

Package ‘rmetasim’

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Title An individual-based population genetic simulation environment

License GPL

Description An interface between R and the metasim simulation engine. Facilitates the use of the metasim engine to build and run individual based population genetics simulations. The simulation environment is documented in: Allan Strand. Metasim 1.0: an individual-based environment for simulating population genetics of complex population dynamics. Mol. Ecol. Notes, 2:373-376, 2002. (Please contact Allan Strand with comments, bug reports, etc). This version represents a significant alteration of function names that hopefully increases consistency and reduces the chances of collisions with other packages naming conventions. For a spatially-explicit, but slower, package with a similar interface, see kernelPop

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is.landscape	<i>Test whether an object is a (fairly) legitimate landscape</i>
--------------	--

Description

Test whether a genuine landscape

Usage

```
is.landscape(Rland = NULL, verb = TRUE, exact = FALSE)
```

Arguments

Rland	the Rmetasim landscape object
verb	print why not a landscape
exact	more strict

Examples

```
exampleland <- landscape.new.example()
is.landscape(exampleland)
rm(exampleland)
```

```
landscape.allelecount
```

Calculate allele numbers (frequency in the statistical sense) at each locus in each population

Description

Calculate allele counts

Usage

```
hetmat <- landscape.exp.het(rland, tbl.out=F)
```

Arguments

rland	the Rmetasim landscape object
tbl.out	return as a (three-dimensional) table if TRUE. If FALSE, return as a dataframe with categorical variables denoting the locus, population and allele.

Value

Depends on the value of tbl.out. See above.

See Also

landscape.allelefreq, landscape.obs.het, landscape.exp.het, landscape.Fwright, landscape.Fst

Examples

```
exampleland <- landscape.new.example()
exampleland <- landscape.simulate(exampleland, 4)
landscape.allelefreq(exampleland, tbl.out=TRUE)
landscape.allelefreq(exampleland, tbl.out=FALSE)
rm(exampleland)
```

```
landscape.allelefreq
```

Calculate allele frequencies at each locus in each population

Description

Calculate allele frequencies

Usage

```
hetmat <- landscape.allelefreq(rland, tbl.out=F)
```

Arguments

<code>rland</code>	the Rmetasim landscape object
<code>tbl.out</code>	return as a (three-dimensional) table if TRUE. If FALSE, return as a dataframe with categorical variables denoting the locus, population and allele.

Value

Depends on the value of `tbl.out`. See above.

See Also

`landscape.obs.het`, `landscape.exp.het`, `landscape.Fwright`, `landscape.Fst`

Examples

```
exampleland <- landscape.new.example()
exampleland <- landscape.simulate(exampleland, 4)
landscape.allelefreq(exampleland, tbl.out=TRUE)
landscape.allelefreq(exampleland, tbl.out=FALSE)
rm(exampleland)
```

```
landscape.amova
```

calculates phi-st for every locus in the landscape

Description

calculates ϕ_{ST} for every locus in the landscape

Usage

```
landscape.amova(rland, np = 24, ns = 24)
```

Arguments

rland	landscape object
np	max number of pops to include
ns	max number of samples to collect

Value

vector of length equal to the number of loci

See Also

[landscape.amova.locus](#), [landscape.amova.pairwise](#)

landscape.amova.locus

uses functions in ade4 to calculate phi-st for a particular locus

Description

Runs an amova on a locus. Does not include information about sequence similarity or ssr size in analysis.

