A tutorial for the spatial Analysis of Principal Components (sPCA) using *adegenet* 1.3-4

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Abstract

This vignette provides a tutorial for the spatial analysis of principal components (sPCA, [1]) using the *adegenet* package [2] for the R software [3]. sPCA is first illustrated using a simple simulated dataset, and then using empirical data of Chamois (*Rupicapra rupicapra*) from the Bauges mountains (France). In particular, we illustrate how sPCA complements classical PCA by being more powerful for retrieving non-trivial spatial genetic patterns.

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1 Introduction

This tutorial goes through the spatial Principal Component Analysis (sPCA, [1]), a multivariate method devoted to the identification of spatial genetic patterns. The purpose of this tutorial is to provide guidelines for the application of sPCA as well as to illustrate its usefulness for the investigation of spatial genetic patterns. After briefly going through the rationale of the method, we introduce the different tools implemented for sPCA in adegenet. This technical overview is then followed by the analysis of an empirical dataset which illustrates the advantage of sPCA over classical PCA for investigating spatial patterns.

1.1 Rationale of sPCA

Mathematical notations used in this tutorial are identical to the original publication [1]. The sPCA analyses a matrix of relative allele frequencies \mathbf{X} which contains genotypes or populations (later refered to as 'entities') in rows and alleles in columns. Spatial information is stored inside a spatial weighting matrix \mathbf{L} which contains positive terms corresponding to some measurement (often binary) of spatial proximity among entities. Most often, these terms can be derived from a connection network built upon a given algorithm (for instance, pp.572-576 in [4]). This matrix is row-standardized (*i.e.*, each of its rows sums to one), and all its diagonal terms are zero. \mathbf{L} can be used to compute the spatial autocorrelation of a given centred variable \mathbf{x} (i.e., with mean zero) with n observations ($\mathbf{x} \in \mathbb{R}^n$) using Moran's I [5, 6, 7]:

$$I(\mathbf{x}) = \frac{\mathbf{x}^T \mathbf{L} \mathbf{x}}{\mathbf{x}^T \mathbf{x}} \tag{1}$$

In the case of genetic data, \mathbf{x} contains frequencies of an allele. Moran's I can be used to measure spatial structure in the values of \mathbf{x} : it is highly positive when values of \mathbf{x} observed at neighbouring sites tend to be similar (positive spatial autocorrelation, referred to as global structures), while it is strongly negative when values of \mathbf{x} observed at neighbouring sites tend to be dissimilar (negative spatial autocorrelation, referred to as local structures).

However, since it is standardized by the variance of \mathbf{x} , Moran's index measures only spatial structures and not genetic variability. The sPCA defines the following function to measure both spatial structure and variability in \mathbf{x} :

$$C(\mathbf{x}) = \text{var}(\mathbf{x})I(\mathbf{x}) = \frac{1}{n}\mathbf{x}^T\mathbf{L}\mathbf{x}$$
 (2)

 $C(\mathbf{x})$ is highly positive when \mathbf{x} has a large variance and exhibits a global structure; conversely, it is largely negative when \mathbf{x} has a high variance and displays a local structure. This function is the criterion used in sPCA, which finds linear combinations of the alleles of \mathbf{X} (denoted $\mathbf{X}\mathbf{v}$) decomposing C from its maximum to its minimum value. Because $C(\mathbf{X}\mathbf{v})$ is a product of variance and autocorrelation, it is important, when interpreting the results, to detail

both components and to compare their value with their range of variation (maximum attainable variance, as well as maximum and minimum I are known analytically). A structure with a low spatial autocorrelation can barely be interpreted as a spatial pattern; similarly, a structure with a low variance would likely not reflect any genetic structure. We will later see how these information can be retrieved from spca results.

1.2 The spca function

The simulated dataset used to illustrate this section has been analyzed in [1], and corresponds to Figure 2A of the article. In *adegenet*, the matrix of alleles frequencies previously denoted **X** exactly corresponds to the **@tab** slot of **genind** or **genpop** objects:

```
> library(adegenet)
> library(adehabitat)
> data(spcaIllus)
 obj <- spcaIllus$dat2A
   #####################
   genotypes of individuals -
S4 class: genind
@call: old2new(object = obj)
@tab: 80 x 192 matrix of genotypes
@ind.names: vector of 80 individual names
@loc.names: vector of 20 locus names
Cloc.nall: number of alleles per locus
Cloc.fac: locus factor for the 192 columns of Ctab
Call.names: list of 20 components yielding allele names for each locus Oploidy: 2
@ploidy:
Otype:
       codom
Optionnal contents:
@pop: factor giving the population of each individual
@pop.names: factor giving the population of each individual
@other: a list containing: xy
> head(truenames(obj[loc="L01"])$tab)
     0035
                                0.5
0352
                                 0.5
                                        0.0
                                                     0.0
                                                            0.5
0423
                0
                    0.0
                             0
                                                 0
                    0.0
                             Ŏ
0289
0053
                                  0.5
```

The object obj is a genind object; note that here, we only displayed the table for the first locus (loc="L01").

