Package: selac (via r-universe)

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Description Sets up and executes a SelAC model (Selection on Amino acids and codons) for testing the presence of selection in amino acid or codon among a set of genes on a fixed phylogeny. Beaulieu et al (2019) <doi:10.1093 molbev="" msy222="">.</doi:10.1093>
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 ${\tt GetAdequateManyReps}$

Parallel model adequacy test

Description

Performs model adequacy test using multiple cores

Usage

Index

```
GetAdequateManyReps(nreps, n.cores, model.to.reconstruct.under = "selac",
  model.to.simulate.under = "gtr", selac.obj.to.reconstruct,
  selac.obj.to.simulate, aa.optim.input = NULL,
  fasta.rows.to.keep = NULL, taxon.to.drop = 2,
  partition.number = 17, numcode = 1, for.gtr.only = NULL)
```

Arguments

nreps Specifies the number of repeated model adequact simulations.

n.cores Specifies the number of cores you want to use.

model.to.reconstruct.under

Specifies the model that the internal nodes are to be reconstructed under assuming a single tip is pruned from the tree.

model.to.simulate.under

Specifies the model that the simulation will be conducted along the pruned tip.

selac.obj.to.reconstruct

The selac output object that contains the model parameters to be used in the reconstruction.

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selac.obj.to.simulate

The selac output object that contains the model parameters to be used in the simulation.

aa.optim.input A list of optimal amino acids with each list element designating a character vector for each gene. The optimal amino acids be the MLE from a selac run (default) or a list of user defined optimal A.A.

fasta.rows.to.keep

Indicates which rows to remove in the input fasta files.

Specifies the tip based on the number in the phy object to be removed and simtaxon.to.drop

ulated.

partition.number

Specifies the partition number to conduct the model adequacy test.

numcode The ncbi genetic code number for translation. By default the standard (num-

code=1) genetic code is used.

for.gtr.only A selac object that can be used as the reference optimal AA for when the ade-

quacy of a GTR+G model is tested only.

Details

Performs a parallelized analysis of the model adequacy test. The test prunes out a user-specified taxon from the tree, performs site data reconstruction for all nodes in the tree under a user-specified model, then simulates the expected data of the pruned taxon according to a user-specified model along uniformly sampled points along the branch. The functionality of the reconstructed sequence is also calculated along the way to see how functionality changes as the simulation reaches the end of the known branch length. The output is a list with elements equally the number of repititions. Each element contains the functionality of the simulated points along equally spaced sampling points along the known branch length (i.e., edge.length * seq(0, 1, by=0.05))

GetAdequateSelac

Model adequacy simulation

Description

Performs a single model adequacy simulation

```
GetAdequateSelac(model.to.reconstruct.under, model.to.simulate.under,
  selac.obj.to.reconstruct, selac.obj.to.simulate, aa.optim.input = NULL,
  fasta.rows.to.keep = NULL, taxon.to.drop = 4,
  partition.number = 55, numcode = 1, for.gtr.only = NULL)
```

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Arguments

model.to.reconstruct.under

Specifies the model that the internal nodes are to be reconstructed under assuming a single tip is pruned from the tree.

model.to.simulate.under

Specifies the model that the simulation will be conducted along the pruned tip.

selac.obj.to.reconstruct

The selac output object that contains the model parameters to be used in the reconstruction.

selac.obj.to.simulate

The selac output object that contains the model parameters to be used in the simulation.

aa.optim.input A list of optimal amino acids with each list element designating a character vector for each gene. The optimal amino acids be the MLE from a selac run (default) or a list of user defined optimal A.A.

fasta.rows.to.keep

Indicates which rows to remove in the input fasta files.

taxon.to.drop Specifies the tip based on the number in the phy object to be removed and sim-

ulated.

partition.number

Specifies the partition number to conduct the model adequacy test.

numcode The ncbi genetic code number for translation. By default the standard (num-

code=1) genetic code is used.

for .gtr.only A selac object that can be used as the reference optimal AA for when the ade-

quacy of a GTR+G model is tested only.