Usage

```
landscape.amova.locus(l = 1, rland)
```

Arguments

l	locus number
rland	landscape object

Details

Should be the same as Weir and Cockerham's θ

Value

list of amova results for a locus

See Also

[landscape.amova](#), [landscape.amova.pairwise](#)

```
landscape.amova.pairwise  
calculates pairwise phi-ST for a landscape
```

Description

pairwise ϕ_{ST} calculator. Kind of slow. use `landscape.sample`

Usage

```
landscape.amova.pairwise(rland)
```

Arguments

rland landscape object

See Also

landscape.amova, landscape.amova.locus

```
landscape.clean    Function to resolve inconsistencies within a landscape
```

Description

Converts a landscape to internal format and back. This can resolve inconsistencies in a 'hand-built' landscape

Usage

```
rland <- landscape.clean(rland)
```

Arguments

rland the Rmetasim landscape object

Examples

```
exampleland <- landscape.new.example()  
exampleland <- landscape.simulate(exampleland, 4)  
exampleland.clean <- landscape.clean(exampleland)  
rm(exampleland)
```

 landscape.coalinput

Add loci and individuals based upon output from SimCoal 2.0

Description

Take rmetasim object and replaces the locus and individual data based on the results of a SimCoal run stored in Arlequin format files

Usage

```
## must be called AFTER integer, switch, and float params have been created
rland <- landscape.coalinput(rland, npp=500, arlseq = "seq.arp", arlms = "ms.arp",
                             msmut = 5e-4, mut.rates = NULL)
```

Arguments

rland	partially created landscape object, required
npp	number per population. Scalar or vector of length equal to number of populations. If scalar, value replicated
arlseq	name of the Arlequin format file containing a single locus of haploid sequence data for any number of populations
arlms	name of the Arlequin format file containing a single locus of diploid microsatellite data for any number of populations
seqsitemut	mutation rate for sequence data
msmut	mutation rate for diploid genotypic data
mut.rates	alternative means to specify mutation rates. Legal values are either NULL or a vector of rates equal to the number of loci to simulate. If NULL, SSR loci are assigned msmut as a mutation rate and sequence-based loci, seqsitemut. If a vector, overrides msmut and seqsitemut

Details

This function provides part of an interface between R and SimCoal, an environment for simulating sequences and microsatellite genotypes from coalescent trees. SimCoal can be used to simulate a standing crop of alleles and their relationships under a wide range of demographies. It returns haplotypes and genotypes of individuals in Arlequin format files.

If either 'arlseq' or 'arlms' are set to NULL, their corresponding data will not be included in the landscape (for example if arlseq=NULL, only diploid genotypes will be imported)

The genotypes in the Arlequin files are used to create rland\$loci objects based upon their frequencies and states. These rland\$loci sub-objects are then used to populate the rland\$individuals sub-object.

The number of populations in the Arlequin files should be the same among genetic locus types (sequence versus microsatellite) and the rland\$intparam\$habitats parameter. The per-population frequency data will be used in creating individuals

Value

an rmetasim object with new loci and individuals

Author(s)

Mark Bravington and Allan Strand

Examples

```
exampleland <- landscape.new.empty()
exampleland <- landscape.new.intparam(exampleland, s=2, h=2)
exampleland <- landscape.new.floatparam(exampleland)
exampleland <- landscape.new.switchparam(exampleland)

# exampleland <- landscape.coalinput(exampleland)
# exampleland$loci
```

landscape.compress *Function to resolve inconsistencies within a landscape, deprecated*

Description

Deprecated, 'clean.landscape()' does the same.

Usage

```
rland <- landscape.compress(rland)
```

Arguments

rland the Rmetasim landscape object

Examples

```
exampleland <- landscape.new.example()
exampleland <- landscape.simulate(exampleland, 4)
exampleland.clean <- landscape.compress(exampleland)
rm(exampleland)
```

```
landscape.democol
```

return largest demographic column from a landscape

Description

return largest demographic column from a landscape

Usage

```
landscape.democol()
```

Details

Useful to write functions that will be insensitive to some changes in the individuals object (mainly addition of non-genetic information)

Value

a scalar integer representing the largest column of demographic information in a landscape's individuals object

See Also

landscape.locus

```
landscape.demography
```

Calculate demographic parameters

Description

Calculate demographic parameters from a landscape: CURRENTLY BROKEN!

