

RClone quickmanual several populations

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“Eager Beginners” Manual for RClone package

RClone data format: several populations

A. Introduction to RClone

RClone is a R package version of *GenClone* program: to analyse data (SSR, SNP, ...), test for clonality and describe spatial clonal organisation.

RClone allows:

1. Description of data set
 - discrimination of MLG (MultiLocus Genotypes);
 - test for reliability of data (in terms of loci and sampling).
2. Determination of MLL (MultiLocus Lineages)
 - psex/psex Fis with pvalue computation;
 - genetic distance matrix computation and threshold definition.
3. Genotypic diversity and evenness indices calculation
 - Simpson complement;
 - Shannon-Wiener diversity and evenness indices;
 - Hill's Simpson reciprocal;
 - Pareto index.

4. Spatial organisation of MLG/MLL

- spatial autocorrelation methods;
- clonal subrange estimation;
- Aggregation index and Edge Effect estimation.

Some of these analysis can be applied to dataset without clones.

B. RClone data format: several population

RClone functions works on diploid/haploid, one or several populations dataset.

If you have only one population in your dataset, go to other vignette *RClone_quickmanual*.

C. General format

If you have haploid data, you can skip to *D. Description of data set*.

An *RClone* table must look like:

```
library(RClone)
data(posidonia)
```

| Po15_1 | Po15_2 | Po4-3_1 | Po4-3_2 | Po5-10_1 | Po5-10_2 | Po5-39_1 | Po5-39_2 |
|--------|--------|---------|---------|----------|----------|----------|----------|
| 137 | 161 | 182 | 188 | 212 | 216 | 234 | 234 |
| 139 | 171 | 182 | 182 | 222 | 226 | 234 | 242 |
| 161 | 161 | 182 | 182 | 210 | 216 | 234 | 234 |
| 161 | 161 | 182 | 182 | 210 | 216 | 234 | 234 |
| 161 | 161 | 182 | 182 | 210 | 216 | 234 | 234 |
| 161 | 161 | 182 | 182 | 210 | 216 | 234 | 234 |
| 161 | 161 | 182 | 182 | 210 | 216 | 234 | 234 |
| 161 | 161 | 182 | 182 | 210 | 216 | 234 | 234 |
| 137 | 157 | 182 | 188 | 208 | 210 | 234 | 234 |
| 137 | 157 | 174 | 180 | 208 | 210 | 234 | 234 |

There is only one allele per column and, per locus, alleles are sorted by increasing order.

This is **mandatory** for all *RClone* functions.

As formatting can be source of error, we included functions to help formatting your diploid data:

1, The classic case: one locus per column

```
data(zostera)
head(zostera)
```

| population | x | y | GA35 | GA2 | GA17H | GA23 | GA12 | GA19 | GA20 | GA16 | GA17D |
|------------|-----|------|--------|--------|--------|--------|--------|--------|--------|--------|--------|
| SaintMalo | 0.0 | 18.0 | 185187 | 102116 | 131135 | 167169 | 131131 | 148148 | 162162 | 168168 | 197197 |
| SaintMalo | 0.0 | 15.5 | 185187 | 102116 | 131135 | 167169 | 131131 | 148148 | 162162 | 168168 | 197197 |
| SaintMalo | 0.0 | 3.5 | 187187 | 102116 | 127133 | 161169 | 131131 | 148148 | 162162 | 168168 | 197197 |
| SaintMalo | 2.0 | 3.5 | 187187 | 102116 | 133135 | 159161 | 131131 | 148148 | 162162 | 168168 | 197197 |
| SaintMalo | 6.5 | 18.0 | 187187 | 102116 | 135141 | 169169 | 131137 | 148150 | 162162 | 168168 | 197197 |
| SaintMalo | 6.5 | 10.0 | 187187 | 116116 | 133133 | 167167 | 131131 | 148148 | 162162 | 168168 | 195197 |

Zostera data is composed of:

- * a first column with population indication;
- * a second and third columns with x/y coordinates;
- * a genotypic dataset.

```
popvec <- zostera[,1] #futur vecpop
coord_zostera <- zostera[,2:3] #futur coordinates
zostera <- zostera[,4:ncol(zostera)] #dataset

zostera <- convert_GC(zostera, 3) #We used "3" because this is the length of each allele.
```

```
head(zostera)
```

| GA35_1 | GA35_2 | GA2_1 | GA2_2 | GA17H_1 | GA17H_2 | GA23_1 |
|--------|--------|-------|-------|---------|---------|--------|
| 185 | 187 | 102 | 116 | 131 | 135 | 167 |
| 185 | 187 | 102 | 116 | 131 | 135 | 167 |
| 187 | 187 | 102 | 116 | 127 | 133 | 161 |
| 187 | 187 | 102 | 116 | 133 | 135 | 159 |
| 187 | 187 | 102 | 116 | 135 | 141 | 169 |
| 187 | 187 | 116 | 116 | 133 | 133 | 167 |

2, The simple case: you already have a one-allele per column table

Just remove the pop/coords informations as above and sort your alleles:

```
sort_all(zostera)
```

3, You already work with Adegenet

It's a kind of like the case number 1, but you have to export your `genind` data into table first:

```
#library(adegenet)
#with data1, a genind object from Adegetnet:

test <- genind2df(data1)
data2 <- convert_GC(test, 3, "/")
#only if yours alleles are of length "3"
```

D. Description of data set

D.1 Discrimination of MLG

List unique alleles per locus:

Basic commands:

```
list_all_tab(zostera, vecpop = popvec)
```

or, for haploid data:

```
list_all_tab(haplodata, haploid = TRUE, vecpop = haplovec)
```

Results:

```
list_all_tab(zostera, vecpop = popvec)
```

#SaintMalo

| locus_1 | locus_2 | locus_3 | locus_4 | locus_5 | locus_6 | locus_7 | locus_8 | locus_9 |
|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| 185 | 102 | 131 | 167 | 131 | 148 | 162 | 168 | 197 |
| 187 | 116 | 127 | 161 | 137 | 150 | | | 195 |
| 189 | | 133 | 159 | | | | | |
| | | 135 | 169 | | | | | |
| | | 137 | 163 | | | | | |
| | | 119 | | | | | | |
| | | 141 | | | | | | |

#Arcouest

| locus_1 | locus_2 | locus_3 | locus_4 | locus_5 | locus_6 | locus_7 | locus_8 | locus_9 |
|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| 187 | 102 | 131 | 161 | 131 | 150 | 162 | 168 | 197 |
| 189 | 116 | 129 | 169 | | 148 | 156 | 166 | |
| 185 | 108 | 141 | 167 | | | 160 | | |
| | 118 | 133 | | | | 166 | | |
| | 120 | 143 | | | | 164 | | |
| | | 135 | | | | | | |

List MLG:

Basic commands:

```
MLG_tab(zostera, vecpop = popvec)
```

or, for haploid data:

```
MLG_tab(haplodata, vecpop = haplovec)
```

Results:

```
MLG_tab(zostera, vecpop = popvec)[[1]]
#SaintMalo
```

| unit_1 | unit_2 | unit_3 | unit_4 | unit_5 |
|--------|--------|--------|--------|--------|
| 1 | 2 | | | |
| 3 | | | | |
| 4 | 11 | | | |
| 5 | 7 | 8 | 9 | 12 |
| 6 | | | | |

Allelic frequencies:

Basic commands:

```
freq_RR(zostera, vecpop = popvec)
```

or, for haploid data:

```
freq_RR(haplodata, haploid = TRUE, vecpop = haplovec)
```

Options:

```
freq_RR(zostera, vecpop = popvec) #on ramets
freq_RR(zostera, vecpop = popvec, genet = TRUE) #on genets
freq_RR(zostera, vecpop = popvec, RR = TRUE) #Round-Robin methods
```

Results:

```
freq_RR(zostera, vecpop = popvec)[[1]]
#SaintMalo
```

| locus | allele | freq_ramet | freq_genet | freq_RR |
|---------|--------|------------|------------|-----------|
| locus_1 | 185 | 0.0517241 | 0.0588235 | 0.0588235 |
| locus_1 | 187 | 0.9137931 | 0.8823529 | 0.8823529 |
| locus_1 | 189 | 0.0344828 | 0.0588235 | 0.0588235 |
| locus_2 | 102 | 0.5000000 | 0.5000000 | 0.5000000 |
| locus_2 | 116 | 0.5000000 | 0.5000000 | 0.5000000 |
| locus_3 | 119 | 0.0172414 | 0.0294118 | 0.0294118 |
| locus_3 | 127 | 0.0689655 | 0.1176471 | 0.1176471 |

D.2 Test for reliability of data

On loci

Basic commands:

```
sample_loci(zostera, vecpop = popvec, nbrepeat = 1000)
```

or, for haploid data:

```
sample_loci(haplodata, haploid = TRUE, vecpop = haplovec, nbrepeat = 1000)
```

Options:

```
sample_loci(zostera, vecpop = popvec, nbrepeat = 1000, He = TRUE) #with He results
sample_loci(zostera, vecpop = popvec, nbrepeat = 1000, graph = TRUE) #graph displayed
sample_loci(zostera, vecpop = popvec, nbrepeat = 1000, bar = TRUE)
#progression bar, could be time consuming
sample_loci(zostera, vecpop = popvec, nbrepeat = 1000, export = TRUE)
#graph export in .eps
```

Results:

```
res <- sample_loci(zostera, vecpop = popvec, nbrepeat = 1000, He = TRUE)
names(res)
```

```
> [1] "SaintMalo" "Arcouest"
```

```
names(res$SaintMalo)
```

```
names(resvigncont2$res2_SU1$SaintMalo)
```

