

Integration of new Methods

PopGenome

Bastian Pfeifer

November 29, 2012

Accessing the class GENOME

The whole framework of PopGenome is based on a single class named **GENOME**. This object contains a huge amount of informations about the observed data and also stores the calculated statistic values. To ensure a whole genome perspective we use the ff-package (<http://cran.r-project.org/web/packages/ff/index.html>) to minimize memory space. Anyway, PopGenome provides an effortless access of the values stored in the class **GENOME**, what facilitates the integration of new methods and the reimplementation of existing methods become redundant. In the next sections we will discuss the integration of new methods on the basis of alignments. Note, this will also work for regions of whole genome SNP data.

Lets implement the Watterson's homozygosity test of neutrality:

$$H = \sum_{i=1}^k x_i^2$$

where k is the total number of haplotypes and x_i the frequency of the i -th haplotype with:

$$\sum_{i=1}^k x_i = 1$$

(1) Reading the data

In this case we read three alignments stored in the folder "Alignments".

```
GENOME.class <- readData("Alignments")
```

(2) Calculation

The FST module calculates the haplotype counts needed for the new statistic.

```
GENOME.class <- F_ST.stats(GENOME.class)
# A faster version would be:
GENOME.class <- F_ST.stats(GENOME.class, mode="haplotype",
                           only.haplotype.counts=TRUE)
```

(3) Getting the results

The haplotype counts of each alignment or region are stored in the sub-class `region.stats`.

```
haplotype.counts <- GENOME.class@region.stats@haplotype.counts
haplotype.counts
[[1]]
    CON RI-0 MR-0 TUL-0 MH-0 ITA-0 CVI-0 COL-2 LA-0 ME-0 GR-5 CHA-0 WS-0
pop.1    4     1     1     2     1     1     1     1     3     1     1     1     1
                    RSCH-0 Alyrata
pop.1        1         1

[[2]] ...
[[3]] ...
```

(4) Writing your own function

```
EW_Test <- function(GENOME.class){
  GENOME.class      <- F_ST.stats(GENOME.class,only.haplotype.counts=TRUE)
  haplotype.counts <- GENOME.class@region.stats@haplotype.counts
  frequencies      <- lapply(haplotype.counts,function(x){return((x/sum(x))^2)})
  EW_values        <- sapply(frequencies,sum)
  return(EW_values)
}

EW_Test(GENOME.class)
[1] 0.09297052 0.18367347 0.07482993
```

Embedding new methods into the PopGenome framework

PopGenome provides a mechanism to fully integrate your own functions into the PopGenome framework. The next subsections will guide you through this mechanism. Let's integrate the Ewens Watterson Test.

(1) Skeleton of a PopGenome function

Use the function `create.PopGenome.method` to generate the new function.

```
# one value for one population
create.PopGenome.method("myFunction", population.specific=TRUE)
# one value for all populations
create.PopGenome.method("myFunction", population.specific=FALSE)
# site specific values
create.PopGenome.method("myFunction", site.specific=TRUE)
```

Now you find the R script `myFunction.R` in your workspace. The script itself describes where to put your function.

```

103      if(NEWPOP, length(popmissing) := 0; respop <- c(1:nops, length(popmissing))
104      popmissing])}else{respop <- 1:nops}}
105
106 # END: DO NOT EDIT
107
108 # define here your own function in the PopGenome framework.
109 # default: bial (biallelic matrix), populations (the defined populations)
110 # ... choose here everthing you want from the GENOME class
111
112 new.value[xx,respop]           <- your_intern_function(bial,populations,...)
113
114
115
116
117 }
118 }
119
120 return(new.value)
121 }

```

Figure 1: *myFunction.R* (population specific)

(2) Writing your own function

Lets fully integrate the Ewens Watterson test described above in the PopGenome framework without accessing the slot `region.stats@haplotype.counts`. This can be neccessary if PopGenome for example would not calculate the haplotype counts. Following parameter are useful:

bial

The parameter `bial` (Biallelic Matrix) contains the SNPs of the alignments. The rownames are the individuals and the columns the positions of the observed SNPs.(Manual:`popGetBial`)

```
bial[1:5,1:5]
```

	12	13	31	44	59
CON	0	1	0	1	0
KAS-1	0	0	0	1	1
RUB-1	1	0	1	1	0
PER-1	0	0	0	0	0
RI-0	0	1	0	0	0

The Biallelic Matrix contains zeros (*major allele*) and ones (*minor allele*) regarding the whole population considered. Because of that the third SNP (position: 44) for example contains 3 minor alleles and 2 major alleles. PopGenome will manage this automatically and will redefine those values for every subset. Nevertheless, you should keep that in mind when yo write your own functions.

populations

The R object `populations` contains the defined populations as rownumbers of the Biallelic Matrix. You should really use this parameter, because sometimes

```

103      if(NEWPOP, length(popmissing) == 1, respop <- 1:nops, popmissing)
104  if(!NEWPOP) {if(length(object@region.data@popmissing[[xx]]) != 0){popmiss-
  popmissing}}else{respop <- 1:nops}
105
106 # END: DO NOT EDIT
107
108 # define here your own function in the PopGenome framework.
109 # default: bial (biallelic matrix), populations (the defined populations)
110 # ... choose here everthing you want from the GENOME class
111
112 new.value[xx,respop]           <- EW_Test(bial,populations)
113
114
115
116
117 }
118 }
119
120 return(new.value)
121 }

```

Figure 2: *myFunction.R* (population specific)

there are not all individuals present in an alignment. (`region.data@populations`)

```
populations[[2]][1:5]
```

```
[1] 1 2 3 4 5
```

In this case the first five individuals of the second ([[2]]) alignment are present.

Implementation

```

EW_test <- function(bial,populations){

  # Lets create the subsets of the Biallelic Matrix
  pop.bial      <- lapply(populations,function(x){return(bial[,x])})
  # Calculate the haplotype counts without accessing the class GENOME
  # calc.haplotype.counts is your own sub-function
  haplotype.counts <- lapply(pop.bial,calc.haplotype.counts)
  frequencies     <- lapply(haplotype.counts,function(x){return((x/sum(x))^2)})
  EW_values       <- sapply(frequencies,sum)

  return(EW_values)
}

```

Loading/Using the function

```

library(PopGenome)
GENOME.class <- readData("Alignments")
create.PopGenome.method("Ewens.Watterson")
# editing
source("Ewens.Watterson.R")
EW_values <- Ewens.Watterson(GENOME.class)

```

Now, you have the whole capability of PopGenome for your function.

1 Parser for new input formats

PopGenome supports also R-objects as an input. Thus you can parse every text file you want and convert it into a special R-object. This R-object is type of a list and contains a matrix including the nucleotides and the individuals as well as the positions of the sites of the matrix. The positions are optional, but might be useful in case of SNP data. The nucleotides are coded as follows:

- $T, U \rightarrow 1$
- $C \rightarrow 2$
- $G \rightarrow 3$
- $A \rightarrow 4$
- $unknown \rightarrow 5$
- $- \rightarrow 6$

Example

```
matrix
[,1] [,2] [,3] [,4]
seq1    1     1     1     2
seq2    1     4     2     1
seq3    1     4     4     1
seq4    1     4     4     1

positions
[1] 25 300 1000 2500

Robject <- list(matrix=matrix, positions=positions)

save(file="Aln", Robject)

# put the the objects in a folder (for example: .../Alignments/Aln.RData)

GENOME.class <- readData("Alignments", format="RData")
```

You also can split huge data in chunks, PopGenome will concatenate them.