The function performing the sPCA is spca; it accepts a bunch of arguments, but only the first two are mandatory to perform the analysis (see ?spca for further information):

> args(spca)

```
function (obj, xy = NULL, cn = NULL, matWeight = NULL, scale = FALSE,
    scale.method = c("sigma", "binom"), scannf = TRUE, nfposi = 1,
    nfnega = 1, type = NULL, ask = TRUE, plot.nb = TRUE, edit.nb = FALSE,
    truenames = TRUE, d1 = NULL, d2 = NULL, k = NULL, a = NULL,
    dmin = NULL)
```

The argument obj is a genind/genpop object. By definition in sPCA, the studied entities are georeferenced. The spatial information can be provided to the function spca in several ways, the first being through the xy argument, which is a matrix of spatial coordinates with 'x' and 'y' coordinates in columns. Alternatively, these coordinates can be stored inside the genind/genpop object, preferably as @other\$xy, in which case the spca function will detect and use this information, and not request an xy argument. Note that obj already contains spatial coordinates at the appropriate place. Hence, we can use the following command to run the sPCA (ask and scannf are set to FALSE to avoid interactivity):

```
> mySpca <- spca(obj, ask=FALSE, type=1, scannf=FALSE)</pre>
```

Note, however, that spatial coordinates are not directly used in sPCA: the spatial information is included in the analysis by the spatial weighting matrix L derived from a connection network (eq. 1 and 2). Technically, the spca function can incorporate spatial weightings as a matrix (argument matWeight), as a connection network with the classes nb or listw (argument cn), both implemented in the spdep package. The function chooseCN is a wrapper for different functions scattered across several packages implementing a variety of connection networks. If only spatial coordinates are provided to spca, chooseCN is called to construct an appropriate graph. See ?chooseCN for more information. Note that many of the spca arguments are in fact arguments for chooseCN: type, ask, plot.nb, edit.nb, d1, d2, k, a, and dmin. For instance, the command:

```
> mySpca <- spca(obj,type=1,ask=FALSE,scannf=FALSE)</pre>
```

performs a sPCA using the Delaunay triangulation as connection network (type=1, see ?chooseCN), while the command:

```
> mySpca <- spca(obj,type=5,d1=0,d2=2,scannf=FALSE)</pre>
```

computes a sPCA using a connection network which defines neighbouring entities based on pairwise geographic distances (type=5), considering as neighbours two entities whose distance between 0 (d1=0) and 2 (d2=2).

Another possibility is of course to provide directly a connection network (nb object) or a list of spatial weights (listw object) to the spca function; this can be done via the cn argument. For instance:

```
> myCn <- chooseCN(obj$other$xy, type=6, k=10, plot=FALSE)
> myCn

Neighbour list object:
Number of regions: 80
Number of nonzero links: 932
Percentage nonzero weights: 14.5625
Average number of links: 11.65

> class(myCn)

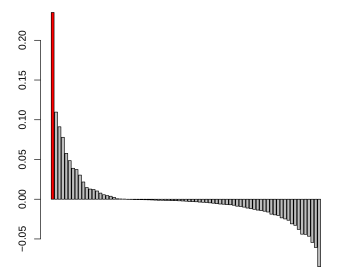
[1] "nb"
> mySpca2 <- spca(obj,cn=myCn,scannf=FALSE)</pre>
```

produces a sPCA using ${\tt myCn}\ (k=10\ {\tt nearest\ neighbours})$ as a connection network.

When used interactively (scannf=TRUE), spca displays a barplot of eigenvalues and asks the user for a number of positive axes ('first number of axes') and negative axes ('second number of axes') to be retained. For the object mySpca, this barplot would be (here we indicate in red the retained eigenvalue):

> barplot(mySpca\$eig,main="Eigenvalues of sPCA", col=rep(c("red","grey"),c(1,100)))

Eigenvalues of sPCA



Positive eigenvalues (on the left) correspond to global structures, while negative eigenvalues (on the right) indicate local patterns. Actual structures should result in more extreme (positive or negative) eigenvalues; for instance, the object mySpca likely contains one single global structure, and no local structure. If one does not want to choose the number of retained axes interactively, the arguments nfposi (number of retained factors with positive eigenvalues) and nfnega (number of retained factors with negative eigenvalues) can be used. Once this information has been provided to spca, the analysis is computed and stored inside an object with the class spca.

1.3 Contents of a spca object

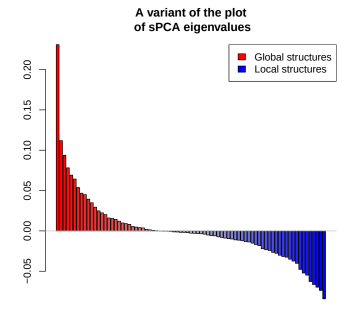
Let us consider a spca object resulting from the analysis of the object obj, using a Delaunay triangulation (type=1) as connection network:

```
> mySpca <- spca(obj,type=1,scannf=FALSE,plot.nb=FALSE,nfposi=1,nfnega=0)</pre>
> class(mySpca)
[1] "spca"
> mySpca
        class: spca
$call: spca(obj = obj, scannf = FALSE, nfposi = 1, nfnega = 0, type = 1,
   plot.nb = FALSE)
$nfposi: 1 axis-components saved
$nfnega: 0 axis-components saved
Positive eigenvalues: 0.2309 0.1118 0.09379 0.07817 0.06911 ...
Negative eigenvalues: -0.08421 -0.07376 -0.06978 -0.06648 -0.06279 ...
  vector length mode
                      content
               numeric eigenvalues
1 $eig 79
  data.frame nrow ncol content
                      principal axes: scaled vectors of alleles loadings
1 $c1
            192 1
                1
1
                      principal components: coordinates of entities ('scores')
2 $1i
            80
3 $1s
            80
                      lag vector of principal components
4 $as
            2
                 1
                      pca axes onto spca axes
$xy: matrix of spatial coordinates
$lw: a list of spatial weights (class 'listw')
other elements: NULL
```

An spca object is a list containing all required information about a performed sPCA. Details about the different components of such a list can be found in the spca documentation (?spca). The purpose of this section is to explicit how the elements described in [1] are stored inside a spca object.