Usage

```
rland <- landscape.demography(rland)
```

Arguments

`rland` the Rmetasim landscape object

Value

A list of length `populations+1`. The first `1..populations` elements are lists comprised of `lambda`, the equilibrium stage-structure, the actual stage structure, a χ^2 value for the test of difference between predicted and actual, and an estimate of significance for that test. The last element of the main list is the same as the previous ones except it refers to the entire landscape

landscape.exp.het *Calculate expected heterozygosity*

Description

Calculate expected heterozygosity from a landscape

Usage

```
hetmat <- landscape.exp.het(rland)
```

Arguments

rland the Rmetasim landscape object

Details

Calculates the expected heterozygosity in each population:

$$1 - \sum_{i_k} p_i^2$$

where p is a vector of allele frequencies for a locus in a population.

Value

A matrix with num loci columns and num populations rows. Each element reflects the expected heterozygosity for that population x locus combination

See Also

landscape.obs.het, Fst.landscape

Examples

```
exampleland <- landscape.new.example()
exampleland <- landscape.simulate(exampleland, 4)
expHet <- landscape.exp.het(exampleland)
rm(exampleland)
```

landscape.Fst	<i>Calculates population structure statistic for the entire landscape</i>
---------------	---

Description

Calculate Fst for each allele at each locus in the landscape. If verb is set to TRUE, the function prints average Fst for loci and overall.

Usage

```
Fstmat <- landscape.Fst(rland, verb=F)
```

Arguments

rland	the Rmetasim landscape object
verb	determines whether there is verbose output

Details

Calculates Fst based upon the ratio of variance in allele frequency across subpopulations to the total variance in that allele's frequency. Does not calculate Wright's other statistics.

Value

A matrix with num alleles columns and num loci rows. Each element reflects the value of Fst for that allelexlocus combination. NA is assigned to alleles that are not present at a locus (either no longer or ever)

See Also

obs.het.landscape, exp.het.landscape, FWright.landscape

Examples

```
exampleland <- landscape.new.example()
exampleland <- landscape.simulate(exampleland, 4)
Fst <- landscape.Fst(exampleland, verb=TRUE)
Fst
rm(exampleland, Fst)
```

landscape.locus *return a matrix containing genotypes for a particular locus*

Description

return a matrix containing genotypes for a particular locus

Usage

```
landscape.locus(lnum=1, Rland)
```

Arguments

lnum	the locus to return
Rland	the Rmetasim landscape object

Details

Returns a matrix with rows = `dim(rland$individuals)[1]`. The first three columns correspond to the class (and two placeholder variables) of an individual. Here rland is a landscape object. The remaining columns (1 if haploid, 2 if diploid) contain the allele indices for the various loci

Value

matrix

See Also

landscape.populations

Examples

```
exampleland <- landscape.new.example()
exampleland <- landscape.simulate(exampleland, 4)
print("Allele frequencies at locus 1")
table(landscape.locus(1, exampleland)[, c(-1:- (landscape.democol()))])
rm(exampleland)
```

```
landscape.locus.states
```

return a matrix containing actual allelic states and their indices

Description

Convenience function to return a matrix containing the states of the alleles and their indices for a particular locus

Usage

```
landscape.locus.states(lnum=1, Rland)
```

Arguments

lnum	the locus to return
Rland	the Rmetasim landscape object

Value

matrix

See Also

landscape.locus, landscape.states

```
landscape.locusvec
```

return a vector with the locus ids for each column in the individuals component of a landscape

Description

return a vector with the locus ids for each column in the individuals component of a landscape

Usage

```
landscape.locusvec(Rland)
```

Arguments

Rland	the Rmetasim landscape object
-------	-------------------------------

Value

vector

```
landscape.modify.epoch
      Modifies one of the landscape's epochs
```

Description

This function updates the demographic parameters in a landscape for a particular epoch

Usage

```
rland <- landscape.modify.epoch(rland, epoch=1, S=NULL, R=NULL, M=NULL, epochprob=NULL)
```

