Package 'tess3r'

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Type Package
Title An R Package for Population Genetics Study
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Description This R package implements the TESS3 program and tools useful to plot program outputs.
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LazyData TRUE
LinkingTo Rcpp
Imports Rcpp, fields, graphics, stats, RColorBrewer
Suggests testthat, knitr
VignetteBuilder knitr
R topics documented:
coord
createGridFromAsciiRaster
getConstraintsFromAsciiRaster
tess3
tess3Project-class
tess3r
Index 8

coord

Input file for tess3

Description

Description of the coord format. The coord format can be used as an input format for coordinates matrices in the functions tess3.

Details

The geno format has one row for each individual. Each row contains the longitude and latitude coordinates of each individual.

Here is an example of a coordiantes matrix using the coord format with 3 individuals and 4 loci:

```
2.5154 5.4390
-8.4293 4.0197
1.3536 5.5852
```

Author(s)

Kevin Caye

createGridFromAsciiRaster

Create grid on which coefficients will be displayed

Description

 $\verb|createGridFromAsciiRaster| returns a grid to use to project clusters on a map. The grid is computed from an ascii raster file.$

Usage

```
createGridFromAsciiRaster(file)
```

Arguments

file

A character string containing a the path to the ascii raster file.

Value

 $\verb|createGridFromAsciiRaster| returns a matrix which represent the grid extract from the raster file.\\$

See Also

This function was made to be used with getConstraintsFromAsciiRaster and maps.

```
getConstraintsFromAsciiRaster
```

Create grid on which coefficients will be displayed

Description

getConstraintsFromAsciiRaster compute a constraint matrix acording to the cell_value_min and cell_value_max values.

Usage

```
getConstraintsFromAsciiRaster(file, cell_value_min = NULL,
    cell_value_max = NULL)
```

Arguments

```
file A character string containing a the path to the ascii raster file.

cell_value_min

All case values inferior to this numeric value will be set to FALSE on the grid.

cell_value_max
```

All case values superior to this numeric value will be set to FALSE on the grid.

Value

 ${\tt getConstraintsFromAsciiRaster} \ \ \textbf{returns} \ \ \textbf{a} \ \ \textbf{matrix} \ \ \textbf{with boolean cells}. \ \ \textbf{Cells is set to} \\ \textbf{TRUE if user wants to project on this map's cell}$

See Also

This function was made to be used with createGridFromAsciiRaster and maps.

maps

Display maps of membership/admixture coefficients.

Description

maps projects membership/admixture coefficients on a grid using Krig. Gradients in coefficients are represented by gradients in colors. if onemap == T & method == "treshold" only coefficients > 0.5 are plotted. if onemap == T & method == "max" at each point the cluster for which the coefficient is maximal is plotted (even if the value is less than 0.5). if onemap == F all values are plotted (since there is one cluster represented on each map, there is no overlap problem).

Usage

```
maps(matrix, coord, grid, constraints = NULL, method = "treshold",
    colorGradientsList = lColorGradients, onemap = T, onepage = T, ...)
```

4 maps

Arguments

See Also

This function was made to used with createGridFromAsciiRaster, maps and tess3

Examples

```
## How maps can be used to plot tess3 function output ##
###################
# runs of tess3 #
###################
# Retrieve data file name
genotype.file <- system.file("extdata/Athaliana", "Athaliana.geno", package = "tess3r")</pre>
coord.file <- system.file("extdata/Athaliana","Athaliana.coord",package = "tess3r")</pre>
# Read coordinate file
coord <- read.coord(coord.file)</pre>
n <- nrow(coord)</pre>
project <- tess3(input.file = genotype.file,</pre>
                 input.coord = coord.file,
                  K = 1:5,
                  ploidy = 1,
                  repetitions = 1,
                  entropy = TRUE,
                  percentage = 0.2,
                  project = "new")
#####################
# Plot result on map#
#####################
asciiFile=system.file("extdata/","lowResEurope.asc",package = "tess3r")
grid=createGridFromAsciiRaster(asciiFile)
# To display only altitudes above 0:
constraints=getConstraintsFromAsciiRaster(asciiFile,cell_value_min=0)
maps (matrix = Q( project, K = 3, run = 1),
     coord = coord,
     grid=grid,constraints=constraints,method="max",main="ancestry coefficient with K = 3
```

tess3 5

tess3	Estimates individual ancestry coefficients, ancestral allele frequencies and an ancestral allele frequency differentiation statistic.