Details

Performs a single model adequacy simulation. The test prunes out a user-specified taxon from the tree, performs site data reconstruction for all nodes in the tree under a user-specified model, then simulates the expected data of the pruned taxon according to a user-specified model along uniformly sampled points along the branch. The functionality of the reconstructed sequence is also calculated along the way to see how functionality changes as the simulation reaches the end of the known branch length. The output is a vector with elements containing the functionality of the simulated points along equally spaced sampling points along the known branch length (i.e., edge.length * seq(0, 1, by=0.05))

GetFunctionality

Calculate functionality

Description

Calculates the functionality of a single gene

GetMarginalAllGenes 5

Usage

```
GetFunctionality(gene.length, aa.data, optimal.aa, alpha, beta, gamma,
   gp = NULL, aa.properties = NULL)
```

Arguments

gene.length Indicates the length of the gene used to calculate functionality.

aa.data A matrix of amino acids

optimal.aa A vector of inferred optimal amino acids.

alpha The inferred Grantham composition paramter
beta The inferred Grantham polarity parameter

gamma The inferred Grantham molecular volume parameter

gp A vector of gamma rates for calculating among site hetergeneity in functionality.

aa.properties User-supplied amino acid distance properties. By default we assume Grantham

(1974) properties.

Details

The purpose of this function is to provide the functionality of a gene based on the inferred parameters from SelAC. The functionality is often used to scale phi.

GetMarginalAllGenes Get marginal reconstruction all genes

Description

Calculates the marginal probability of each codon at all sites across all genes

Usage

```
GetMarginalAllGenes(selac.obj, aa.optim.input = NULL,
  fasta.rows.to.keep = NULL, taxon.to.drop, partition.number = NULL)
```

Arguments

selac.obj An object of class SELAC.

aa.optim.input A list of optimal amino acids with each list element designating a character vector for each gene. The optimal amino acids be the MLE from a selac run

(default) or a list of user defined optimal A.A.

fasta.rows.to.keep

Indicates which rows to remove in the input fasta files.

taxon. to.drop A single taxon (defined by number in phy object) to be removed from the recon-

struction.

partition.number

If only a single gene is desired to be reconstructed, the input is the partition number in the selac object.

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Details

Provides marginal probabilities for all nodes across all genes. The function is fairly simple to use as it only requires as input the selac output object and the working directory that the original analysis took place.

GetPartitionOrder

Get data partiion order

Description

Provides the order of the partitions after the data is read into SELAC.

Usage

```
GetPartitionOrder(codon.data.path)
```

Arguments

codon.data.path

Provides the path to the directory containing the gene specific fasta files of coding data. Must have a ".fasta" line ending.

Details

Provides the order of the partitions when the data is read into SELAC. This function is mainly useful for when users want to supply their own optimal amino acid list into SELAC.

GetSelacPhiCat

Phi rate category information under SELAC+gamma

Description

Provides likelihood information and best rates across sites and across genes under SELAC+gamma

```
GetSelacPhiCat(selac.obj, codon.data.path, aa.optim.input = NULL,
  fasta.rows.to.keep = NULL, n.cores.by.gene.by.site = 1)
```

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Arguments

selac.obj An object of class SELAC.

codon.data.path

Provides the path to the directory containing the gene specific fasta files of coding data.

aa.optim.input A list of optimal amino acids with each list element designating a character vector for each gene. The optimal amino acids be the MLE from a selac run (default) or a list of user defined optimal A.A.

fasta.rows.to.keep

Indicates which rows to remove in the input fasta files.

n.cores.by.gene.by.site

The number of cores to decidate to parallelize analyses by site WITHIN a gene. Note n.cores.by.gene*n.cores.by.gene.by.site is the total number of cores dedicated to the analysis.

Details

The purpose of this function is to determine which rate category best fits each site across genes. The output is a list object, with each list entry designating the optimal rate category across sites for that gene.

GetSelacSiteLikelihoods

Calculate site likelihoods under SelAC

Description

Calculates the likelihoods across sites and across genes under SELAC

Usage

```
GetSelacSiteLikelihoods(selac.obj, codon.data.path,
  aa.optim.input = NULL, fasta.rows.to.keep = NULL)
```

Arguments

An object of class SELAC. selac.obj

codon.data.path

Provides the path to the directory containing the gene specific fasta files of coding data.

aa.optim.input A list of optimal amino acids with each list element designating a character vector for each gene. The optimal amino acids be the MLE from a selac run (default) or a list of user defined optimal A.A.

fasta.rows.to.keep

Indicates which rows to remove in the input fasta files.