Arguments

rland	landscape object, required
epoch	the epoch to modify, default 1
S	(default=NULL) Survivability matrix for epoch, NULL leaves unchanged
R	(default=NULL) female Reproduction matrix for epoch, NULL leaves unchanged
M	(default=NULL) Male reproduction matrix for epoch, NULL leaves unchanged
epochprob	(default=NULL) probability of choosing this epoch, NULL leaves unchanged
startgen	(default=NULL) generation in which this epoch starts, NULL leaves unchanged
extinct	(default=NULL) vector of extinction probabilities per generation for each sub-population, NULL leaves unchanged
carry	(default=NULL) vector of carrying capacities for each subpopulation, must be <code>rland\$intparam\$habitats</code> in length, NULL leaves unchanged
localprob	(default=NULL) vector of probabilities for choosing local demographics, must be <code>length(rland\$demography\$localdem)</code> in length, NULL leaves unchanged

```
landscape.new.epoch
      Create an Epoch
```

Description

Create an epoch for a Rmetasim landscape object

Usage

```
## must be called AFTER integer, switch, and float params have
##   been created and after the demography has been created
## S, R, and M matrices must be square matrices of size X by X
##   where X = rland$intparam$stages*rland$intparam$habitats
```

```
rland <- landscape.new.epoch(rland, S=Smatrix, R=Rmatrix, M=Mmatrix, epochprob=1, startgen=1)
```



```

exampleland<- landscape.new.epoch(exampleland,exampleS,exampleR,exampleM)

exampleland$demography$epochs[[1]]

rm(exampleS)
rm(exampleR)
rm(exampleM)
rm(exampleland)

```

```

landscape.new.example
      Create a Default Landscape

```

Description

Create a Rmetasim landscape with all default parameters.

Usage

```
rland <- landscape.new.example()
```

Arguments

None

Examples

```

## Only usage
landscape.new.example()

```

```

landscape.new.floatparam
      Create a set of floating point parameters

```

Description

Create a set of floating point parameters for a Rmetasim landscape.

Usage

```

## must be called AFTER landscape.new.empty()
rland <- landscape.new.floatparam(rland,s=0)

```

Arguments

rland	skeleton of landscape object, required
s	selfing (default=0), the selfing rate of the species

Examples

```
## Defaults
exampleland <- landscape.new.empty()
exampleland <- landscape.new.floatparam(exampleland)
exampleland$floatparam

## .5 selfing rate
exampleland <- landscape.new.empty()
exampleland <- landscape.new.floatparam(exampleland, s=0.5)
exampleland$floatparam

rm(exampleland)
```

```
landscape.new.individuals
      Fill a landscape with individuals
```

Description

Create a set of individuals for a Rmetasim landscape object.

Usage

```
## must be called AFTER integer, switch, and float params, demography,
## epochs, and loci have been created

rland <- landscape.new.individuals(rland, PopulationSizes)
```

Arguments

```
rland          nearly complete landscape object, required
PopulationSizes
                vector of integers denoting how many individuals are in which stage and in
                which subpopulation, vector is ordered as: (pop1 stage1, pop1 stage2, ..., pop2
                stage1, pop2stage2, ...), must be of length  $rlandintparamhabitats * rlandintparamstages$ 
```

Examples

```
exampleS <- matrix(c(0.1, 0, 0.5, 0.3), nrow = 2)
exampleR <- matrix(c(0, 1.1, 0, 0), nrow = 2)
exampleM <- matrix(c(0, 0, 0, 1), nrow = 2)

exampleland <- landscape.new.empty()
exampleland <- landscape.new.intparam(exampleland, s=2, h=2)
exampleland <- landscape.new.floatparam(exampleland)
exampleland <- landscape.new.switchparam(exampleland)
exampleland <- landscape.new.local.demo(exampleland, exampleS, exampleR, exampleM)
```

```

## nonsense matrices
exampleS <- matrix(c(rep(0,4),
                    rep(1,4),
                    rep(0,4),
                    rep(1,4)), nrow = 4)
exampleR <- matrix(c(rep(0.5,4),
                    rep(0,4),
                    rep(0.5,4),
                    rep(0,4)), nrow = 4)
exampleM <- matrix(c(rep(0,4),
                    rep(.25,4),
                    rep(0,4),
                    rep(0,4)), nrow = 4)

exampleland<- landscape.new.epoch(exampleland,exampleS,exampleR,exampleM)
exampleland <- landscape.new.locus(exampleland,type=2,ploidy=2,mutationrate=.001,numallele
exampleland <- landscape.new.locus(exampleland,type=1,ploidy=1,mutationrate=.001,numallele
exampleland <- landscape.new.locus(exampleland,type=0,ploidy=2,mutationrate=.004,numallele

exampleland <- landscape.new.individuals(exampleland,
                                         c(5,20,7,15))

exampleland$individuals

rm(exampleS)
rm(exampleR)
rm(exampleM)
rm(exampleland)