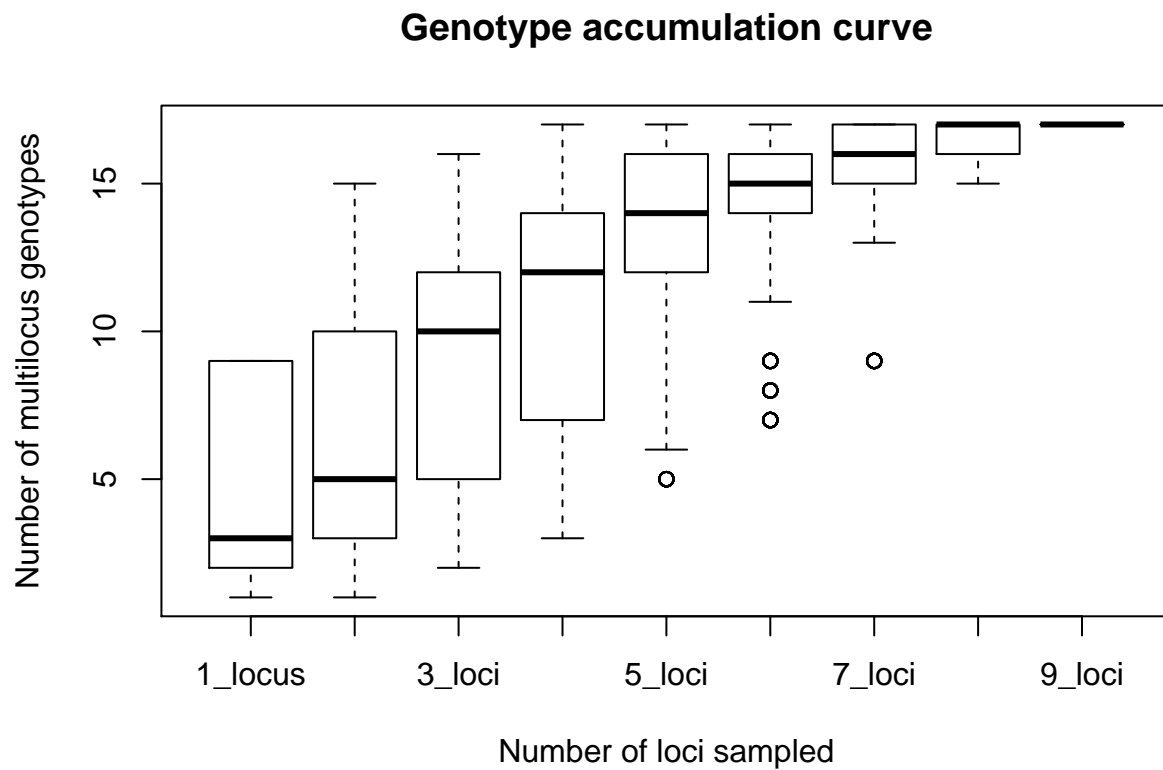
```
#Results: MLG
res$Arcouest$res_MLG
```

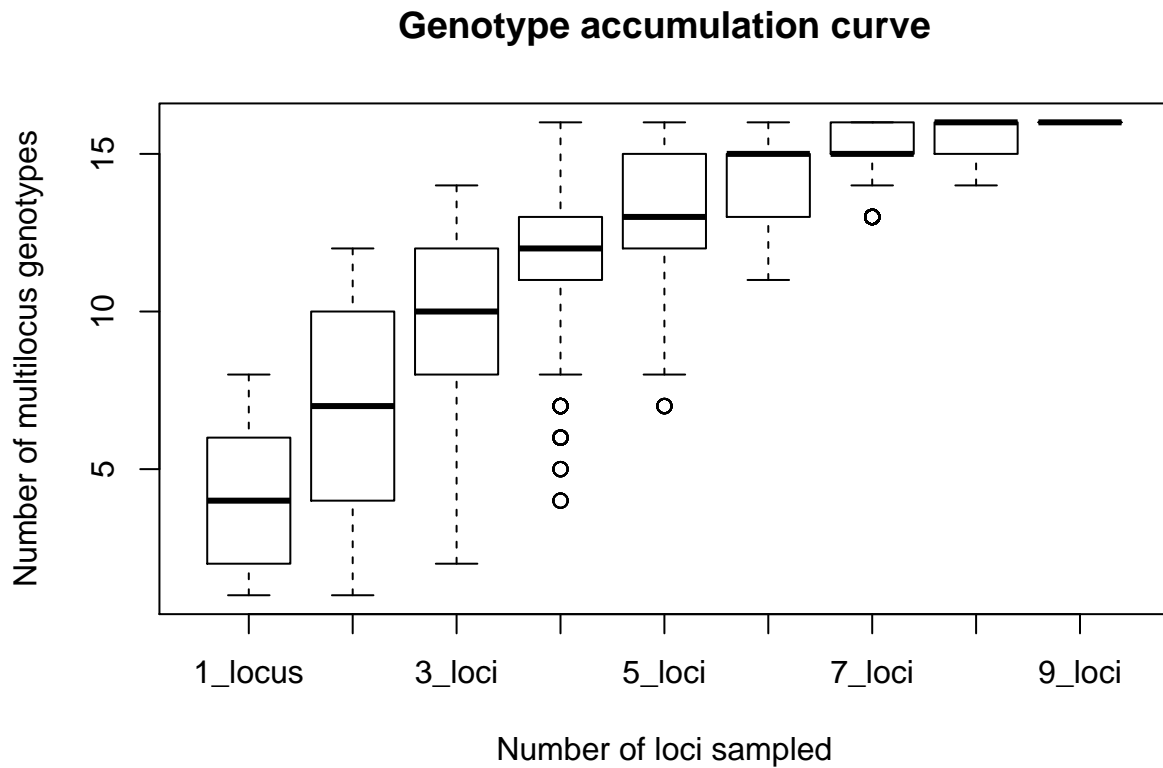
| nb_loci | min | max | mean_MLG | SE |
|---------|-----|-----|----------|-----------|
| 1 | 1 | 8 | 3.861 | 0.0819780 |
| 2 | 1 | 12 | 6.997 | 0.1021539 |
| 3 | 2 | 14 | 9.800 | 0.0873248 |
| 4 | 4 | 16 | 11.757 | 0.0678864 |
| 5 | 7 | 16 | 13.157 | 0.0555647 |
| 6 | 11 | 16 | 14.299 | 0.0414405 |
| 7 | 13 | 16 | 15.061 | 0.0307933 |
| 8 | 14 | 16 | 15.576 | 0.0210396 |
| 9 | 16 | 16 | 16.000 | 0.0000000 |

```
#Results: alleles  
res$Arcouest$res_alleles
```

```
#Results: raw data  
#res$Arcouest$raw_He  
#res$Arcouest$raw_MLG  
#res$Arcouest$raw_all
```

```
boxplot(res$SaintMalo$raw_MLG, main = "Genotype accumulation curve",  
        xlab = "Number of loci sampled", ylab = "Number of multilocus genotypes")
```





Same on units

Basic commands:

```
sample_units(zostera, vecpop = popvec, nbrepeat = 1000)
```

or, for haploid data:

```
sample_units(haplodata, haploid = TRUE, vecpop = haplovec, nbrepeat = 1000)
```

E. Determination of MLL

E.1 psex/psex Fis with pvalue computation

pgen, psex and p-values

Basic commands:

```
pgen(zostera, vecpop = popvec)
data(factor) #for psex
psex(zostera, vecpop = popvec)
```

or, for haploid data:


```
pgen(haplodata, haploid = TRUE, vecpop = haplovec)
data(factorR) #for psex
psex(haplodata, haploid = TRUE, vecpop = haplovec)
```

Options: (*idem on psex and pgen*)

```
#allelic frequencies computation:
psex(zostera, vecpop = popvec) #psex on ramets
psex(zostera, vecpop = popvec, genet = TRUE) #psex on genets
psex(zostera, vecpop = popvec, RR = TRUE) #psex with Round-Robin method
#psex computation
psex(zostera, vecpop = popvec) #psex with one psex per replica
psex(zostera, vecpop = popvec, MLGsim = TRUE) #psex MLGsim method
#pvalues:
psex(zostera, vecpop = popvec, nbrepeat = 100) #with p-values
psex(zostera, vecpop = popvec, nbrepeat = 1000, bar = TRUE)
#with p-values and a progression bar
```

Results:

```
data(factorR)
res <- psex(zostera, vecpop = popvec, RR = TRUE, nbrepeat = 1000)
res$Arcouest[[1]]
#if nbrepeat != 0, res contains a table of psex values and a vector of sim-psex values
```

| pgen | genet | psex | pvalue |
|-----------|-------|---------------------|---------------------|
| 0.0001692 | | | |
| 0.0000416 | | | |
| 0.0000900 | | | |
| 0.0000416 | 2 | 0.00124641891362029 | 0.00571428571428571 |
| 0.0000000 | | | |
| 0.0001800 | | | |

```
res$Arcouest[[2]] #a part of sim-psex values
```

```
> [1] 0.005507812 0.080799100 0.073342047 0.080799100 0.019908965
> [6] 0.008798312 0.002194897 0.003586792 0.046359147 0.116134553
```

Fis, pgen Fis, psex Fis and p-values

Not for haploid data !