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Details

The purpose of this function is to provide the site likelihoods across genes. It is also flexible in that it allows different hypotheses about optimal acids across genes and/or site. The output is a list object, with each list entry designating 1) the tot.likelihood for that gene, and 2) the site likelihoods for that gene.

Index matrix

Example datasets

Description

Matrix index

NucSimulator

Simulate DNA under General-Time Reversible model

Description

Simulates nucleotide data based on parameters under the GTR+G model

Usage

```
NucSimulator(phy, pars, nsites, nuc.model, base.freqs,
  include.gamma = TRUE, gamma.type = "median", ncats = 4,
  start.vals_array = NULL, user.rate.cats = NULL, user.rates = NULL)
```

Arguments

phy	The phylogenetic tree	with branch lengths.
P11.9	The phylogenetic tree	with oranien lengths.

pars A vector of parameters used for the simulation. They are ordered as follows:

gamma shape and the rates for the nucleotide model.

nsites The number of sites to simulate.

nuc.model Indicates what type nucleotide model to use. There are three options: "JC",

"GTR", or "UNREST".

base frequencies for A C G T (in that order).

include.gamma Boolean on whether to use a gamma model

gamma.type How the gamma bins are used

ncats The number of discrete gamma categories.

start.vals_array

A vector of nucleotides to be used as the starting nucleotide for each site in the

simulation.

user.rate.cats The user supplied gamma categories to use instead of choosing at random.

user.rates The user supplied rates to use instead of choosing categories at random.

Details

Simulates a nucleotide matrix using parameters under the GTR+G model. Note that the output can be written to a fasta file using the write.dna() function in the ape package.

PlotEquilbriumCodonDistribution

Function to plot a distribution of frequencies of codons given a 3d array of equilibrium frequency matrices

Description

Function to plot a distribution of frequencies of codons given a 3d array of equilibrium frequency matrices

Usage

```
PlotEquilbriumCodonDistribution(eq.freq.matrices, values,
  palette = "Set1", lwd = 2, ...)
```

Arguments

eq.freq.matrices

A 3d array of eq.freq.matrix returned from ComputeEquilibriumFrequencies

values The vector of labels for each matrix (i.e., different Phi values)

palette Color palette to use from RColorBrewer

lwd Line width

... Other paramters to pass to plot()

PlotExpectedFitness

Function to plot a distribution of fitnesses based on codon equilibrium freqs

Description

Function to plot a distribution of fitnesses based on codon equilibrium freqs

```
PlotExpectedFitness(codon.fitnesses.matrices, codon.eq.matrices, values,
  optimal.aa = NULL, palette = "Set1", lwd = 2,
  include.stop.codon = FALSE, type = "histogram", fitness = TRUE,
  numcode = 1, ...)
```

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Arguments

codon.fitnesses.matrices

A 3d array of aa.fitness.matrix returned from ComputeEquilibriumAAFitness

(first element in return)

codon.eq.matrices

A 3d array of codon equilibrium frequencies

values The vector of labels for each matrix (i.e., different Phi values)

optimal.aa Single letter code for the optimal aa. If NULL, integrates across aa.

palette Color palette to use from RColorBrewer

lwd Line width

include.stop.codon

Include stop codons

type If "histogram", do a histogram plot; if "density", do a density plot

fitness If TRUE, plot W; if FALSE, plot S (= 1 - W)

numcode The genetic code

... Other paramters to pass to plot()

PlotGeneSiteInfo Function to plot info by site in a gene

Description

Function to plot info by site in a gene

Usage

PlotGeneSiteInfo(all.info, aa.properties = NULL, mean.width = 10)

Arguments

all.info The output of GetGeneSiteInfo

aa.properties The aa.properties you want to use; if NULL, uses Grantham

mean.width Sliding window width

PlotMutationFitnessSpectra

Plot fitness of mutations, weighted by frequency of those mutations

Description

Plot fitness of mutations, weighted by frequency of those mutations

Usage

```
PlotMutationFitnessSpectra(mutation.fitness.object.list, values,
  optimal.aa = NULL, palette = "Set1", lwd = 2, ...)
```

Arguments

mutation.fitness.object.list

List that contains multiple objects from ComputeMutationFitnesses() calls

values The vector of labels for each matrix (i.e., different Phi values)

optimal.aa Single letter code for the optimal aa. If NULL, integrates across aa.

palette Color palette to use from RColorBrewer

lwd Line width

... other arguments to pass to plot()

PlotPerAAFitness Function to plot a distribution of fitnesses W or selection coefficients S for a given optimal aa and other terms.