```

landscape.new.intparam

Create a set of integer parameters

Description

Create a set of integer parameters for a Rmetasim landscape.

Usage

```

## must be called AFTER landscape.new.empty()
rland <- landscape.new.intparam(rland,h=2,s=1,cg=0,ce=0,totgen=500,maxland=10000)

```

Arguments

rland	skeleton of landscape object, required
h	habitats (default=1), the number of different subpopulations within the landscape
s	stages (default=1), the number of stages in the life cycle of the organism

cg	currentgen (default=0), the current generation the simulation has reached
ce	currentepoch (default=0), the current epoch the simulation has reached
totgen	totoalgens (default=1000), the total number of generations to simulate
maxland	maxlandsize(default=200000), the maxium number of individuals that can exist in the simulation

Examples

```
## Defaults
exampleland <- landscape.new.empty()
exampleland <- landscape.new.intparam(exampleland)
exampleland$intparam

## 2 habitats, 3 stage lifecycle, 1000000 generations, maximum 1000000 individuals
exampleland <- landscape.new.empty()
exampleland <- landscape.new.intparam(exampleland,h=2,s=2,totgen=1000000,maxland=1000000)
exampleland$intparam

rm(exampleland)
```

landscape.new.landscape

Create a Skeletal Landscape

Description

Create a skeletal Rmetasim landscape ready to be configured

Usage

```
rland <- landscape.new.empty()
```

Arguments

None

Examples

```
## Only usage
landscape.new.empty()
```

```
landscape.new.local.demo
      Create a Local Demography
```

Description

Create a local demography for an Rmetasim Landscape object

Usage

```
## must be called AFTER integer, switch, and float params have been created
## S, R, and M matrices must be square matrices of size rland$intparam$stages
rland <- landscape.new.local.demo(rland, S=Smatrix, R=Rmatrix, M=Mmatrix, k=0)
```

Arguments

rland	partially created landscape object, required
S	Survivability matrix for demography, required
R	female Reproduction matrix for demography, required
M	Male reproduction matrix for demography, required
k	flag for type of matrix, 0=demography at zero population density, 1=demography at carrying capacity

Details

The local demography objects encapsulate demography within a particular region. Multiple such objects can be defined to account for different demographies across space. The flag, k, can indicate whether the matrices represent demography at zero population growth and at carrying capacity, if density-dependence is modeled

Examples

```
exampleS <- matrix(c(0.1, 0, 0.5, 0.3), nrow = 2)
exampleR <- matrix(c(0, 1.1, 0, 0), nrow = 2)
exampleM <- matrix(c(0, 0, 0, 1), nrow = 2)

exampleland <- landscape.new.empty()
exampleland <- landscape.new.intparam(exampleland, s=2)
exampleland <- landscape.new.floatparam(exampleland)
exampleland <- landscape.new.switchparam(exampleland)
exampleland <- landscape.new.local.demo(exampleland, exampleS, exampleR, exampleM)

exampleland$demography$localdem

rm(exampleS)
rm(exampleR)
```

```
rm(exampleM)
rm(exampleland)
```

```
landscape.new.locus
```