Fis

Basic commands:

```
Fis(zostera, vecpop = popvec)
```

Options:

```
Fis(zostera, vecpop = popvec) #Fis on ramets
Fis(zostera, vecpop = popvec, genet = TRUE) #Fis on genets
Fis(zostera, vecpop = popvec, RR = TRUE) #Fis with Round-Robin methods
#RR = TRUE contains two results : a table with allelic frequencies
#and a table with Fis results
```

Results:

```
Fis(zostera, vecpop = popvec, RR = TRUE)$Arcouest[[2]]
```

| locus | Hobs | Hatt | Fis |
|---------|-----------|-----------|------------|
| locus_1 | 0.2666667 | 0.3300242 | 0.1919786 |
| locus_2 | 0.7500000 | 0.6995968 | -0.0720461 |
| locus_3 | 0.6250000 | 0.7721774 | 0.1906005 |
| locus_4 | 0.6250000 | 0.5383065 | -0.1610487 |
| locus_5 | 0.0000000 | 0.0000000 | NaN |
| locus_6 | 0.2000000 | 0.1862069 | -0.0740741 |
| locus_7 | 0.2857143 | 0.3772941 | 0.2427280 |
| locus_8 | 0.1875000 | 0.1754032 | -0.0689655 |
| locus_9 | 0.0000000 | 0.0000000 | NaN |

pgen Fis, psex Fis and p-values

Basic commands: (*idem* for *pgen_Fis* and *psex_Fis*)

```
pgen_Fis(zostera, vecpop = popvec)
```

Options:

```
#allelic frequencies:
psex_Fis(zostera, vecpop = popvec) #psex Fis on ramets
psex_Fis(zostera, vecpop = popvec, genet = TRUE) #psex Fis on genets
psex_Fis(zostera, vecpop = popvec, RR = TRUE) #psex Fis with Round-Robin method
#psex computation
psex_Fis(zostera, vecpop = popvec) #psex Fis, one for each replica
psex_Fis(zostera, vecpop = popvec, MLGsim = TRUE) #psex Fis with MLGsim method
#pvalues
psex_Fis(zostera, vecpop = popvec, nbrepeat = 100) #with p-values
psex_Fis(zostera, vecpop = popvec, nbrepeat = 1000, bar = TRUE)
#with p-values and a progression bar
```

Results:

```
data(factorR)
res <- psex_Fis(zostera, vecpop = popvec, RR = TRUE, nbrepeat = 1000)
res$Arcouest[[1]]
#if nbrepeat != 0, res contains a table of psex values and a vector of sim-psex Fis values
```

| p _{gen} Fis | genet | p _{sex} Fis | pvalue |
|----------------------|-------|----------------------|--------------------|
| 0.0001459 | | | |
| 0.0000587 | | | |
| 0.0000708 | | | |
| 0.0000587 | 2 | 0.00175772228659339 | 0.0154639175257732 |
| 0.0000003 | | | |
| 0.0001417 | | | |

```
res$Arcouest[[2]] #a part of sim psex Fis values
```

```
> [1] 0.046691095 0.065845604 0.047964606 0.030472230 0.029045389
> [6] 0.009682317 0.101570928 0.200020973 0.004458226 0.033269017
```

E.2 MultiLocus Lineages

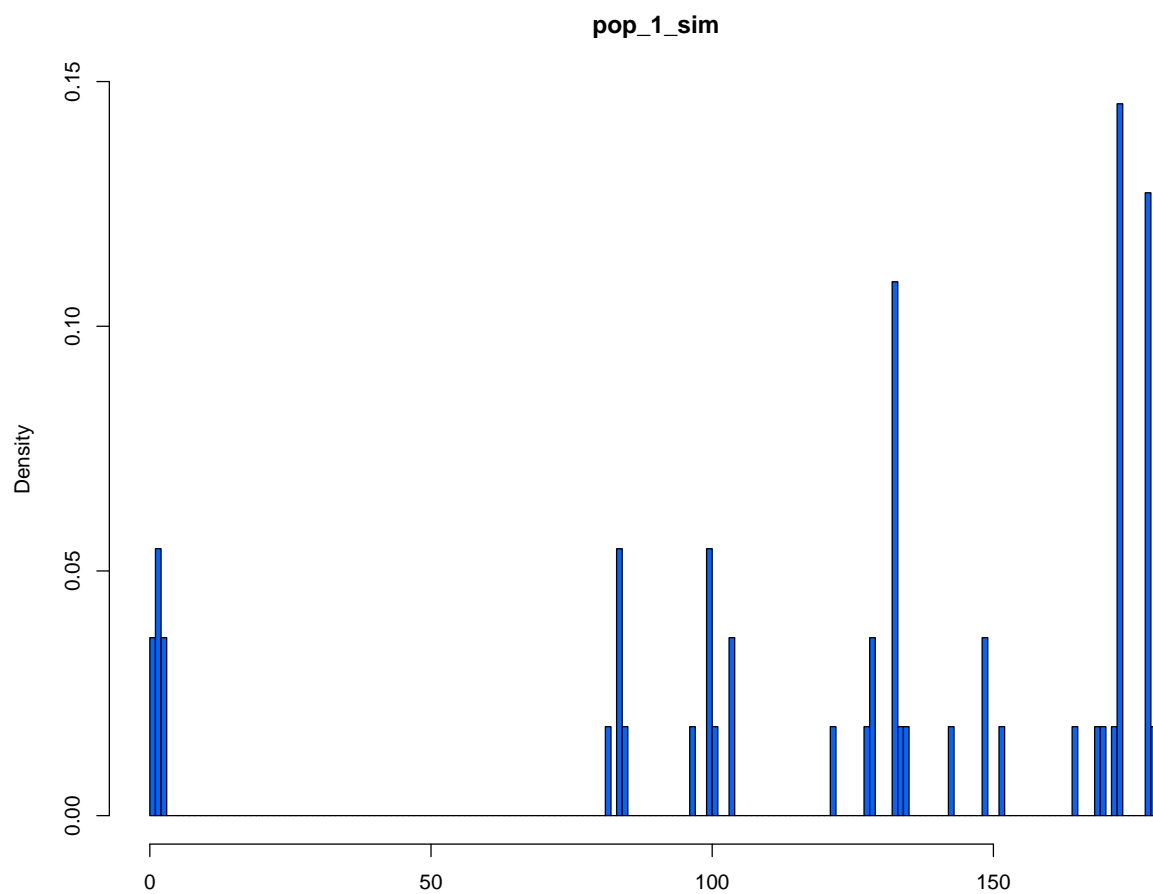
Genetic distance matrix computation and threshold definition

On a theoretical diploid population with $c = 0.9999$ (c , clonality rate).

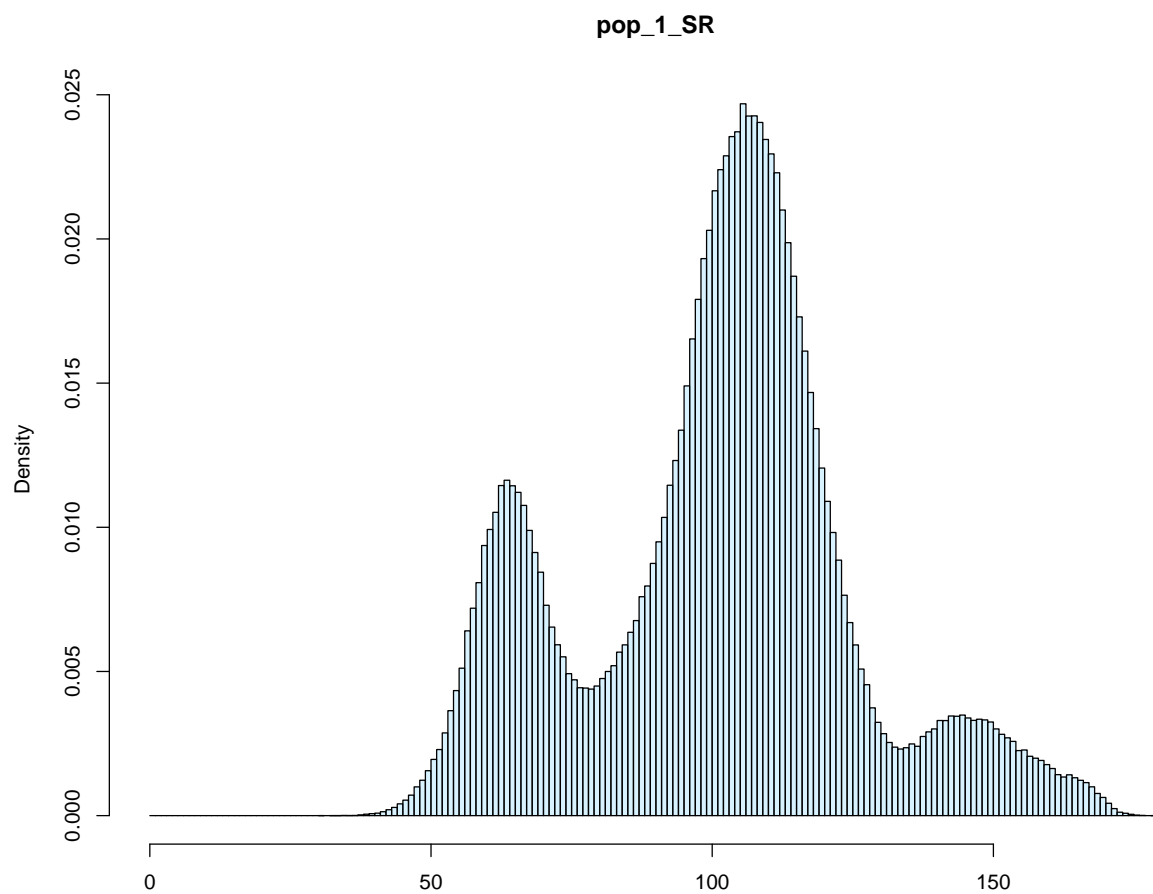
```
data(popsim)
vecsim <- c(rep(1,50), rep(2,50))

#genetic distances computation, distance on allele differences:
respop <- genet_dist(popsim, vecpop = vecsim)
ressim <- genet_dist_sim(popsim, vecpop = vecsim, nbrepeat = 1000)
#theoretical distribution: sexual reproduction
ressimWS <- genet_dist_sim(popsim, vecpop = vecsim, genet = TRUE, nbrepeat = 1000)
#idem, without selfing

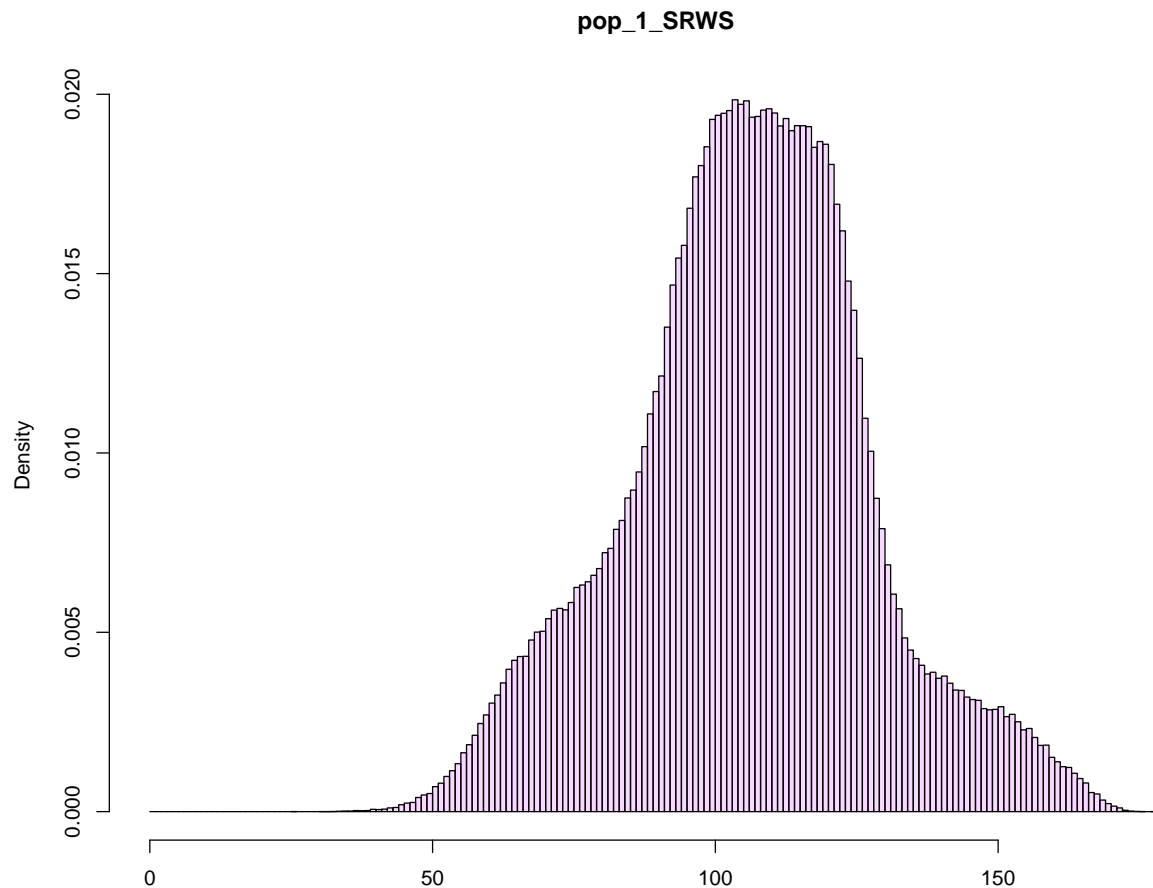
#graph prep.:
#first pop:
p1 <- hist(respop[[1]]$distance_matrix, freq = FALSE, col = rgb(0,0.4,1,1),
           breaks = seq(0, max(respop[[1]]$distance_matrix)+1, 1),
           main = "pop_1_sim", xlab = "")
```



```
p2 <- hist(ressim[[1]]$distance_matrix, freq = FALSE, col = rgb(0.7,0.9,1,0.5),
           breaks = seq(0, max(ressim[[1]]$distance_matrix)+1, 1),
           main = "pop_1_SR", xlab = "")
```