Description

Function to plot a distribution of fitnesses W or selection coefficients S for a given optimal aa and other terms.

```
PlotPerAAFitness(aa.fitness.matrices, values, optimal.aa = NULL,
  palette = "Set1", lwd = 2, include.stop.codon = FALSE,
  type = "histogram", fitness = TRUE, scale.x.axis.by.Ne = FALSE,
  legend.title = NULL, Ne = 10^6, ...)
```

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Arguments

aa.fitness.matrices

A 3d array of aa.fitness.matrix returned from ComputeEquilibriumAAFitness

(first element in return)

values The vector of labels for each matrix (i.e., different Phi values)

optimal.aa Single letter code for the optimal aa. If NULL, integrates across aa.

palette Color palette to use from RColorBrewer

lwd Line width

include.stop.codon

Include stop codons

type If "histogram", do a histogram plot; if "density", do a density plot

fitness If TRUE, plot fitness W; if FALSE, plot selection coefficient S (= W- 1)

scale.x.axis.by.Ne

if TRUE, x axis is transformed from S to S*Ne; if FALSE no scaling is done

legend.title Sets the title of the figure legend.

Ne used to scale x axis when scale.x.axis.by.Ne is TRUE

... Other paramters to pass to plot()

selac example

Example yeast dataset

Description

Example gene, tree, and model output file.

SelacHMMOptimize

Efficient optimization of a Hidden Markov SELAC model

Description

Efficient optimization of model parameters under a HMM SELAC model

```
SelacHMMOptimize(codon.data.path, n.partitions = NULL, phy,
  data.type = "codon", codon.model = "selac",
  edge.length = "optimize", edge.linked = TRUE, nuc.model = "GTR",
  estimate.aa.importance = FALSE, include.gamma = FALSE,
  gamma.type = "quadrature", ncats = 4, numcode = 1,
  diploid = TRUE, k.levels = 0, aa.properties = NULL,
  verbose = FALSE, n.cores.by.gene = 1, n.cores.by.gene.by.site = 1,
  max.tol = 0.001, max.tol.edges = 0.001, max.evals = 1e+06,
```