Add a locus

Description

Add a locus to a Rmetasim landscape object

Usage

```
## must be called AFTER integer, switch, and float params have been created
rland <- landscape.new.locus(rland,type=0,ploidy=1,mutationrate=0,transmission=1,
```

Arguments

rland	partially created landscape object, required
type	(default=0) type of locus, 0=Infinite Allele mutation model (Integer), 1=Step-wise mutation model (Integer) state, 2=DNA base (variable length string state)
ploidy	(default=1) locus ploidy, 1 or 2
mutationrate	(default=0) probability of mutation per generation, less than or equal to 1
transmission	(default=1) 1=uniparental inheritance, 0=biparental inheritance
numalleles	(default=2) number of different alleles at the time of creation
allelesize	(default=50) length of DNA strings if type=2
frequencies	(default=NULL) vector of frequencies for each allele, must be numalleles long and add up to 1, if NULL frequencies are equally distributed

Examples

```
exampleland <- landscape.new.empty()
exampleland <- landscape.new.intparam(exampleland, s=2, h=2)
exampleland <- landscape.new.floatparam(exampleland)
exampleland <- landscape.new.switchparam(exampleland)

exampleland <- landscape.new.locus(exampleland,type=2,ploidy=2,mutationrate=.001,numalleles=2)

exampleland$loci

rm(exampleland)
```

```
landscape.new.switchparam
```

Create a set of boolean parameters

Description

Create a set of boolean (1 or 0) parameters for a Rmetasim landscape.

Usage

```
## must be called AFTER landscape.new.empty()
rland <- landscape.new.switchparam(rland, re=1, rd=1, mp=1, dd=0)
```

Arguments

rland	skeleton of landscape object, required
re	randepoch (default=0), 1=randomly pick a new epoch (from the epochs listed in the landscape) after an epoch completes, 0=epochs are chosen in order
rd	randdemo (default=0), 1=randomly choose a demography (from the demographies listed in the landscape) for each subpopulation, 0=demographies are assigned in order
mp	multp (default=1), 1=multiple paternity, 0=entire families from a single mating
dd	density dependence. If dd=1, then two of each local demography matrix must be defined, the first set using new.local.demo with k=0 and representing demography at low density and again with k=1 for demography at high population density.

Examples

```
## Defaults
exampleland <- landscape.new.empty()
exampleland <- landscape.new.switchparam(exampleland)
exampleland$switchparam

## Random epochs, random demographies, and no multiple paternity
exampleland <- landscape.new.empty()
exampleland <- landscape.new.switchparam(exampleland, re=1, rd=1, mp=0)
exampleland$switchparam

rm(exampleland)
```

`landscape.obs.het` *Calculate observed heterozygosity*

Description

Calculate observed heterozygosity from a landscape

Usage

```
hetmat <- landscape.obs.het(rland)
```

Arguments

`rland` the Rmetasim landscape object

Value

A matrix with num loci columns and num populations rows. Each element reflects the observed heterozygosity for that population x locus combination

See Also

`landscape.exp.het`, `landscape.Fst`

Examples

```
exampleland <- landscape.new.example()
exampleland <- landscape.simulate(exampleland, 4)
obshet <- landscape.obs.het(exampleland)
rm(exampleland)
```

`landscape.ploidy` *return a vector with the ploidy of each locus*

Description

return a vector with the ploidy of each locus in the order they appear in the landscape

Usage

```
landscape.ploidy(Rland)
```

Arguments

`Rland` the Rmetasim landscape object

Value

vector

See Also

landscape.populations

Examples

```
exampleland <- landscape.new.example()
landscape.ploidy(exampleland)
rm(exampleland)
```

landscape.populations

return a vector of population IDs from a landscape

Description

return a vector of population IDs from a landscape

Usage

```
landscape.populations(Rland)
```

Arguments

Rland the Rmetasim landscape object

Details

Returns a vector of length `dim(rland$individuals)[1]` where `rland` is a landscape object. The vector classifies individuals into populations (or habitats)

Value

a vector

See Also

landscape.locus, landscape.ploidy

Examples

```
exampleland <- landscape.new.example()
exampleland <- landscape.simulate(exampleland, 4)
plot(table(landscape.populations(exampleland)), main="Distribution of population size in la
rm(exampleland)
```

landscape.read *Load a landscape from a file*

Description

Load a Rmetasim landscape from a file into an R object.

