A short manual for TESS3: a program to estimate spatial population structure

(command-line and R wrapper)

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Please, print this reference manual only if it is necessary.

This short manual aims to help users to run TESS3 command-line engine and R wrapper function on Mac, Linux and Windows.

1 Description

Inference of spatial population structure are commonly perform with computer program based on intensive stochastic simulation. These methods do not scale with the dimension of the data sets generated from nextgeneration sequencing technologies. The computer program TESS3 has functionalities similar to the spatial bayesian clustering program TESS [?], but has run-times several order faster than those of TESS 2.3. The method is based on geographically constrained non-negative matrix factorization, and provides similar results that with TESS 2.3. In addition TESS3 computes an ancestral allele frequency differentiation statistic that can be used to perform selection scans.

2 Installation

The source code of TESS3 can be download on github: https://github.com/cayek/TESS3/archive/ master.zip. Then you just need to follow instructions on the github project page https://github.com/ cayek/TESS3.

3 Data format

input file 3.1

The TESS3 input file consists of a genotype file in the **geno** format and coordinate file in the **coord** format. **geno** (example.geno)

The geno format has one row for each SNP. Each row contains 1 character per individual: 0 means zero copies of the reference allele. 1 means one copy of the reference allele. 2 means two copies of the reference allele. 9 means missing data.

Below, an example of a geno file for n=3 individuals and L=4 loci.

```
112
010
091
```

coord (example.coord)

The coord format has one row for each individual. Each row contains the longitude latitude information of the individual.

Below, an example of a geno file for n=3 individuals and L=4 loci.

```
2.515455200203690111e+01 5.439077948590419709e+01
-8.429345306090160861e+00 4.019713388759510053e+01
1.351129055178800087e+01 5.585331731335860184e+01
```

Other formats for genotype data sets can be used thanks to the LEA R package [?]. Indeed, LEA enable to convert into geno format the following usual formats: ped, ancestrymap, vcf and lfmm.

3.2 output files

There are three main **output files**.

- The file with the extension the \mathbf{Q} contains individual admixture coefficients. It contains a matrix with n rows (the number of individuals) and K columns (the number of ancestral populations).
- The file with the extension G contains the ancestral genotypic frequencies. It contains a matrix with $n_a \times L$ lines (the number of alleles times the number of SNPs) and K columns (the number of ancestral populations). For a diploid SNP, the first line contains the ancestral frequencies for the number of allele equals to 0, the second line contains the ancestral frequencies for the number allele equals to 1, the third line contains the ancestral frequencies for the number of alleles equal to 2.
- The file with the extension **.Fst** contains ancestral allele frequency differentiation statistic. In this file each row is the statistic estimate for each loci.

There also are less important **output** files, but that can be useful to have information on the graph computed, the least-squared criterion and the cross-entropy criterion.

- The file with the extension the .W contains the edge weight matrix of the nearest-neighbor graph computed from coordinates data.
- The file with the extension the .sum contains the value of least-squared criterion, the cross entropy criteria
 on all data and only on masked data if the user asked it.

4 Run the program

4.1 Command-line

The TESS3 program can be executed from a command line. The format is:

./TESS3 -x genotype_file.geno -K number_of_ancestral_populations -r coordinates_file

Only these three options are mandatory. There is no ordering for the options in the command line. Here is a description of these options :

- -x genotype_file.geno is the path to the genotype file (in .geno format).
- -K number_of_ancestral_populations is the number of ancestral populations.
- -r coordinates_file is the path to the coordinate file (in .coord format).

Additional options are available:

- -a alpha is the value of the normalized regularization parameter (by default : 0.001). This parameter control the spatial regularity of the ancestry estimates.
- -W edge_weight_input is the path to the file that contains a edge weight matrix of the spatial graph. The file has to contain each element of the matrix separates by a space row by row. The program use this graph in place of the one computed if no edge weight input file is given.
- -q output_Q is the path for the output file containing the ancestry coefficients. By default, the name of
 the output file is the same name as the input file with the extension .K.Q.
- -g output_G is the path to the output file containing the ancestral genotype frequencies. By default, the name of the output file is the same name as the input genotype file with the extension .K.G.
- -f output_FST is the path to the output file containing the ancestral allele frequency differentiation statistic. By default, the name of the output file is the same name as the input genotype file with the extension .K.Fst.
- -y output_FST is the path to the output file containing the value of least-squared criterion, the cross entropy criteria on all data and only on masked data. By default, the name of the output file is the same name as the input genotype file with the extension .K.sum.
- -c perc is the percentage of masked genotypes. If this option is set, the cross-entropy criterion is calculated (see [?] for more details on the cross-entropy criterion). The default percentage is 5%.
- -e tolerance is the tolerance error in the TESS3 optimization algorithm (by default: 0.0000001).
- -i iteration_number is the max number of iterations of the algorithm (default : 200).
- I nb_SNPs starts the algorithm with a run of TESS3 using a subset of nb_SNPs random SNPs. This
 option can speed up TESS3 estimation for very large data sets.
- -Q input_Q is the path to an initial file for the Q matrix containing individual admixture coefficients. If both -I and -Q are set, -Q is chosen.
- -s seed is a seed to initialize the random number generator.
- -m ploidy 1 if haploid, 2 if diploid (default : 2).
- -p p is the number of CPUs to use when the algorithm is run on a multiprocessor system. Be aware that the number of processes has to be lower or equal to the number of CPU units available on your computer (default : 1).