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```
max.restarts = 3, user.optimal.aa = NULL,
fasta.rows.to.keep = NULL, recalculate.starting.brlen = TRUE,
output.by.restart = TRUE, output.restart.filename = "restartResult",
user.supplied.starting.param.vals = NULL, tol.step = 1,
optimizer.algorithm = "NLOPT_LN_SBPLX", max.iterations = 6)
```

Arguments

codon.data.path

Provides the path to the directory containing the gene specific fasta files of cod-

ing data. Must have a ".fasta" line ending.

n.partitions The number of partitions to analyze. The order is based on the Unix order of the

fasta files in the directory.

phy The phylogenetic tree to optimize the model parameters.

data.type The data type being tested. Options are "codon" or "nucleotide".

codon.model The type of codon model to use. There are four options: "none", "GY94",

"FMutSel0", "selac".

edge.length Indicates whether or not edge lengths should be optimized. By default it is set

to "optimize", other option is "fixed", which user-supplied branch lengths.

edge.linked A logical indicating whether or not edge lengths should be optimized separately

for each gene. By default, a single set of each lengths is optimized for all genes.

nuc.model Indicates what type nucleotide model to use. There are three options: "JC",

"GTR", or "UNREST".

estimate.aa.importance

Indicates whether gene specific importance of distance parameter is to be esti-

mate.

include.gamma A logical indicating whether or not to include a discrete gamma model.

gamma.type Indicates what type of gamma distribution to use. Options are "quadrature"

after the Laguerre quadrature approach of Felsenstein 2001 or median approach

of Yang 1994.

ncats The number of discrete categories.

numcode The ncbi genetic code number for translation. By default the standard (num-

code=1) genetic code is used.

diploid A logical indicating whether or not the organism is diploid or not.

k.levels Provides how many levels in the polynomial. By default we assume a single

level (i.e., linear).

aa.properties User-supplied amino acid distance properties. By default we assume Grantham

(1974) properties.

verbose Logical indicating whether each iteration be printed to the screen.

n.cores.by.gene

The number of cores to dedicate to parallelize analyses across gene.

n.cores.by.gene.by.site

The number of cores to decidate to parallelize analyses by site WITHIN a gene. Note n.cores.by.gene*n.cores.by.gene.by.site is the total number of cores dedicated to the analysis.

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max.tol Supplies the relative optimization tolerance.

max.tol.edges Supplies the relative optimization tolerance for branch lengths only. Default is

that is the same as the max.tol.

max.evals Supplies the max number of iterations tried during optimization.

max.restarts Supplies the number of random restarts.

user.optimal.aa

If optimal.aa is set to "user", this option allows for the user-input optimal amino acids. Must be a list. To get the proper order of the partitions see "GetPartitionOrder" documentation.

fasta.rows.to.keep

Indicates which rows to remove in the input fasta files.

recalculate.starting.brlen

Whether to use given branch lengths in the starting tree or recalculate them.

output.by.restart

Logical indicating whether or not each restart is saved to a file. Default is TRUE.

output.restart.filename

Designates the file name for each random restart.

user.supplied.starting.param.vals

Designates user-supplied starting values for C.q.phi.Ne, Grantham alpha, and Grantham beta. Default is NULL.

tol.step If > 1, makes for coarser tolerance at earlier iterations of the optimizer optimizer.algorithm

The optimizer used by nloptr.

max.iterations Sets the number of cycles to optimize the different parts of the model.

optimal.aa Indicates what type of optimal.aa should be used. There are four options: "none", "majrule", "optimize", or "user".

Details

A hidden Markov model which no longers optimizes the optimal amino acids, but instead allows for the optimal sequence to vary along branches, clades, taxa, etc. Like the original function, we optimize parameters across each gene separately while keeping the shared parameters, alpha, beta, edge lengths, and nucleotide substitution parameters constant across genes. We then optimize alpha, beta, gtr, and the edge lengths while keeping the rest of the parameters for each gene fixed. This approach is potentially more efficient than simply optimizing all parameters simultaneously, especially if fitting models across 100's of genes.

SelacHMMSimulator

Simulate DNA under the SELAC model

Description

Simulates nucleotide data based on parameters under the SELAC model

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Usage

```
SelacHMMSimulator(phy, pars, nsites, codon.freq.by.aa = NULL,
  codon.freq.by.gene = NULL, numcode = 1, aa.properties = NULL,
  nuc.model, include.gamma = FALSE, gamma.type = "quadrature",
  ncats = 4, k.levels = 0, diploid = TRUE, site.cats.vector = NULL)
```

Arguments

phy The phylogenetic tree with branch lengths.

pars A vector of parameters used for the simulation. They are ordered as follows:

C.q.phi, alpha, beta, Ne, base.freqs for A C G, and the rates for the nucleotide model the very last parameter is always the switching rate of the optimal AA.

nsites Length of the alignment to be simulated

codon.freq.by.aa

A matrix of codon frequencies for each possible optimal amino acid. Rows are

aa (including stop codon), cols are codons.

codon.freq.by.gene

A matrix of codon frequencies for each gene.

numcode The ncbi genetic code number for translation. By default the standard (num-

code=1) genetic code is used.

aa.properties User-supplied amino acid distance properties. By default we assume Grantham

(1974) properties.

nuc.model Indicates what type nucleotide model to use. There are three options: "JC",

"GTR", or "UNREST".

include.gamma A logical indicating whether or not to include a discrete gamma model.

gamma.type Indicates what type of gamma distribution to use. Options are "quadrature"

after the Laguerre quadrature approach of Felsenstein 2001 or median approach

of Yang 1994.

ncats The number of discrete categories.

k.levels Provides how many levels in the polynomial. By default we assume a single

level (i.e., linear).

diploid A logical indicating whether or not the organism is diploid or not.

site.cats.vector

A vector designating the rate category for phi when include.gamma=TRUE.

Details

Simulates a nucleotide matrix using parameters under the SELAC model. Note that the output can be written to a fasta file using the write.dna() function in the ape package.

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SelacOptimize	Efficient of	optimization	of the SE	LAC model
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Description

Efficient optimization of model parameters under the SELAC model

Usage