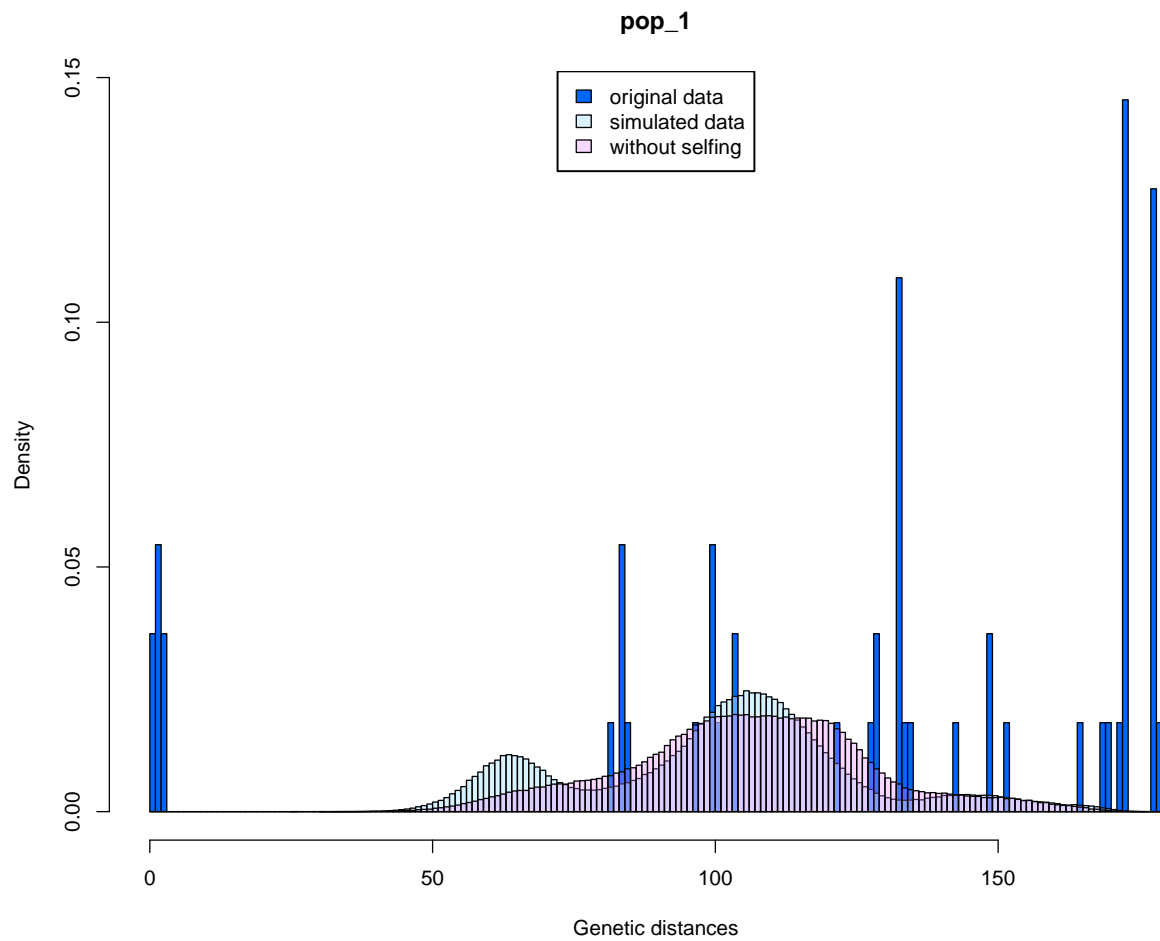
```
p3 <- hist(ressimWS[[1]]$distance_matrix, freq = FALSE, col = rgb(0.9,0.5,1,0.3),  
           breaks = seq(0, max(ressimWS[[1]]$distance_matrix)+1, 1),  
           main = "pop_1_SRWS", xlab = "")
```



```
limx <- max(max(respop[[1]]$distance_matrix), max(ressim[[1]]$distance_matrix),
            max(ressimWS[[1]]$distance_matrix))

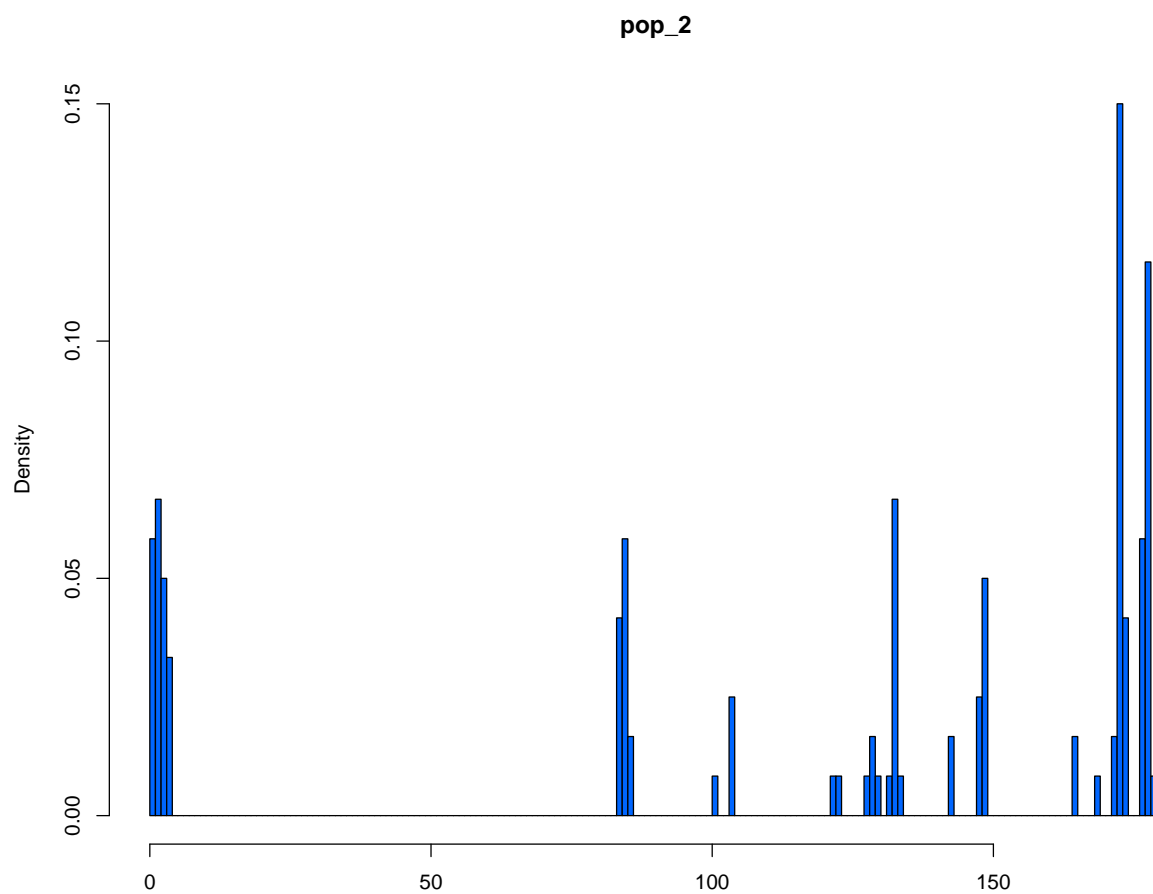
#graph superposition:
plot(p1, col = rgb(0,0.4,1,1), freq = FALSE, xlim = c(0,limx),
     main = paste("pop", unique(vccsim)[[1]], sep = "_"),
     xlab = "Genetic distances")
plot(p2, col = rgb(0.7,0.9,1,0.5), freq = FALSE, add = TRUE)
plot(p3, col = rgb(0.9,0.5,1,0.3), freq = FALSE, add = TRUE)

#adding a legend:
leg.txt <- c("original data", "simulated data", "without selfing")
col <- c(rgb(0,0.4,1,1), rgb(0.7,0.9,1,0.5), rgb(0.9,0.5,1,0.3))
legend("top", fill = col, leg.txt, plot = TRUE, bty = "o", box.lwd = 1.5,
      bg = "white")
```