First, eigenvalues of the analysis are stored inside the **\$eig** component as a numeric vector stored in decreasing order:

```
> head(mySpca$eig)
```



The axes of the analysis, denoted ${\bf v}$ in eq. (4) [1] are stored as columns inside the \$c1 component. Each column contains loadings for all the alleles:

> head(mySpca\$c1)

```
Axis 1
L01.1 1.268838e-02
L01.2 2.220446e-16
L01.3 -1.119979e-01
L01.4 -4.440892e-16
L01.5 -2.766095e-02
L01.6 -4.477031e-02
```

```
> tail(mySpca$c1)

Axis 1
L20.3 0.28715850
L20.4 0.01485180
L20.5 -0.01500353
L20.6 0.01659481
L20.7 -0.14260743
L20.8 -0.15388988

> dim(mySpca$c1)
```

[1] 192 1

The entity scores, denoted $\psi = \mathbf{X}\mathbf{v}$ in the article, are stored in columns in the \$1i component:

> head(mySpca\$li)

```
Axis 1
0035 -0.4367748
0352 -0.8052723
0423 -0.4337114
0289 0.1434650
0487 -0.4802931
0053 -0.5421831
```

> tail(mySpca\$li)

```
Axis 1
1074 -0.06178196
1187 -0.08144162
1260 0.41491795
1038 0.25643986
1434 0.35618737
1218 0.21433977
```

> dim(mySpca\$li)

[1] 80 1

The lag vectors of the scores can be used to better perceive global structures. Lag vectors are stored in the \$1s component:

> head(mySpca\$ls)

```
Axis 1
0035 -0.7076732
0352 -0.6321654
0423 -0.4822952
0289 0.3947791
0487 -0.2803381
0053 -0.4848376
```

> tail(mySpca\$ls)

```
Axis 1
1074 0.4930238
1187 -0.8384871
1260 0.6887072
1038 0.3665794
1434 0.3109197
1218 0.3329688

> dim(mySpca$ls)
```

Lastly, we can compare the axes of an classical PCA (denoted \mathbf{u} in the paper) to the axes of the sPCA (\mathbf{v}). This is achieved by projecting \mathbf{u} onto \mathbf{v} , but this projection is a particular one: because both \mathbf{u} and \mathbf{v} are centred to mean zero and scaled to unit variance, the value of the projection simply is the correlation between both axes. This information is stored inside the \$as component:

```
> mySpca$as
```

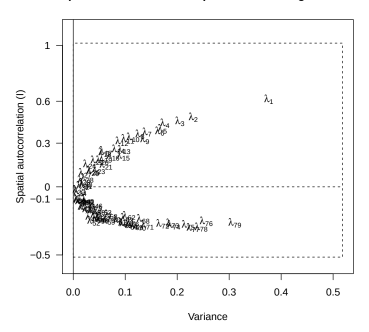
```
Axis 1
PCA Axis1 -0.7363595
PCA Axis2 0.3395674
```

1.4 Graphical display of spca results

The information contained inside a spca object can be displayed in several ways. While we have seen that a simple barplot of sPCA eigenvalues can give a first idea of the global and local structures to be retained, we have also seen that each eigenvalue can be decomposed into a *variance* and a *spatial autocorrelation* (Moran's I) component. This information is provided by the summary function, but it can also be represented graphically. The corresponding function is screeplot, and can be used on any spca object:

```
> screeplot(mySpca)
```

Spatial and variance components of the eigenvalues



The resulting figure represents eigenvalues of sPCA (denoted λ_i with $i=1,\ldots,r$, where λ_1 is the highest positive eigenvalue, and λ_r is the highest negative eigenvalue) according the their variance and Moran's I components. These eigenvalues are contained inside a rectangle indicated in dashed lines. The maximum attainable variance by a linear combination of alleles is the one from an ordinary PCA, indicated by the vertical dashed line on the right. The two horizontal dashed lines indicate the range of variation of Moran's I, given the spatial weighting matrix that was used. This figure is useful to assess whether a given score of entities contains relatively enough variability and spatial structuring to be interpreted. For instance, here, λ_1 clearly is the largest eigenvalue in terms of variance and of spatial autocorrelation, and can be well distinguished from all the other eigenvalues. Hence, only the first global structure, associated to λ_1 , should be interpreted.