```
SelacOptimize(codon.data.path, n.partitions = NULL, phy,
  data.type = "codon", codon.model = "selac",
  edge.length = "optimize", edge.linked = TRUE,
  optimal.aa = "optimize", nuc.model = "GTR", include.gamma = FALSE,
  gamma.type = "quadrature", ncats = 4, numcode = 1,
  diploid = TRUE, k.levels = 0, aa.properties = NULL,
  verbose = FALSE, n.cores.by.gene = 1, n.cores.by.gene.by.site = 1,
  max.tol = 0.001, max.tol.edges = 0.001, max.evals = 1e+06,
  max.restarts = 3, user.optimal.aa = NULL,
  fasta.rows.to.keep = NULL, recalculate.starting.brlen = TRUE,
  output.by.restart = TRUE, output.restart.filename = "restartResult",
  user.supplied.starting.param.vals = NULL, tol.step = 1,
  optimizer.algorithm = "NLOPT_LN_SBPLX", start.from.mle = FALSE,
  mle.matrix = NULL, partition.order = NULL, max.iterations = 6,
  dt.threads = 1)
```

Arguments

codon.data.pat	h	
	Provides the path to the directory containing the gene specific fasta files of coding data. Must have a ".fasta" line ending.	
n.partitions	The number of partitions to analyze. The order is based on the Unix order of the fasta files in the directory.	
phy	The phylogenetic tree to optimize the model parameters.	
data.type	The data type being tested. Options are "codon" or "nucleotide".	
codon.model	The type of codon model to use. There are four options: "none", "GY94", "YN98", "FMutSel0", "FMutSel", "selac".	
edge.length	Indicates whether or not edge lengths should be optimized. By default it is set to "optimize", other option is "fixed", which is the user-supplied branch lengths.	
edge.linked	A logical indicating whether or not edge lengths should be optimized separately for each gene. By default, a single set of each lengths is optimized for all genes.	
optimal.aa Indicates what type of optimal.aa should be used. There are five options: "non "majrule", "averaged, "optimize", or "user".		
nuc.model	Indicates what type nucleotide model to use. There are three options: "JC", "GTR", or "UNREST".	

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include gamma A logical indicating whether or not to include a discrete gamma model.

gamma.type Indicates what type of gamma distribution to use. Options are "quadrature"

after the Laguerre quadrature approach of Felsenstein 2001 or median approach

of Yang 1994 or "lognormal" after a lognormal quadrature approach.

ncats The number of discrete categories.

numcode The ncbi genetic code number for translation. By default the standard (num-

code=1) genetic code is used.

diploid A logical indicating whether or not the organism is diploid or not.

k.levels Provides how many levels in the polynomial. By default we assume a single

level (i.e., linear).

aa.properties User-supplied amino acid distance properties. By default we assume Grantham

(1974) properties.

verbose Logical indicating whether each iteration be printed to the screen.

n.cores.by.gene

The number of cores to dedicate to parallelize analyses across gene.

n.cores.by.gene.by.site

The number of cores to decidate to parallelize analyses by site WITHIN a gene. Note n.cores.by.gene*n.cores.by.gene.by.site is the total number of cores dedi-

cated to the analysis.

max.tol Supplies the relative optimization tolerance.

max.tol.edges Supplies the relative optimization tolerance for branch lengths only. Default is

that is the same as the max.tol.

max.evals Supplies the max number of iterations tried during optimization.

max.restarts Supplies the number of random restarts.

user.optimal.aa

If optimal.aa is set to "user", this option allows for the user-input optimal amino acids. Must be a list. To get the proper order of the partitions see "GetParti-

tionOrder" documentation.

fasta.rows.to.keep

Indicates which rows to remove in the input fasta files.

recalculate.starting.brlen

Whether to use given branch lengths in the starting tree or recalculate them.

output.by.restart

Logical indicating whether or not each restart is saved to a file. Default is TRUE.

output.restart.filename

Designates the file name for each random restart.

user.supplied.starting.param.vals

Designates user-supplied starting values for C.q.phi.Ne, Grantham alpha, and

Grantham beta. Default is NULL.

tol.step If > 1, makes for coarser tolerance at earlier iterations of the optimizer

optimizer.algorithm

The optimizer used by nloptr.

start.from.mle If TRUE, will start optimization from the MLE. Default is FALSE.

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mle.matrix The user-supplied matrix of parameter values for when start.from.mle is set to TRUE.

partition.order

Allows for a specialized order of the partitions to be gathered from the working directory.

max.iterations Sets the number of cycles to optimize the different parts of the model.

dt.threads Indicates how many available threads to allow data.table to use. Default is zero.

Details

Here we optimize parameters across each gene separately while keeping the shared parameters, alpha, beta, edge lengths, and nucleotide substitution parameters constant across genes. We then optimize alpha, beta, gtr, and the edge lengths while keeping the rest of the parameters for each gene fixed. This approach is potentially more efficient than simply optimizing all parameters simultaneously, especially if fitting models across 100's of genes.

Examples