Usage

```
rland <- landscape.read(fn="mylandscape.lnd")
```

Arguments

fn the path and name of the file containing the landscape

Examples

```
## Needs write access to the current directory, files created!!
landscape.write(landscape.new.example(), "exampleland.lnd")
exampleland <- landscape.read("exampleland.lnd")
exampleland
rm(exampleland)
```

landscape.sample *simulates sampling for genetics on the landscape*

Description

Randomly pulls a max of ns individuals from a max of np populations and returns a landscape object that could be used for further simulation, but is usually used for analyses and summary statistics calculatiuons. If one needs a sample of specific populations/habitats, then these should be given in the vector pvec

Usage

```
landscape.sample(rland, np = NULL, ns = NULL, pvec = NULL)
```

Arguments

rland landscape object
np number populations
ns number samples per population
pvec vector of population id numbers to sample

Value

landscape object

Examples

```
l <- landscape.new.example()
l <- landscape.simulate(l,1)
l.samp <- landscape.sample(l,np=3,ns=24)
landscape.amova.pairwise(l.samp)
l.samp2 <- landscape.sample(l,ns=24,pvec=c(1,3))
landscape.amova.pairwise(l.samp2)
```

landscape.simulate *Run a simulation for a single landscape through time*

Description

Simulate a Rmetasim landscape for a number of generations.

Usage

```
rland <- landscape.simulate(rland=l,numit=100,seed=-1,compress=FALSE,adj.lambda=0)
```

Arguments

rland	the Rmetasim landscape object
numit	the number of generations/iterations to simulate, note that landscapes will not run past the <code>rland\$ntparam\$totalgens</code> value
seed	The default value of seed uses the seed set in the calling environment. Any other value for seed uses 'set.seed()' to reset the random number generator. landscape.simulate uses the RNG selected by the calling environment.
compress	If true, landscape.simulate executes a survival and carrying capacity step before returning. In demographies with high reproductive potential, this can significantly reduce the size of R objects returned
adj.lambda	Tries to apply a correction to population growth that makes the observed growth rate more closely approximate that predicted from standard analysis eigensystem of the sum of the survival and reproduction Lefkovitch matrices

Examples

```
exampleland <- landscape.new.example()
exampleland <- landscape.simulate(exampleland, 4)
exampleland
rm(exampleland)
```

landscape.states *return a matrix containing actual genotypes for a particular locus*

Description

return a matrix containing the states of the alleles in genotypes for a particular locus

Usage

```
landscape.states(lnum=1, Rland)
```

Arguments

lnum	the locus to return
Rland	the Rmetasim landscape object

Details

Returns a matrix with `rows = dim(rland$individuals)[1]`. The columns 1:landscape.democol() correspond to demographic variables for an individual. The columns are: state, placeholder, birthyear, id, mother's id, and father's id. Here rland is a landscape object. The remaining columns (1 if haploid, 2 if diploid) contain the states of the alleles for the selected loci

Value

matrix

See Also

landscape.locus

Examples

```
exampleland <- landscape.new.example()
exampleland <- landscape.simulate(exampleland, 4)
print("Allele frequencies at locus 1")
table(landscape.states(1, exampleland)[, c(-1:-landscape.democol())])
rm(exampleland)
```

landscape.theta.h *Calculate theta using heterozygosity*

Description

Calculate theta from a landscape based upon heterozygosity.

Usage

```
theta.h.mat <- landscape.theta.h(rland)
```

Arguments

rland the Rmetasim landscape object

Details

Uses routines in the package 'ape'

Value

A matrix with num loci columns and num populations rows. Each element reflects the estimated theta for that population x locus combination

See Also

landscape.theta.k, landscape.theta.s

Examples

```
exampleland <- landscape.new.example()
exampleland <- landscape.simulate(exampleland, 4)

theta.h.mat <- landscape.theta.h(exampleland)
theta.h.mat
```

landscape.theta.k *Calculate theta using the number of alleles*

Description

Calculate theta using number of alleles from a landscape.