The global and local tests proposed in [1] can be used to reinforce the decision of interpreting or not interpreting global and local structures. Each test can detect the presence of one kind of structure. We can apply them to the object obj, used in our sPCA:

```
> myGtest <- global.rtest(obj$tab,mySpca$lw,nperm=99)
> myGtest

Monte-Carlo test
Call: global.rtest(X = obj$tab, listw = mySpca$lw, nperm = 99)
```

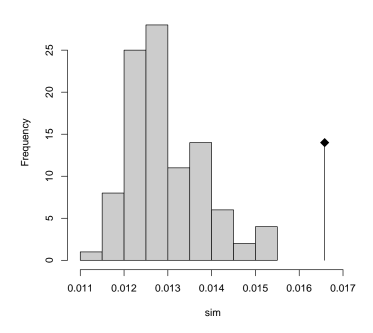
```
Observation: 0.01658103

Based on 99 replicates
Simulated p-value: 0.01
Alternative hypothesis: greater

Std.Obs Expectation Variance
4.181857e+00 1.295912e-02 7.501292e-07
```

> plot(myGtest)

Histogram of sim



The produced object is a randtest object (see ?randtest), which is the class of objects for Monte-Carlo tests in the ade4 package. As shown, such object can be plotted using a plot function: the resulting figure shows an histogram of permuted test statistics and indicates the observed statistics by a black dot and a segment. Here, the plot clearly shows that the oberved test statistic is larger than most simulated values, leading to a likely rejection of the null hypothesis of absence of spatial structure. Note that because 99 permutations were used, the p-value cannot be lower than 0.01. In practice, more permutations should be used (like 999 or 9999 for results intended to be published).

The same can be done with the local test, which here we do not expect to be significant:

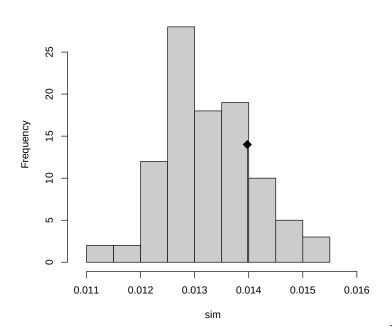
```
> myLtest <- local.rtest(obj$tab,mySpca$lw,nperm=99)
> myLtest
```

```
Monte-Carlo test
Call: local.rtest(X = obj$tab, listw = mySpca$lw, nperm = 99)
Observation: 0.01397349
Based on 99 replicates
Simulated p-value: 0.21
Alternative hypothesis: greater

Std.Obs Expectation Variance
8.943942e-01 1.323669e-02 6.786467e-07
```

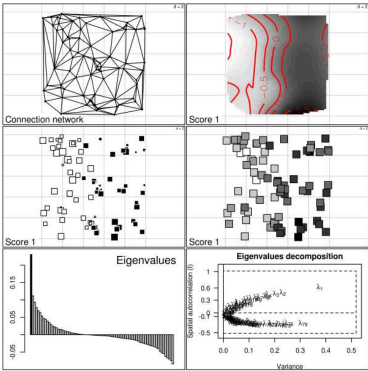
> plot(myLtest)

Histogram of sim



Once we have an idea of which structures shall be interpreted, we can try to visualize spatial genetic patterns. There are several ways to do so. The first, most simple approach is through the function plot (see ?plot.spca):

> plot(mySpca)



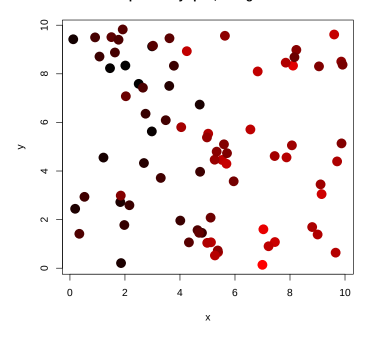
This figure displays various information, that we detail from the top to bottom and from left to right (also see ?plot.spca). The first plot shows the connection network that was used to define spatial weightings. second, third, and fourth plots are different representations of a score of entities in space, the first global score being the default (argument axis). In each, the values of scores (\$li[,axis] component of the spca object) are represented using black and white symbols (a variant being grey levels): white for negative values, and black for positive values. The second plot is a local interpolation of scores (function s.image in ade4), using grey levels, with contour lines. The closer the contour lines are from each other, the stepest the genetic differentiation is. The third plot uses different sizes of squares to represent different absolute values (s.value in ade4): large black squares are well differentiated from large white squares, but small squares are less differentiated. The fourth plot is a variant using grey levels (s.value in ade4, with 'greylevel' method). Here, all the three representations of the first global score show that genotypes are splitted in two genetical clusters, one in the west (or left) and one in the east (right). The last two plots of the plot.spca function are the two already seen displays of eigenvalues.

While the default plot function for spca objects provides a useful summary of the results, more flexible tools are needed e.g. to map the principal components onto the geographic space. This can be achieved using the

colorplot function. This function can summarize up to three scores at the same time by translating each score into a channel of color (red, green, and blue). The obtained values are used to compose a color using the RGB system. See ?colorplot for details about this function. The original idea of such representation is due to [8]. Despite the colorplot clearly is more powerful to represent more than one score on a single map, we can use it to represent the first global structure that was retained in mySpca:

> colorplot(mySpca,cex=3,main="colorplot of mySpca, first global score")

colorplot of mySpca, first global score

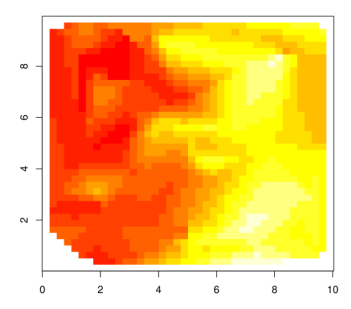


See examples in ?colorplot and ?spca for more examples of applications of colorplot to represent sPCA scores.