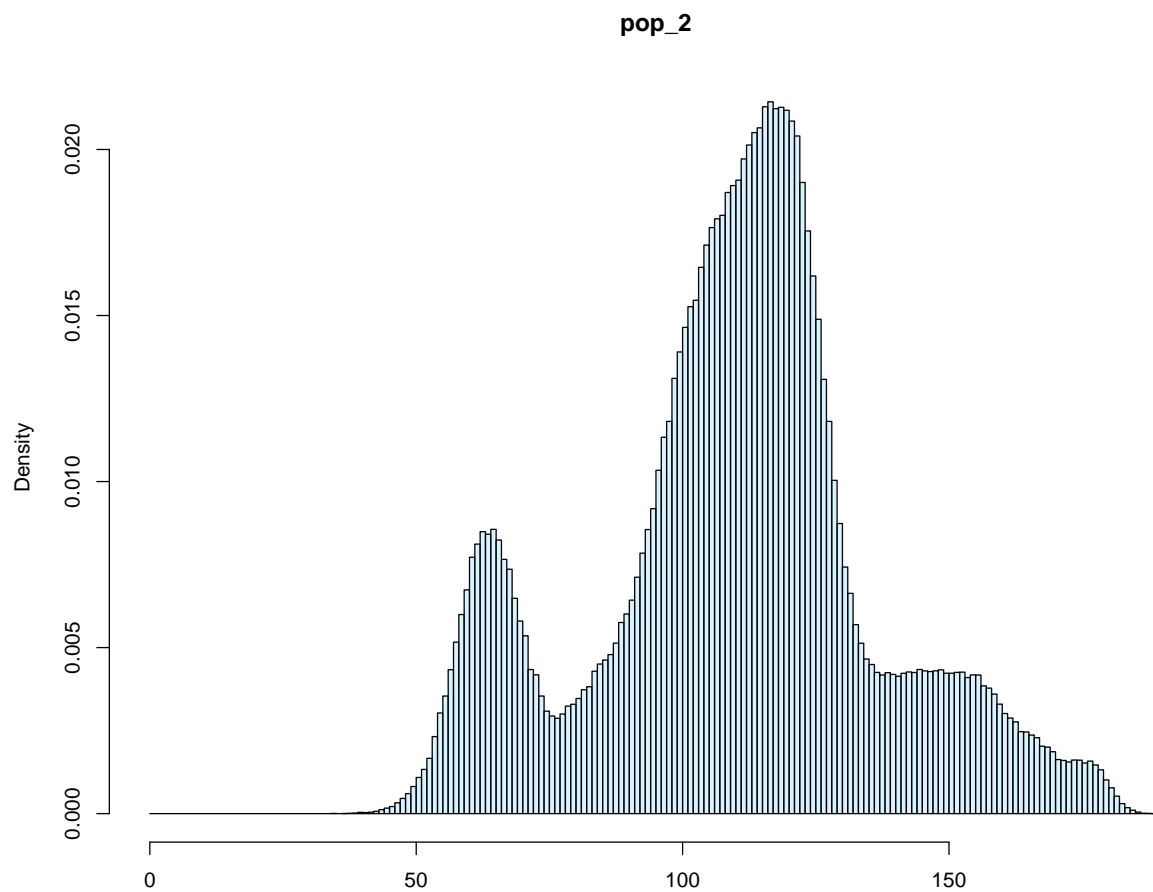


```
#second pop:
p <- 2 #useful if several populations: just change *p* and run lines

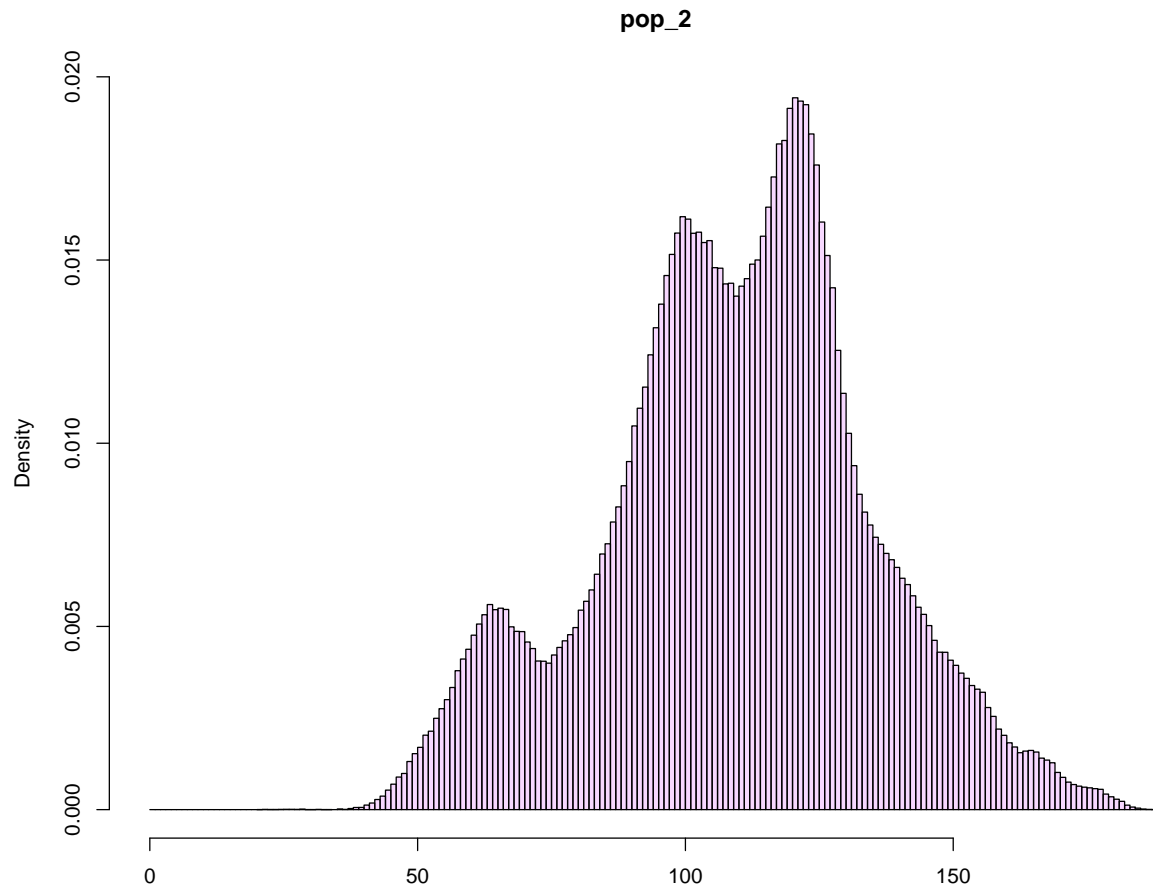
p1 <- hist(respop[[p]]$distance_matrix, freq = FALSE, col = rgb(0,0.4,1,1),
           breaks = seq(0, max(respop[[p]]$distance_matrix)+1, 1),
           main = paste("pop", p, sep = "_"), xlab = "")
```



```
p2 <- hist(ressim[[p]]$distance_matrix, freq = FALSE, col = rgb(0.7,0.9,1,0.5),  
           breaks = seq(0, max(ressim[[p]]$distance_matrix)+1, 1),  
           main = paste("pop", p, sep = "_"), xlab = "")
```