Description

tess3 estimates admixture coefficients using a graph based Non-Negative Matrix Factorization algorithms, and provide STRUCTURE-like outputs. tess3 also computes an ancestral allele frequency differentiation statistic

Usage

```
tess3(input.file, input.coord, K, project = "continue", repetitions = 1,
   alpha = 0.001, tolerance = 1e-05, entropy = FALSE, percentage = 0.05,
   I = 0, iterations = 200, ploidy = 1, seed = -1, CPU = 1,
   Q.input.file = "")

## S4 method for signature 'tess3Project, ANY'
plot(x, y, ...)
```

Arguments

ploidy

guments	
input.file	A character string containing a the path to the input genotype file, a genotypic matrix in the geno format.
input.coord	A character string containing a the path to the input coordinate file, a coordinate matrix in the coord format.
K	An integer vector corresponding to the number of ancestral populations for which the tess3 algorithm estimates have to be calculated.
project	A character string among "continue", "new", and "force". If "continue", the results are stored in the current project. If "new", the current project is removed and a new one is created to store the result. If "force", the results are stored in the current project even if the input file has been modified since the creation of the project.
repetitions	An integer corresponding with the number of repetitions for each value of K.
alpha	A numeric value corresponding to the tess3 regularization parameter. The results depend on the value of this parameter.
tolerance	A numeric value for the tolerance error.
entropy	A boolean value. If true, the cross-entropy criterion is calculated.
percentage	A numeric value between 0 and 1 containing the percentage of masked genotypes when computing the cross-entropy criterion. This option applies only if entropy == TRUE.
I	The number of SNPs to initialize the algorithm. It starts the algorithm with a run of snmf using a subset of nb.SNPs random SNPs. If this option is set with nb.SNPs, the number of randomly chosen SNPs is the minimum between 10000 and 10 % of all SNPs. This option can considerably speeds up snmf estimation for very large data sets.
iterations	An integer for the maximum number of iterations in algorithm.

1 if haploid, 2 if diploid, n if n-ploid.

6 tess3

seed A seed to initialize the random number generator. By default, the seed is ran-

domly chosen.

CPU A number of CPUs to run the parallel version of the algorithm. By default, the

number of CPUs is 1.

Q.input.file A character string containing a path to an initialization file for Q, the individual

admixture coefficient matrix.

Value

tess3 returns an object of class tess3Project. The following methods can be applied to the object of class tess3Project:

 ${\tt plot} \qquad \qquad {\tt Plot \ the \ minimal \ cross-entropy \ in \ function \ of \ K.}$

show Display information about the analyses.

summary Summarize the analyses.

Q Return the admixture coefficient matrix for the chosen run with K ancestral pop-

ulations.

G Return the ancestral allele frequency matrix for the chosen run with K ancestral

populations.

FST Return ancestral allele frequency differentiation statistic matrix for the chosen

run with K ancestral populations.

cross.entropy

Return the cross-entropy criterion for the chosen runs with K ancestral popula-

tions.

load.snmfProject(file.tess3Project)

Load the file containing an tess3Project objet and return the tess3Project object.

remove.snmfProject(file.tess3Project)

 $\label{lem:condition:condition:all the files associated with the } Erase \ a \ \text{tess3Project object.} \ Caution: \ All \ the \ files \ associated \ with \ the$

object will be removed.

Examples

tess3Project-class 7

```
# get the cross-entropy of each run for K = 4
ce = cross.entropy(project, K = 4)

# select the run with the lowest cross-entropy
best = which.min(ce)

# plot the best run for K = 3 (ancestry coefficients).
barplot(t(Q(project, K = 3, run = best)), col = c(2:4))
```

tess3Project-class An S4 class to represent a tess3 project.

Description

An S4 class to represent a tess3 project.

See Also

You can see how to use method of this class in tess3

tess3r

tess3r: An R Package for Population Genetics Study

Description

This R package implements the TESS3 program and tools useful to plot program outputs.

Index

```
*Topic format
   coord, 2
coord, 2, 5
createGridFromAsciiRaster, 2, 3, 4
cross.entropy (tess3), 5
example_coord(coord), 2
FST (tess3), 5
G(tess3), 5
geno, 5
getConstraintsFromAsciiRaster, 2,
Krig, 3
load.snmfProject (tess3), 5
maps, 2, 3, 3, 4
plot, tess3Project, ANY-method
       (tess3), 5
Q(tess3), 5
remove.snmfProject(tess3),5
tess3, 2, 4, 5, 5, 7
tess3Project-class,7
tess3r,7
tess3r-package (tess3r), 7
```