Another common practice is interpolating principal components to get maps of genetic clines. Note that it is crucial to perform this interpolation after the analysis, and not before, which would add artefactual structures to the data. Interpolation is easy to realize using interp from the akima package, and image, or filled.contour to display the results:

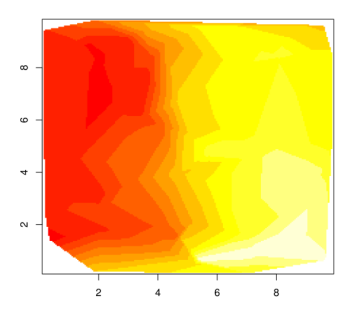
```
> library(akima)
> x <- other(obj)$xy[,1]
> y <- other(obj)$xy[,2]

> temp <- interp(x, y, mySpca$li[,1])
> image(temp)
```



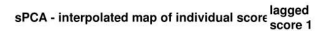
Note that for better clarity, we can use the lagged principal scores (\$1s) rather than the original scores (\$1i); we also achieve a better resolution using specific interpolated coordinates:

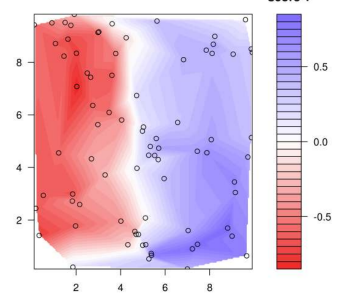
```
> interpX <- seq(min(x),max(x),le=200)
> interpY <- seq(min(y),max(y),le=200)
> temp <- interp(x, y, mySpca$ls[,1], xo=interpX, yo=interpY)
> image(temp)
```



Alternatively, filled.contour can be used for the display, and a customized color palette can be specified:

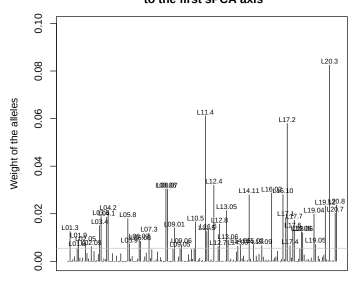
```
> myPal <- colorRampPalette(c("firebrick2", "white", "lightslateblue"))
> annot <- function(){
+         title("sPCA - interpolated map of individual scores")
+         points(x,y)
+ }
> filled.contour(temp, color.pal=myPal, nlev=50, key.title=title("lagged \nscore 1"), plot.title=annot())
```





Besides assessing spatial patterns, it is sometimes valuable to assess which alleles actually exhibit the structure of interest. In sPCA, the contribution of alleles to a specific structure is given by the corresponding squared loading. We can look for the alleles contributing most to e.g. the first axis of sPCA, using the function loadingplot (see ?loadingplot for a description of the arguments):

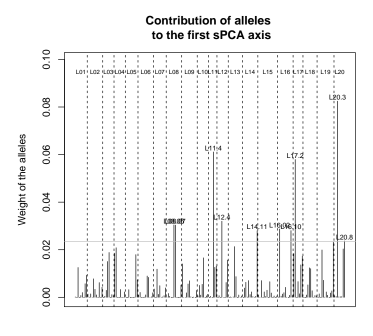
Contribution of alleles to the first sPCA axis



Alleles

See ?loadingplot for more information about this function, in particular for the definition of the threshold value above which alleles are annotated. Note that it is possible to also separate the alleles by markers, using the fac argument, to assess if all markers have comparable contributions to a given structure. In our case, we would only have to specify fac=obj@loc.fac; also note that loadingplot invisibly returns information about the alleles whose contribution is above the threshold. For instance, to identify the 5% of alleles with the greatest contributions to the first global structure in mySpca, we need:

```
 \begin{tabular}{ll} temp <- loading plot(myLoadings, threshold=quantile(myLoadings, 0.95), & xlab="Alleles", ylab="Weight of the alleles", & the alleles", & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading plot(myLoadings, 0.95), & the alleles is a constant. \\ \begin{tabular}{ll} temp <- loading p
                                                                                                                                         main="Contribution of alleles \n to the first sPCA axis", fac=obj$loc.fac, cex.fac=0.6)
          temp
 $threshold
95%
0.02345973
  $var.names
[1] "L08.06"
[9] "L20.3"
                                                                                        "L08.07"
                                                                                                                                                 "L11.4"
                                                                                                                                                                                                         "L12.4"
                                                                                                                                                                                                                                                              "L14.11" "L16.02" "L16.10" "L17.2"
                                                                                        "L20.8
L12.4 L14.11 L16.02 L16.10
105 130 146 154
                                                                                                                                                                                                                                                                                                                                                                                    187
  $var.values
L08.06 L08.07 L11.4 L12.4 L14.11 L16.02 L16.10 0.03044687 0.03037709 0.06111338 0.03199067 0.02799529 0.02873923 0.02806079
                               L17.2
                                                                                                   L20.3
                                                                                                                                                                        L20.8
 0.05793290 0.08246000 0.02368209
```

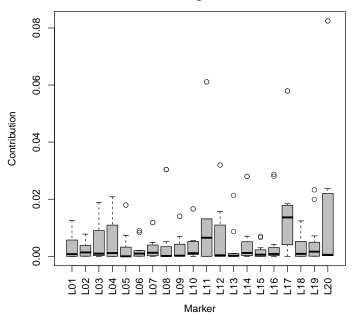


But to assess the average contribution of each marker, the boxplot probably is a better tool:

Alleles

```
> boxplot(myLoadings~obj$loc.fac, las=3, ylab="Contribution", xlab="Marker",
+ main="Contributions by markers \nto the first global score", col="grey")
```

Contributions by markers to the first global score



2 Case study: spatial genetic structure of the chamois in the Bauges mountains

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.