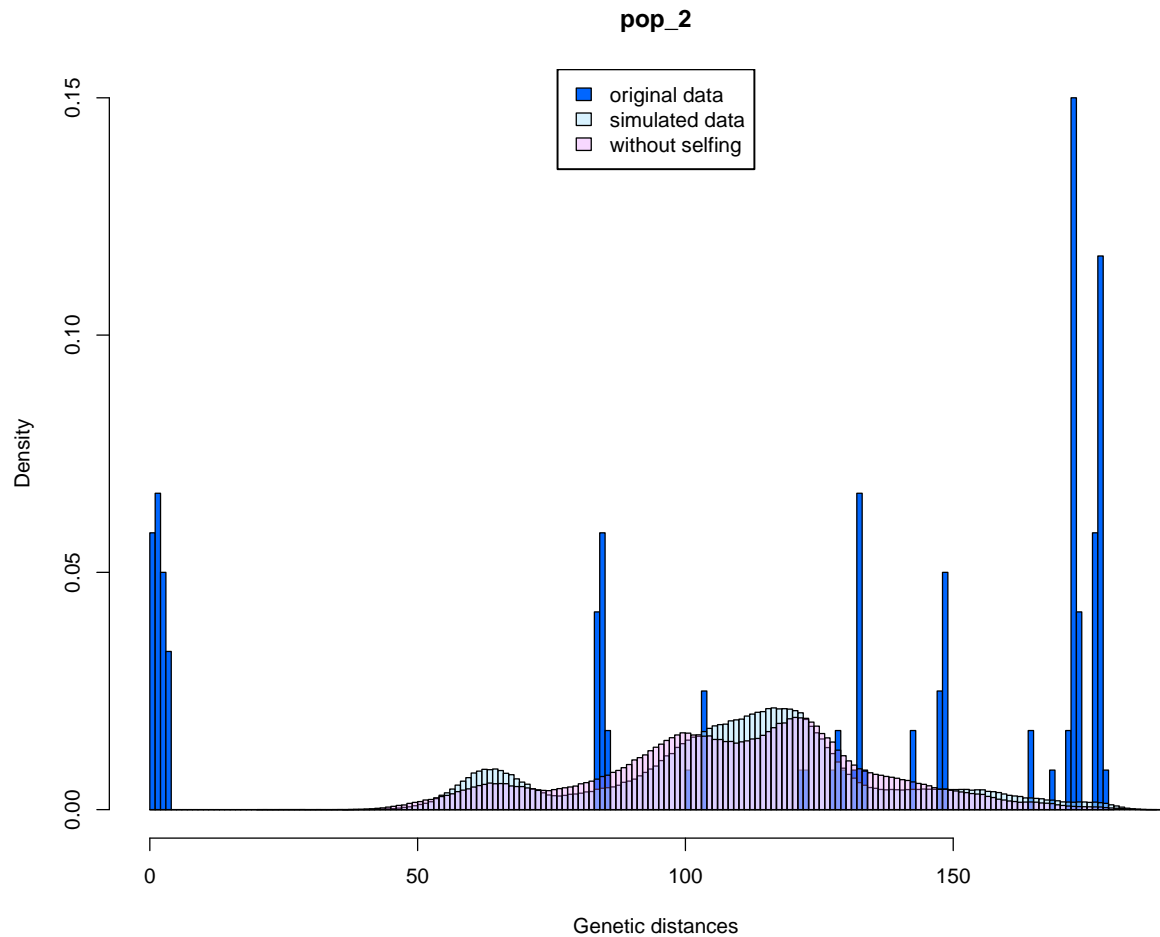
```
p3 <- hist(ressimWS[[p]]$distance_matrix, freq = FALSE, col = rgb(0.9,0.5,1,0.3),  
           breaks = seq(0, max(ressimWS[[p]]$distance_matrix)+1, 1),  
           main = paste("pop", p, sep = "_"), xlab = "")
```



```
limx <- max(max(respop[[p]]$distance_matrix), max(ressim[[p]]$distance_matrix),
            max(ressimWS[[p]]$distance_matrix))

#graph superposition:
plot(p1, col = rgb(0,0.4,1,1), freq = FALSE, xlim = c(0,limx),
     main = paste("pop", unique(vccsim)[[p]], sep = "_"),
     xlab = "Genetic distances")
plot(p2, col = rgb(0.7,0.9,1,0.5), freq = FALSE, add = TRUE)
plot(p3, col = rgb(0.9,0.5,1,0.3), freq = FALSE, add = TRUE)

#adding a legend:
leg.txt <- c("original data", "simulated data", "without selfing")
col <- c(rgb(0,0.4,1,1), rgb(0.7,0.9,1,0.5), rgb(0.9,0.5,1,0.3))
legend("top", fill = col, leg.txt, plot = TRUE, bty = "o", box.lwd = 1.5,
      bg = "white")
```



```
#determining alpha2
table(respop[[1]]$distance_matrix)
>
> 1  2  3 82 84 85 97 100 101 104 122 128 129 133 134 135 143 149
> 2  3  2  1  3  1  1  3  1  2  1  1  2  6  1  1  1  2
> 152 165 169 170 172 173 178 179
> 1  1  1  1  1  8  7  1
#alpha2 = 3
```

```
#creating MLL list:
MLLlist <- MLL_generator(popsim, vecpop = vecsim, alpha2 = c(3,0))
##This will create a list of MLL (alpha2 = 3) and MLG (alpha2 = 0) !

#or
res <- genet_dist(popsim, vecpop = vecsim, alpha2 = c(3,0))
MLLlist <- MLL_generator2(list(res[[1]]$potential_clones,
  res[[2]]$potential_clones), MLG_list(popsim, vecpop = vecsim), vecpop = vecsim)
```

For haploid data, theoretical example:

```

respop <- genet_dist(haplodata, haploid = TRUE, vecpop = vecchaplo)
ressim <- genet_dist_sim(haplodata, haploid = TRUE, vecpop = vecchaplo,
                        nbrepeat = 1000)
MLLlist <- MLL_generator(haplodata, haploid = TRUE, vecpop = vecchaplo,
                        alpha2 = c(3,0))
#or
res <- genet_dist(haplodata, haploid = TRUE, vecpop = vecchaplo, alpha2 = c(3,0))
MLLlist <- MLL_generator2(list(res[[1]]$potential_clones, res[[2]]$potential_clones),
                        haploid = TRUE, MLG_list(haplodata, vecpop = vecchaplo), vecpop = vecchaplo)

```

F. Genotypic diversity and evenness indices calculation

F.1 Classic genotypic indices

Basic commands:

```
clonal_index(zostera, vecpop = popvec)
```

or, with MLL:

```
clonal_index(popsim, vecpop = vecsim, listMLL = MLLlist)
```

or, for haploid data:

```
clonal_index(haplodata, vecpop = vecchaplo)
```

Results:

```
clonal_index(zostera, vecpop = popvec)
```

| | G | R | H'' | J' | D | V | Hill |
|-----------|----|-----------|----------|-----------|-----------|-----------|-----------|
| SaintMalo | 17 | 0.5714286 | 2.671294 | 0.9428497 | 0.9507389 | 0.8559028 | 20.300000 |
| Arcouest | 16 | 0.5172414 | 2.268605 | 0.8182264 | 0.8413793 | 0.3918367 | 6.304348 |

F.2 Pareto index

Basic commands:

```
Pareto_index(zostera, vecpop = popvec)
```

or, with MLL:

```
Pareto_index(popsim, vecpop = vecsim, listMLL = MLLlist)
```

or, for haploid data:

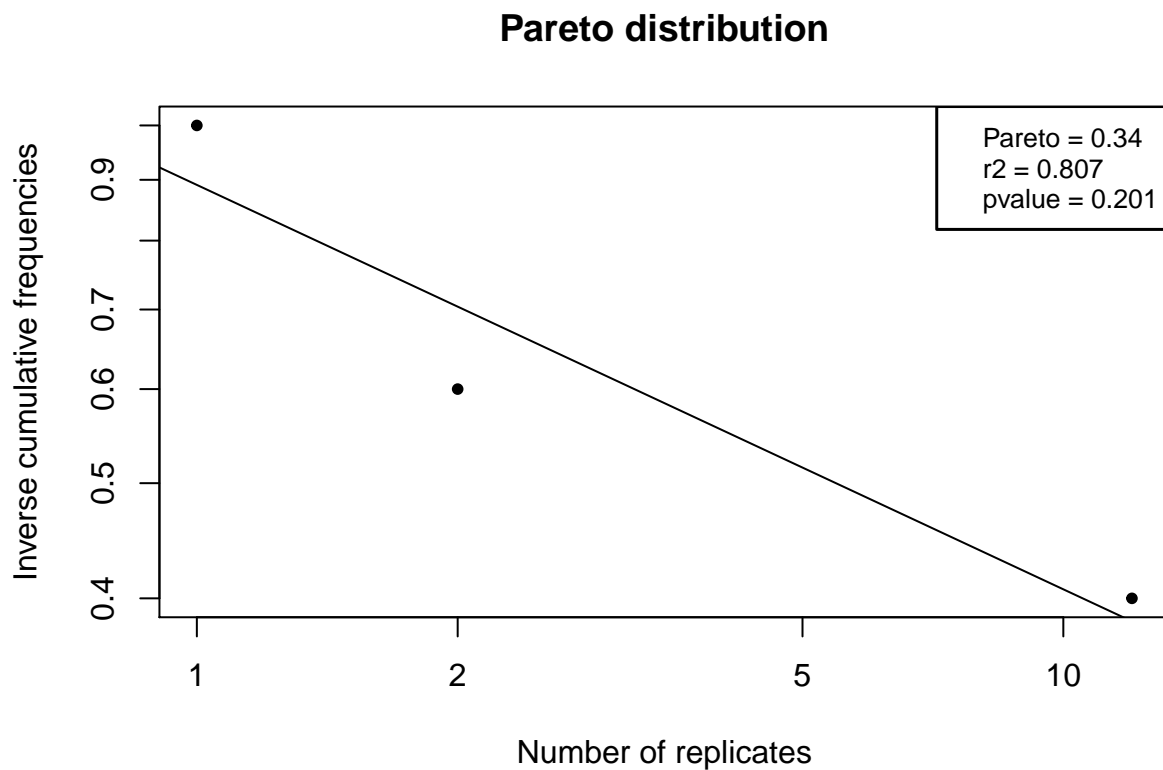
```
Pareto_index(haplodata, vecpop = vechaplo)
```

Options:

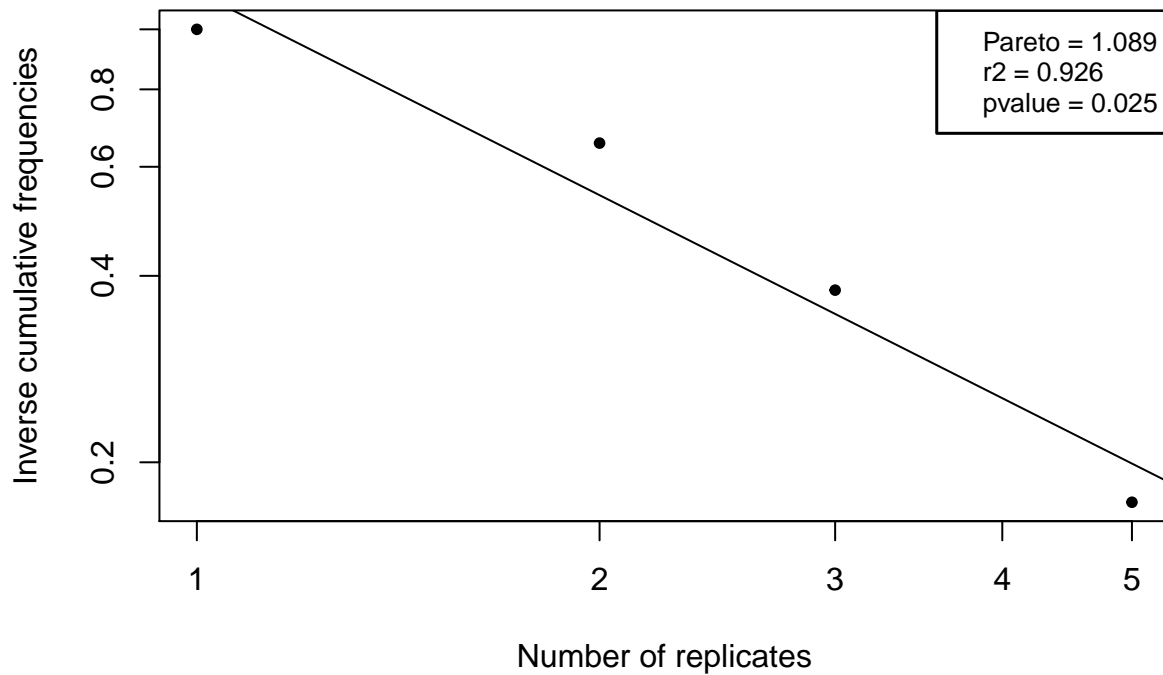
```
Pareto_index(zostera, vecpop = popvec, graph = TRUE) #classic graphic  
Pareto_index(zostera, vecpop = popvec, legends = 2, export = TRUE)  
                                                    #export option  
Pareto_index(zostera, vecpop = popvec, full = TRUE) #all results
```

Results:

```
res <- Pareto_index(zostera, vecpop = popvec, full = TRUE, graph = TRUE, legends = 2)
```



Pareto distribution



```
names(res$SaintMalo)
```

```
> [1] "Pareto"          "c_Pareto"        "regression_results"  
> [4] "coords_Pareto"
```

```
res$SaintMalo$Pareto
```

```
> [1] 0.3403204
```

```
res$SaintMalo$c_Pareto
```

```
> [1] 1.34032
```

```
#res$SaintMalo$regression_results  
#res$SaintMalo$coords_Pareto #points coordinates
```

G. Spatial description of clonality

G.1 Spatial autocorrelation

Basic commands:

```
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec, Loiselle = TRUE)
```

or, with MLL:

```
autocorrelation(popsim, coords = coord_sim, Loiselle = TRUE, listMLL = MLLlist)
```

or, for haploid data:

```
autocorrelation(haplodata, haploid = TRUE, coords = coord_haplo, Loiselle = TRUE)
```

Lot's of options:

```
#kinship distances:
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec, Loiselle = TRUE)
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec, Ritland = TRUE)

#ramets/genets methods:
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec, Loiselle = TRUE)
                                                                    #ramets
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec,, Loiselle = TRUE,
                genet = TRUE, central_coords = TRUE)
                                                                    #genets, central coordinates of each MLG
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec, Loiselle = TRUE,
                genet = TRUE, random_unit = TRUE)
                                                                    #genets, one random unit per MLG
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec, Loiselle = TRUE,
                genet = TRUE, weighted = TRUE)
                                                                    #genets, with weighted matrix on kinships

#distance classes construction:
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec, Loiselle = TRUE)
                                                                    #10 equidistant classes
distvec <- c(0,10,15,20,30,50,70,76.0411074)
                                                                    #with 0, min distance and 76.0411074, max distance
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec, Loiselle = TRUE,
                vecdist = distvec) #custom distance vector
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec, Loiselle = TRUE,
                class1 = TRUE, d = 7) #7 equidistant classes
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec, Loiselle = TRUE,
                class2 = TRUE, d = 7)
                                                                    #7 distance classes with the same number of units in each

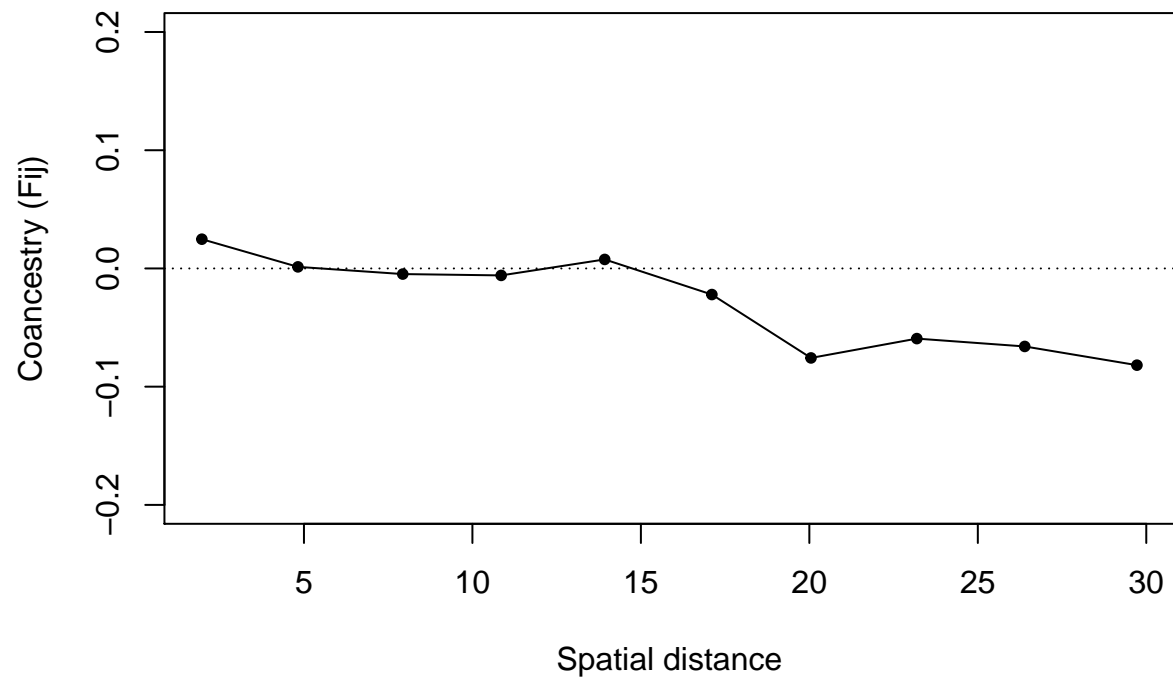
#graph options:
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec, Ritland = TRUE,
                graph = TRUE) #displays graph
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec, Ritland = TRUE,
                export = TRUE) #export graph

#pvalues computation
autocorrelation(zostera, coords = coord_zostera, vecpop = popvec, Ritland = TRUE,
                nbrepeat = 1000)
```

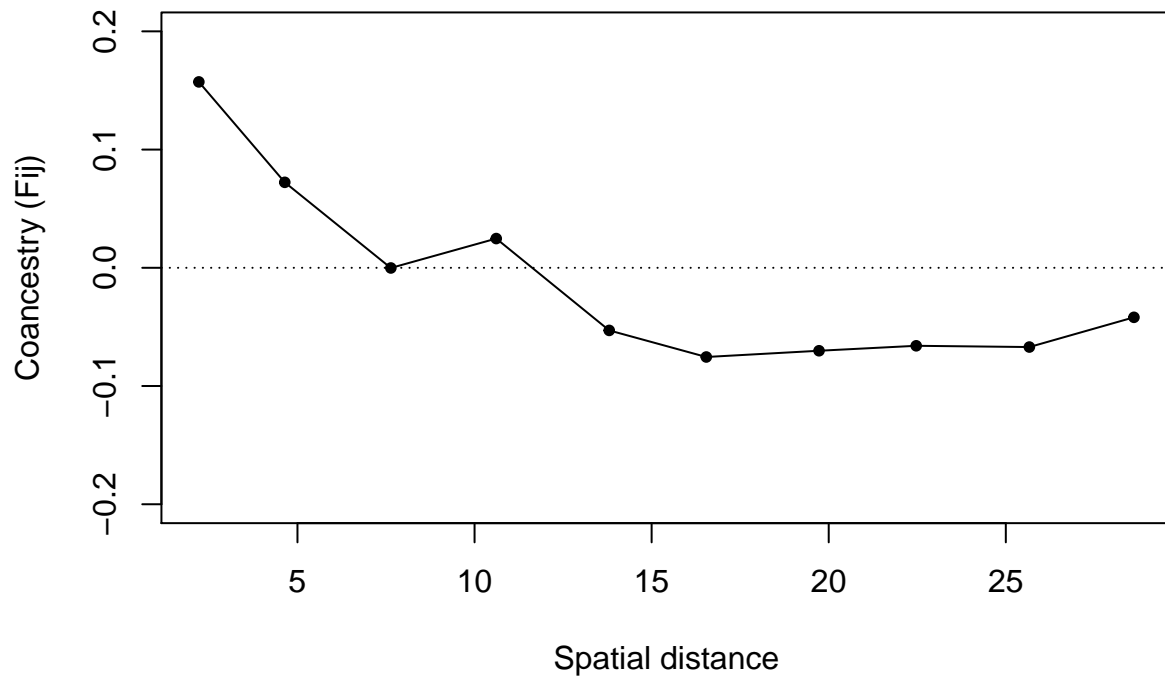
Results:

```
res <- autocorrelation(zostera, coords = coord_zostera, vecpop = popvec,  
                      Ritland = TRUE, nbrepeat = 1000, graph = TRUE)
```