Usage

```
theta.k.mat <- landscape.theta.k(rland)
```

Arguments

rland the Rmetasim landscape object

Details

Uses routines in the package 'ape'

Value

A matrix with num loci columns and num populations rows. Each element reflects the estimated theta for that population x locus combination

See Also

landscape.theta.h, landscape.theta.s

Examples

```
exampleland <- landscape.new.example()
exampleland <- landscape.simulate(exampleland, 4)

theta.k.mat <- landscape.theta.k(exampleland)
theta.k.mat
```

landscape.theta.s *Calculate theta using segregating sites*

Description

Calculate theta from a landscape based upon the number of segregating sites.

Usage

```
theta.s.mat <- landscape.theta.s(rland)
```

Arguments

rland the Rmetasim landscape object

Details

Uses routines in the package 'ape'

Value

A matrix with num loci columns and num populations rows. Each element reflects the estimated theta for that population x locus combination

See Also

theta.k.landscape, theta.h.landscape

Examples

```
exampleland <- landscape.new.example()  
exampleland <- landscape.simulate(exampleland, 4)  
theta.s.mat <- landscape.theta.s(exampleland)  
theta.s.mat
```

landscape.to.rtheta

Converts genetic marker data in a landscape into a format suitable for analysis using ANOVA

Description

This function converts Rmetasim landscapes into a format that can be analyzed with ANOVA to calculate Weir's theta.

Usage

```
genin <- landscape.to.rtheta(Rland, numi)
```

Arguments

Rland the Rmetasim landscape object
numi number of individuals to sample at random from each population/habitat. The default value of 0 takes all individuals in each population. CAUTION this could take a long time.

Value

A matrix with columns representing the population, class, individual, locus, allelic position, allele id.

See Also

popstruct

landscape.write *Save a landscape to a file*

Description

Save a Rmetasim landscape object to a file

Usage

```
rland <- landscape.write(rland=l, fn="mylandscape.lnd")
```

Arguments

rland	the Rmetasim landscape object
fn	the path and name of the file to save the landscape to

Examples

```
## Needs write access to the current directory, files created!!
exampleland <- landscape.new.example()
landscape.write(exampleland, "exampleland.lnd")

rm(exampleland)
```

landscape.write.foreign
 Save a landscape to a file in a foreign format

Description

Save a Rmetasim landscape object to a file in a suite of output formats

Usage

```
rland <- landscape.write.foreign(rland, fn="foreign", numi=24, fmt="GDA")
```

Arguments

rland	the Rmetasim landscape object
fn	the path and name of the file to save the landscape to
numi	number of individuals sampled per population for inclusion in subsequent analyses
fmt	the output format for the landscape: Can take the following values: "arlequin", "arlequinhap", "biosys", "gen

Examples

```
## Needs write access to the current directory, files created!!
exampleland <- landscape.new.example()
landscape.write.foreign(exampleland, fn="exampleland.nex", fmt="GDA")
rm(exampleland)
```

SimulationComponents

Code components to simulate a landscape

Description

These functions can be used to construct custom simulations of landscapes. Each conducts only a single generations worth of change

Usage

```
rland <- landscape.advance(rland=1)
rland <- landscape.carry(rland=1)
rland <- landscape.extinct(rland=1)
rland <- landscape.reproduce(rland=1)
rland <- landscape.survive(rland=1)
```

Arguments

rland the Rmetasim landscape object

Details

landscape.advance() merely advances the generation counter and selects the new generations demographic conditions if such conditions can vary. The other functions implement carrying capacity, local extinction, reproduction, and survival/growth, respectively. The function landscape.simulate() bundles the functionality of these components into a single function (and executes it slightly faster all within linked C++ code).

See Also

landscape.simulate

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