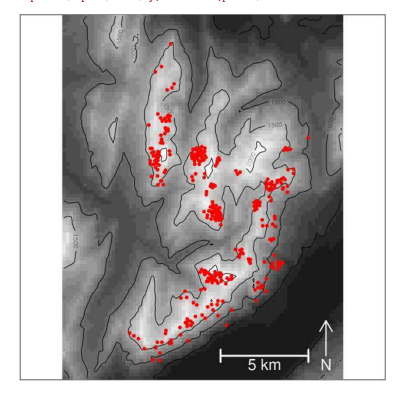
2.1 An overview of the data

We first load the data:

- > data(rupica)
- > rupica

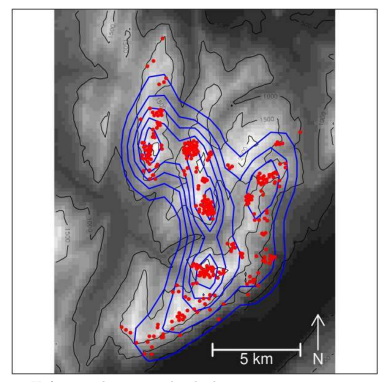
rupica is a genind object, that is, the class of objects storing genotypes (as opposed to population data) in *adegenet*. rupica also contains topographic information about the sampled area, which can be displayed by calling rupica\$other\$showBauges. Altitude maps are displayed using the *adehabitat* package [9]. The spatial distribution of the sampling can be displayed as follows:

> rupica\$other\$showBauges()
> points(rupica\$other\$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.

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



Unfortunately, geographical clustering is not strong enough to assign unambiguously each individual to a group. Therefore, we need to carry all analyses at the individual level, which precludes the use of most population genetics tools.

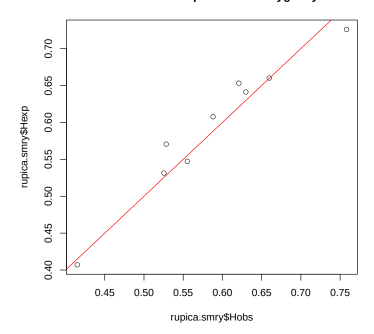
2.2 Summarising the genetic diversity

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)</pre>
```

- # Total number of genotypes: 335
- # Population sample sizes:

Observed vs expected heterozygosity



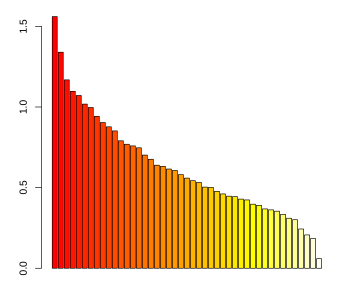
The red line indicate identity between both quantities. Observed heterozygosity do not seem to deviate massively from theoretical expectations. This is confirmed by a classical pairwise t-test::

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

We can seek a global picture of the genetic diversity among genotypes using a Principal Component Analysis (PCA, function dudi.pca in the ade4 package). The analysis is performed on a table of standardized alleles frequencies, obtained by scaleGen (use the binomial scaling option). Note that we disable the scaling option when performing the PCA, which would otherwise re-scale the data and therefore erase the previous scaling of scaleGen. The function dudi.pca displays a barplot of eigenvalues and asks for a number of retained principal components:

```
> rupica.X <- scaleGen(rupica, method="binom")
> rupica.pca1 <- dudi.pca(rupica.X, cent=FALSE, scale=FALSE, scannf=FALSE, nf=2)
> barplot(rupica.pca1$eig, main="Rupica dataset - PCA eigenvalues",
+ col=heat.colors(length(rupica.pca1$eig)))
```

Rupica dataset - PCA eigenvalues



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 = FALSE, scale = FALSE, scannf = FALSE,
    nf = 2)
$nf: 2 axis-components saved
$rank: 45
eigen values: 1.561 1.34 1.168 1.097 1.071 ...
  vector length mode
                        content
1 $cw
         55
                 numeric column weights
2 $1w
         335
                 numeric row weights
3 $eig
         45
                 numeric eigen values
  data.frame nrow ncol content
$tab 335 55 modifie
1 $tab
                        modified array
              335
2 $1i
                   2
                        row coordinates
3 $11
4 $co
              335
                         row normed scores
              55
                         column coordinates
```

column normed scores

In general, eigenvalues represent the amount of genetic diversity — as measured by the multivariate method being used — represented by each principal component (PC). An abrupt decrease in eigenvalues is likely to indicate the boundary between true patterns and non-interpretable structures. In this case, the first two PCs may contain some relevant biological signal.

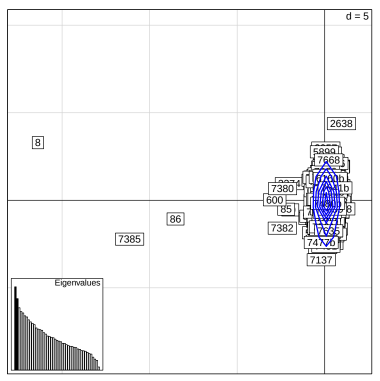
We can use s.label to display the two first components of the analysis. Kernel density estimation (s.kde2d) is used for a better assessment of the distribution of the genotypes onto the principal axes:

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

5 \$c1

55

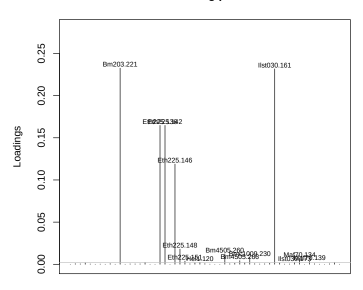
other elements: cent norm



This scatterplot shows that the only structure identified by PCA points to a few outliers. loadingplot confirms that this corresponds to the possession of a few original alleles:

> loadingplot(rupica.pca1\$c1^2)

Loading plot



Variables

We can go back to the genotypes for the concerned markers (e.g., Bm203) to check whether the highlighted genotypes are uncommon. truenames extracts the table of allele frequencies from a genind object (restoring original labels for markers, alleles, and individuals):

```
> X <- truenames(rupica)
> class(X)

[1] "matrix"
> dim(X)

[1] 335 55

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

bm203.221

0 0.00597014925373134
330 1
```

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==0.5]
```

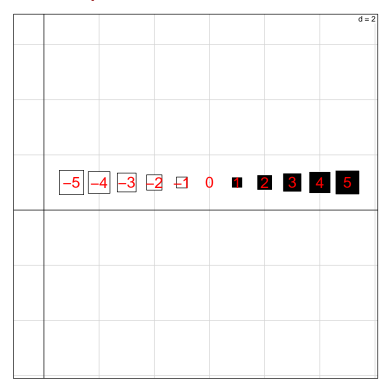
```
001 019 029 276
"8" "86" "600" "7385"
```

These are indeed our outliers. From the point of view of PCA, this would be the only structure in the data. However, further analyses show that more is to be seen...

2.3 Mapping and testing PCA results

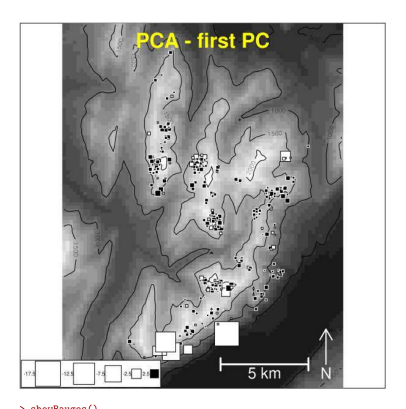
A frequent practice in spatial genetics is mapping the first principal components (PCs) onto the geographic space. ade4's function s.value is well-suited to do so, using black and white squares of variable size for positive and negative values. To give a legend for this type of representation:

```
> 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)
```

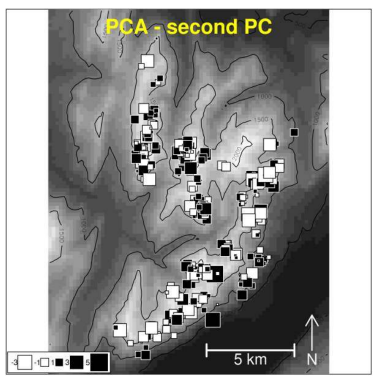


We apply this graphical representation to the first two PCs of the PCA:

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



```
> showBauges()
> s.value(rupica$other$xy, rupica.pca1$1i[,2], add.p=TRUE, csize=0.7)
> title("PCA - second PC",col.main="yellow",line=-2, cex.main=2)
```



As we can see, none of these PCs seems to display a particular spatial pattern. This visual assessment can be complemented by a test of spatial autocorrelation in these PCs. This can be achieved using Moran's I test. We use spdep's function ${\tt moran.mc}$ to perform these two tests. We first need 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, modelled as disks with a radius of 1150m. ${\tt chooseCN}$ is used to create the corresponding connection network:

```
> rupica.graph <- chooseCN(rupica$other$xy,type=5,d1=0,d2=2300, plot=FALSE, res="listw")
```

The connection network should ressemble this:

> rupica.graph

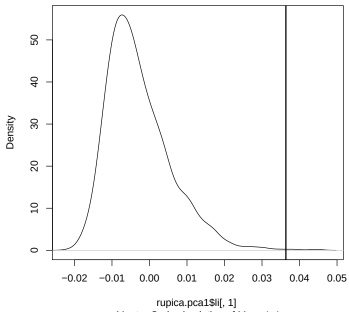
rupica.graph



We perform Moran's test for the first two PCs, and plot the results.

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

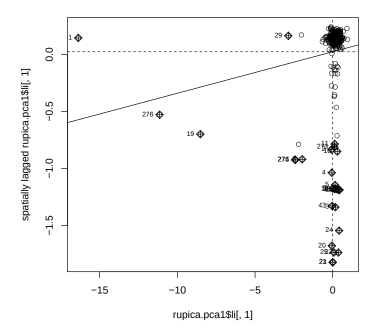
Density plot of permutation outcomes



Monte-Carlo simulation of Moran's I

This result is surprisingly significant. Why is this? Moran's plot (moran.plot) represents the tested variable against its lagged vector; we apply it to the first PC:

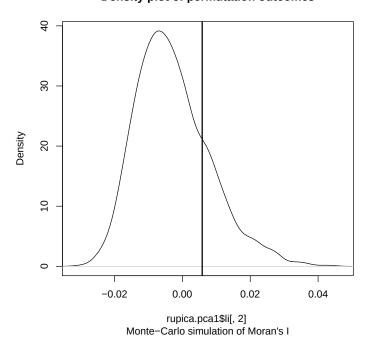
> moran.plot(rupica.pca1\$li[,1], rupica.graph)



Positive autocorrelation corresponds to a positive correlation between a variable and its lag vector. Here, we can see that this relation is entirely driven by the previously identified outliers, which turn out to be neighbours. This is therefore a fairly trivial and uninteresting pattern. Results obtained on the second PC are less surprisingly non-significant:

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

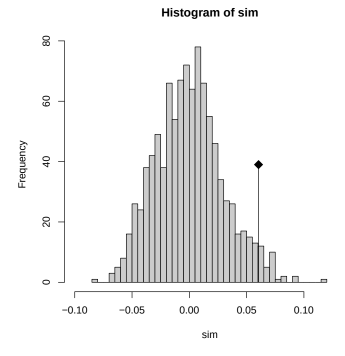
Density plot of permutation outcomes



2.4 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(rupica$other$xy))
> plot(mtest, nclass=30)
```



