegenet.r-forge.r-project.org/
```

or by typing:

```
adegenetTutorial("basics")
adegenetTutorial("spca")
```

## References

- [1] L. L. Cavalli-Sforza, P. Menozzi, and A. Piazza. Demic expansions and human evolution. *Science*, 259:639–646, 1993.
- [2] I. T. Joliffe. *Principal Component Analysis*. Springer Series in Statistics. Springer, second edition, 2004.
- [3] T. Jombart, S. Devillard, A.-B. Dufour, and D. Pontier. Revealing cryptic spatial patterns in genetic variability by a new multivariate method. *Heredity*, 101:92–103, 2008.
- [4] K. Pearson. On lines and planes of closest fit to systems of points in space. *Philosophical Magazine*, 2:559–572, 1901.