```
## Not run:
phy <- ape::read.tree(file=system.file("extdata", "rokasYeast.tre", package="selac"))
result <- SelacOptimize(codon.data.path = paste0(find.package("selac"), '/extdata/'),
n.partitions=1, phy=phy, max.evals=10)
print(result)
## End(Not run)</pre>
```

SelacSimulator

Simulate DNA under the SELAC model

Description

Simulates nucleotide data based on parameters under the SELAC model

Usage

```
SelacSimulator(phy, pars, aa.optim_array, codon.freq.by.aa = NULL,
  codon.freq.by.gene = NULL, numcode = 1, aa.properties = NULL,
  nuc.model, include.gamma = FALSE, gamma.type = "quadrature",
  ncats = 4, k.levels = 0, diploid = TRUE, site.cats.vector = NULL)
```

Arguments

phy The phylogenetic tree with branch lengths.

pars A vector of parameters used for the simulation. They are ordered as follows:

C.q.phi, alpha, beta, Ne, base.freqs for A C G, and the rates for the nucleotide

model.

aa.optim_array A vector of optimal amino acids for each site to be simulated.

codon.freq.by.aa

A matrix of codon frequencies for each possible optimal amino acid. Rows are aa (including stop codon), cols are codons.

codon.freq.by.gene

A matrix of codon frequencies for each gene.

numcode The ncbi genetic code number for translation. By default the standard (num-

code=1) genetic code is used.

aa.properties User-supplied amino acid distance properties. By default we assume Grantham

(1974) properties.

nuc.model Indicates what type nucleotide model to use. There are three options: "JC",

"GTR", or "UNREST".

include.gamma A logical indicating whether or not to include a discrete gamma model.

gamma.type Indicates what type of gamma distribution to use. Options are "quadrature"

after the Laguerre quadrature approach of Felsenstein 2001 or median approach

of Yang 1994.

ncats The number of discrete categories.

k.levels Provides how many levels in the polynomial. By default we assume a single

level (i.e., linear).

diploid A logical indicating whether or not the organism is diploid or not.

site.cats.vector

A vector designating the rate category for phi when include.gamma=TRUE.

Details

Simulates a nucleotide matrix using parameters under the SELAC model. Note that the output can be written to a fasta file using the write.dna() function in the ape package.

SelacSimulatorEvolvingRates

Simulate DNA under the SELAC model and evolving rates

Description

Simulates nucleotide data based on parameters under the SELAC model but assumes either Phi or Ne evolves along the tree.