If you need a summary of options, you can use the -h option by typing the following command

```
./TESS3 -h
```

4.2 R wrapper

The TESS3 program can be executed using a wrapper in R software environment. The wrapper and helper functions are defined in the R script src/Rwrapper/TESS3.R, the user can directly source this script in a R session. We now present function define in this script:

- TESS3

Description The wrapper function that call the command-line program. This function create a directory TESS3_workingDirectory to store input and output file.

Usage

Arguments

- genotype : genotype R matrix of size n individual by L loci or the .geno format file name.
- spatialData: coordinate R matrix of size n individual by 2 or the .coord format file name.
- K: vector of number of ancestral cluster. The TESS3 program is run for each element of this vector.
- ploidy: 1 if haploid, 2 if diploid.
- rep: number of run for each number of ancestral population.
- maskedProportion : if maskedProportion > 0, the cross-entropy criterion is calculated for this percentage of masked genotypes.
- alpha: value of the normalized regularization parameter. This parameter control the spatial regularity
 of the ancestry estimates.
- Getter

Description Functions useful to fetch results of TESS3 R function.

Usage

```
getQ( project, K, run = "best" )
getG( project, K, run = "best" )
getFst( project, K, run = "best" )
getCrossEntropy( project, func = mean )
getLeastSquared( project, func = mean )
```

Arguments

- project : object returned by the TESS3 R function.
- K: number of ancestral cluster.
- run: number of the run, or "best" if you want the best result with respect to the least-squared criterion.
- func : function used to summarize data over all run.
- Reader

Description Function useful to read data from file.

```
Usage
read.coord( file )
```

Arguments

- file name of the file to read.

A full example in R is available at the end of this note.

5 Tutorial

5.1 Data set

The data set that we analyse in this tutorial is a simulated dataset from a Arabidopsis thaliana Data set used in [?]. We obtain a haploid data set of n = 170 individual and L = 26943 loci.

5.2 Example of analysis using TESS3

This tutorial example describe how to run TESS3 in R for different values of K the number of ancestral populations. Then we plot the cross entropy criterion with respect to K. Finally we use R script available on POPS website (see http://membres-timc.imag.fr/Olivier.Francois/pops.html) to plot the ancestry coefficient with the K found with the cross entropy criteria [?]. You need to change TESS3_directory and POPS_directory by the directory path of TESS3 and POPS.

```
source("TESS3_directory/src/Rwrapper/TESS3.R")
library(LEA)
setwd( "TESS3_directory/data/simulated/Athaliana" )
# Run TESS3 on a data set simualted from an Arabidopsis Athalina data set #
#read data
spatialData = read.coord("Athaliana.coord")
n = nrow(spatialData)
project = TESS3( genotype = "Athaliana.geno",
            spatialData = "Athaliana.coord",
            K = 1:5,
            ploidy = 1,
            rep = 1,
            maskedProportion = 0.2)
# Chose of K with cross-entropy criterion #
plot(1:5,
    getCrossEntropy( project ),
    main = "Cross entropy",
    type="b",
    xlab = "K",
    ylab = "corss entropy" )
# Plot result on map for K = 3 #
# R script available on http://membres-timc.imag.fr/Olivier.Francois/pops.html
source("POPS_direction/R/POPSutilities.r")
asciiFile="/home/cayek/Projects/TESS3/data/simulated/Athaliana/down_etopo1.asc"
```

5.3 Example of genome-scan for selection using TESS3

This tutorial show an example in R of use of the ancestral allele frequency differentiation statistic computed by TESS3. We first run TESS3 with K=3 ancestral populations. Then using standard population genetic theory the F_{ST} -statistic is transformed into squared t-score and p-value are computed using a fisher distribution. The test inflation due to neutral population structure is corrected by recalibrating the t^2 -score to have uniform distribution under the null hypothesis. Finally, we control the false discovery rate using a Benjamini Hochberg procedure and plot the Manhattan plot. You need to change TESS3_directory by the directory path of TESS3.

```
source("TESS3_directory/src/Rwrapper/TESS3.R")
library(LEA)
setwd( "TESS3_directory/data/simulated/Athaliana" )
# Run TESS3 on a data set simualted from an Arabidopsis Athalina data set #
#read data
spatialData = read.coord("Athaliana.coord")
genotype = read.geno("Athaliana.geno")
n = nrow(spatialData)
project = TESS3( genotype = genotype,
              spatialData = spatialData,
              K = 3,
              ploidy = 1,
              rep = 5)
# Genome scan for selection #
####################################
#### Fst with TESS3
Fst = getFst( project, K = 3 )
Fst[Fst < 0.0] = 0.0
#### Convert Fst into t score
squared.t.scores = Fst*(n-2)/(1-Fst)
#### recalibrated p-values
gif = 25
adj.p.values = pf( squared.t.scores/gif , df1 = 2, df2 = n-3, lower = FALSE )
hist(adj.p.values,prob=TRUE)
#### Benjamini Hochberg procedure
alpha = 1e-10
L = length(adj.p.values)
# return a list of candidates with an expected FDR of alpha.
w = which(sort(adj.p.values) < alpha * (1:L) / L)</pre>
```

6 Contact

If you need assistance, do not hesitate to send us an email (kevin.caye@imag.fr or olivier.francois@imag.fr).