Spatial autocorrelation analysis



Spatial autocorrelation analysis



```
names(res$Arcouest)
```

```
> [1] "Main_results"          "Slope_and_Sp_index"
> [3] "Slope_resample"        "Kinship_resample"
> [5] "Matrix_kinship_results" "Class_kinship_results"
> [7] "Class_distance_results"
```

```
res$Arcouest$Main_results #enables graph reproduction
```

| dist_min | dist_max | dist_mean | ln(dist_mean) | nb_pairs | mean_Ritland | pval_kin |
|----------|----------|-----------|---------------|----------|--------------|----------|
| 1.11803 | 3.00000 | 2.215687 | 0.7955625 | 23 | 0.1572378 | 0.000 |
| 3.04138 | 6.02080 | 4.641553 | 1.5350490 | 35 | 0.0723040 | 0.000 |
| 6.10328 | 9.05539 | 7.636613 | 2.0329541 | 62 | -0.0001518 | 0.292 |
| 9.12414 | 12.09339 | 10.612723 | 2.3620535 | 61 | 0.0247131 | 0.014 |
| 12.16553 | 15.13275 | 13.802695 | 2.6248639 | 58 | -0.0529723 | 0.022 |
| 15.18223 | 18.06931 | 16.543032 | 2.8059650 | 50 | -0.0754410 | 0.000 |
| 18.24829 | 21.10095 | 19.726507 | 2.9819633 | 54 | -0.0701760 | 0.000 |
| 21.18962 | 24.08319 | 22.470382 | 3.1121981 | 28 | -0.0659711 | 0.034 |
| 24.41311 | 27.22591 | 25.664587 | 3.2451121 | 20 | -0.0670280 | 0.068 |
| 27.45906 | 30.26962 | 28.617915 | 3.3540329 | 15 | -0.0418956 | 0.436 |

```
apply(res$Arcouest$Main_results, 2, mean)[6] #mean Fij
```

```
> mean_Ritland
> -0.0119381
```

```
res$Arcouest$Slope_and_Sp_index #gives b and Sp indices
```

| | b | b_log | Sp | Sp_log |
|-------------|------------|------------|------------|------------|
| obs_value | -0.0069054 | -0.0877033 | 0.0081938 | 0.1040665 |
| mean_sim | 0.0000230 | 0.0002994 | -0.0000088 | -0.0000719 |
| sd_sim | 0.0009620 | 0.0107159 | 0.0009555 | 0.0106701 |
| 0.95_inf | -0.0021680 | -0.0232025 | -0.0015771 | -0.0177145 |
| 0.95_sup | 0.0016229 | 0.0187066 | 0.0021527 | 0.0233927 |
| 0.9_inf | -0.0017065 | -0.0190059 | -0.0014359 | -0.0155080 |
| 0.9_sup | 0.0014775 | 0.0162403 | 0.0017695 | 0.0190726 |
| pval_upper | 0.0000000 | 0.0000000 | 1.0000000 | 1.0000000 |
| pval_lower | 1.0000000 | 1.0000000 | 0.0000000 | 0.0000000 |
| pval_2sides | 0.0000000 | 0.0000000 | 0.0000000 | 0.0000000 |

```
#raw data:
#res$Arcouest$Slope_resample
#res$Arcouest$Kinship_resample
#res$Arcouest$Matrix_kinship_results
#res$Arcouest$Class_kinship_results
#res$Arcouest$Class_distance_results
```

G.2 Clonal subrange

Basic commands:

```
clonal_sub(zostera, coords = coord_zostera, vecpop = popvec)
```

or, with MLL:

```
clonal_sub(popsim, coords = coord_sim, listMLL = MLLlist)
```

or, for haploid data:

```
clonal_sub(haplodata, haploid = TRUE, coords = coord_haplo)
```

Options: same distance classes definition as *autocorrelation*:

```
clonal_sub(posidonia, coords = coord_posidonia) #basic, with 10 equidistant classes
distvec <- c(0,10,15,20,30,50,70,76.0411074)
#with 0, min distance and 76.0411074, max distance
clonal_sub(zostera, coords = coord_zostera, vecpop = popvec, vecdist = distvec)
```

```

                                #custom distance classes
clonal_sub(zostera, coords = coord_zostera, vecpop = popvec, class1 = TRUE, d = 7)
                                #7 equidistant classes
clonal_sub(zostera, coords = coord_zostera, vecpop = popvec, class1 = TRUE, d = 7)
                                #7 distance classes with the same number of units in each

```

Results:

```

res <- clonal_sub(zostera, coords = coord_zostera, vecpop = popvec)
res$Arcouest[[1]] #Global clonal subrange
> [1] 19.10497

```

```

res$Arcouest$clonal_sub_tab #details per class

```

| nb_pairs | dist_min | dist_max | dist_mean | Fr | log(Fr) |
|----------|-----------|----------|-----------|------------|------------|
| 19 | 0.7071068 | 2.915476 | 1.969185 | 0.3157895 | -0.5006024 |
| 49 | 3.201562 | 6.184658 | 4.821457 | 0.3265306 | -0.4860761 |
| 49 | 6.264982 | 9.219544 | 7.93246 | 0.1836735 | -0.7359536 |
| 74 | 9.340771 | 12.36932 | 10.85383 | 0.1756757 | -0.7552884 |
| 77 | 12.5 | 15.43535 | 13.92621 | 0.1688312 | -0.7725474 |
| 70 | 15.5 | 18.5809 | 17.10734 | 0.1285714 | -0.8908555 |
| 36 | 18.72165 | 21.59282 | 20.04924 | 0.08333333 | -1.079181 |
| 31 | 21.70829 | 24.69818 | 23.18994 | 0 | -Inf |
| 25 | 24.82438 | 27.85678 | 26.39228 | 0 | -Inf |
| 5 | 28.41215 | 30.99193 | 29.72322 | 0 | -Inf |

G.3 Aggregation index

Basic commands:

```

agg_index(zostera, coords = coord_zostera, vecpop = popvec)

```

or, with MLL:

```

agg_index(popsim, coords = coord_sim, listMLL = MLLlist)

```

or, for haploid data:

```

agg_index(haplodata, coords = coord_haplo)

```

Options:

```

agg_index(zostera, coords = coord_zostera, vecpop = popvec, nbrepeat = 100)
                                #pvalue computation
agg_index(zostera, coords = coord_zostera, vecpop = popvec, nbrepeat = 1000,
                                bar = TRUE) #could be time consuming

```

Results:

```
res <- agg_index(zostera, coords = coord_zostera, vecpop = popvec, nbrepeat = 1000)
```

```
res$SaintMalo$results #Aggregation index
```

| Ac | pval | nbrepeat |
|-----------|-------|----------|
| 0.1965314 | 0.006 | 1000 |

```
#res$SaintMalo$simulation #vector of sim aggregation index
```

G.4 Edge Effect

Basic commands:

```
#for zostera, centers of quadra is at 15,10  
edge_effect(zostera, coords = coord_zostera, vecpop = popvec,  
            center = rep(c(15,10),2))
```

or, with MLL:

```
edge_effect(popsim, coords = coord_sim, center = rep(c(15,10),2), listMLL = MLLlist)
```

or, for haploid data:

```
edge_effect(haplodata, coords = coord_haplo, center = rep(c(15,10),2))
```

Options:

```
edge_effect(zostera, coords = coord_zostera, vecpop = popvec, center = rep(c(15,10),2),  
            nbrepeat = 100) #pvalue computation  
edge_effect(zostera, coords = coord_zostera, vecpop = popvec, center = rep(c(15,10),2),  
            nbrepeat = 1000, bar = TRUE) #could be time consuming
```

Results:

```
res <- edge_effect(zostera, coords = coord_zostera, vecpop = popvec,  
                  center = rep(c(15,10),2), nbrepeat = 100) #better put 1000 nbrepeat at least
```

```
res$SaintMalo$results #Aggregation index
```

| Ee | pval_Ee | nbrepeat |
|-----------|---------|----------|
| 0.0288465 | 0.798 | 1000 |

```
#res$SaintMalo$simulation #vector of sim aggregation index
```

H. BONUS

Summary function:

Basic commands:

```
genclone(zostera, coords = coord_zostera, vecpop = popvec)
```

or, with MLL:

```
genclone(popsim, coords = coord_sim, listMLL = MLLlist)
```

or, for haploid data:

```
genclone(haplodata, haploid = TRUE, coords = coord_haplo)
```

Options:

```
genclone(zostera, coords = coord_zostera, vecpop = popvec, nbrepeat = 100) #pvalues
genclone(zostera, coords = coord_zostera, vecpop = popvec, nbrepeat = 1000, bar = TRUE)
#could be time consuming
```

Results:

```
genclone(zostera, coords = coord_zostera, vecpop = popvec)
```

| | N | Lineage | nb_L | nb_all | SE | Fis | pval_2sides | Fis_WR | pval_2sides.1 | R |
|-----------|----|---------|------|----------|-----------|-----|-------------|--------|---------------|-----------|
| SaintMalo | 29 | MLG | 17 | 2.777778 | 0.6620208 | NaN | NA | NaN | NA | 0.5714286 |
| Arcouest | 30 | MLG | 16 | 3.111111 | 0.6111111 | NaN | NA | NaN | NA | 0.5172414 |

| | Pareto_index | Sp_Loiselle | pval_2sides | Sp_L_WR | pval_2sides.1 | Sp_Ritland | pval_2sides.2 |
|-----------|--------------|-------------|-------------|-------------|---------------|-------------|---------------|
| SaintMalo | 1.088812 | 0.008698691 | NA | 0.00542055 | NA | 0.008193788 | NA |
| Arcouest | 0.3403204 | 0.007036951 | NA | 0.003786488 | NA | 0.003920472 | NA |

| | Sp_R_WR | pval_2sides | H'' | J' | D | V | Hill |
|-----------|-------------|-------------|----------|-----------|-----------|-----------|----------|
| SaintMalo | 0.004776727 | NA | 2.671294 | 0.9428497 | 0.9507389 | 0.8559028 | 20.3 |
| Arcouest | 0.001466866 | NA | 2.268605 | 0.8182264 | 0.8413793 | 0.3918367 | 6.304348 |

Arcouest and SaintMalo have *NaN* values for Fis and Fis_WR because some loci are homozygous.