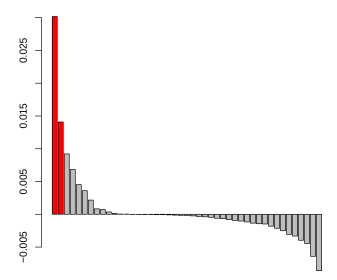
Interestingly, this test turns out to be marginally significant, and would encourage us to look for spatial patterns. This is the role of the spatial Principal Component Analysis.

2.5 Spatial Principal Component Analysis

We apply an sPCA to the \mathtt{rupica} dataset, using the connection network used previously in Moran's I tests:

```
> 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)), main="rupica dataset - sPCA eigenvalues")
```

rupica dataset - sPCA eigenvalues



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

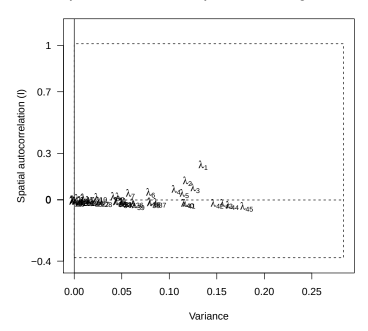
> rupica.spca1

```
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: 0.03018 0.01408 0.009211 0.006835 0.004529 ...
Negative eigenvalues: -0.008611 -0.006414 -0.004451 -0.003963 -0.003329 ...
  vector length mode
                        content
                numeric eigenvalues
1 $eig
  data.frame nrow ncol content
$c1 55 2 principa
                       principal axes: scaled vectors of alleles loadings
1 $c1
                       principal components: coordinates of entities ('scores')
2 $1i
             335 2
             335
3 $1s
                 2
                       lag vector of principal components
4 $as
             2
                       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)

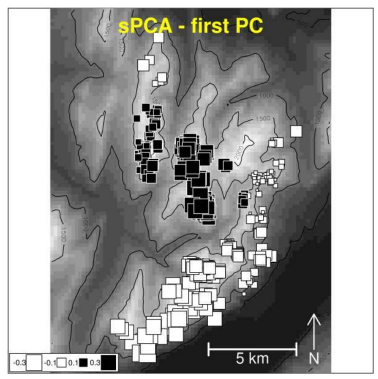
Spatial and variance components of the eigenvalues



While λ_1 indicates with no doubt a structure, the second eigenvalue, λ_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.

We map the sPCA results using s.value and lagged scores (\$1s) instead of the PC (\$1i), which are a 'denoisified' version of the PCs.

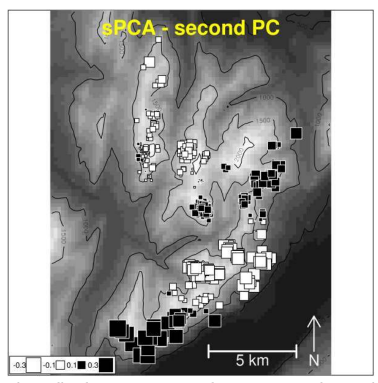
```
> showBauges()
> s.value(rupica$other$xy, rupica.spca1$ls[,1], add.p=TRUE, csize=0.7)
> title("sPCA - first PC",col.main="yellow",line=-2, cex.main=2)
```



This first PC shows a remarkably clear structure opposing two high-altitude areas separated by a valley, which is thought to be an obstacle to the dispersal of Chamois (due to higher exposition to predation in low-altitude areas).

The second PC of sPCA shows an equally interesting structure:

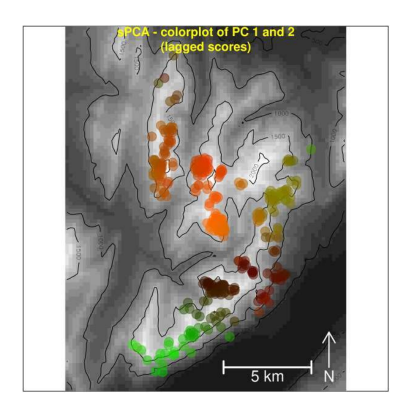
```
> showBauges()
> s.value(rupica$other$xy, rupica.spca1$ls[,2], add.p=TRUE, csize=0.7)
> title("sPCA - second PC",col.main="yellow" ,line=-2, cex.main=2)
```



The smaller clusters appearing on this map correspond to social units identified by direct observation in the field. Therefore, this genetic structure is merely a reflect of the social behaviour of these individuals.

Both genetic structures can be represented altogether using colorplot. The final figure should ressemble this (although colors may change from one computer to another):

```
> showBauges()
> colorplot(rupica$other$xy, rupica.spca1$ls, axes=1:2, transp=TRUE, add=TRUE, cex=3)
> title("sPCA - colorplot of PC 1 and 2\n(lagged scores)", col.main="yellow", line=-2, cex=2)
```



References

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