```
SelacSimulatorEvolvingRates(phy, pars, aa.optim_array,
  root.codon.frequencies, numcode = 1, aa.properties = NULL, nuc.model,
  k.levels = 0, diploid = TRUE, pars.to.evolve = "phi",
  evolve.type = "BM", evolve.pars = c(1, 0), Ne.vals.evolved = NULL)
```

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Arguments

phy The phylogenetic tree with branch lengths.

pars A vector of parameters used for the simulation. They are ordered as follows:

C.q.phi, alpha, beta, and Ne.

aa.optim_array A vector of optimal amino acids for each site to be simulated.

root.codon.frequencies

A vector of codon frequencies for each possible optimal amino acid. Thus, the

vector is of length 64x64.

numcode The The ncbi genetic code number for translation. By default the standard (num-

code=1) genetic code is used.

aa.properties User-supplied amino acid distance properties. By default we assume Grantham

(1974) properties.

nuc.model Indicates what type nucleotide model to use. There are three options: "JC",

"GTR", or "UNREST".

k.levels Provides how many levels in the polynomial. By default we assume a single

level (i.e., linear).

diploid A logical indicating whether or not the organism is diploid or not.

pars.to.evolve Indicates which parameters to assume evolve along the tree. Only two options:

"phi" or "Ne".

evolve. type The process by which the focal parameter evovles. There are two options: Brow-

nian motion ("BM") or Ornstein-Uhlenbeck ("OU").

evolve.pars The process parameters used to simulate focal parameter evolution. Under

"BM", the order is root.state, rate; under "OU", the order is alpha, sigma.sq,

and the mean.

Ne.vals.evolved

Under selac we assume a global Ne for all genes. Thus, when the focal parameter to evolve is "Ne", then a user specified vector of simulated Ne values are

provided here.

Details

Simulates a nucleotide matrix using parameters under the SELAC model, but allows either Phi or Ne to evolve along the tree. Note that the output can be written to a fasta file using the write.dna() function in the ape package.

selon example Example archosaur dataset

Description

Example tree and model output file.

SelonHMMOptimize 21

SelonHMMOptimize	Optimize parameters under the HMM SELON model	
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Description

Optimizes model parameters under the HMM SELON model

Usage

```
SelonHMMOptimize(nuc.data.path, n.partitions = NULL, phy,
  edge.length = "optimize", edge.linked = TRUE, nuc.model = "GTR",
  global.nucleotide.model = TRUE, diploid = TRUE, verbose = FALSE,
  n.cores = 1, max.tol = .Machine$double.eps^0.5, max.evals = 1e+06,
  cycle.stage = 12, max.restarts = 10, output.by.restart = TRUE,
  output.restart.filename = "restartResult", fasta.rows.to.keep = NULL)
```

Arguments

fasta.rows.to.keep

nuc.data.path	Provides the path to the directory containing the gene specific fasta files that contains the nucleotide data.
n.partitions	The number of partitions to analyze. The order is based on the Unix order of the fasta files in the directory.
phy	The phylogenetic tree to optimize the model parameters.
edge.length	Indicates whether or not edge lengths should be optimized. By default it is set to "optimize", other option is "fixed", which user-supplied branch lengths.
edge.linked	A logical indicating whether or not edge lengths should be optimized separately for each gene. By default, a single set of each lengths is optimized for all genes.
nuc.model	Indicates what type nucleotide model to use. There are three options: "JC", "GTR", or "UNREST".
global.nucleoti	ide.model
	assumes nucleotide model is shared among all partitions
diploid	A logical indicating whether or not the organism is diploid or not.
verbose	Logical indicating whether each iteration be printed to the screen.
n.cores	The number of cores to run the analyses over.
max.tol	Supplies the relative optimization tolerance.
max.evals	Supplies the max number of iterations tried during optimization.
cycle.stage	Specifies the number of cycles per restart. Default is 12.
max.restarts	Supplies the number of random restarts.
output.by.resta	art
	Logical indicating whether or not each restart is saved to a file. Default is TRUE.
output.restart	
	Designates the file name for each random restart.

Indicates which rows to remove in the input fasta files.

SelonOptimize SelonOptimize

Details

SELON stands for SELection On Nucleotides. This function takes a user supplied topology and a set of fasta formatted sequences and optimizes the parameters in the SELON model. Selection is based on selection towards an optimal nucleotide at each site, which is based simply on the majority rule of the observed data. The strength of selection is then varied along sites based on a Taylor series, which scales the substitution rates. Still a work in development, but so far, seems very promising.

SelonOptimize

Optimize parameters under the SELON model

Description

Optimizes model parameters under the SELON model

Usage

```
SelonOptimize(nuc.data.path, n.partitions = NULL, phy,
  edge.length = "optimize", edge.linked = TRUE,
  optimal.nuc = "majrule", nuc.model = "GTR", set.Ne = 10000,
  diploid = TRUE, verbose = FALSE, n.cores = 1,
  max.tol = .Machine$double.eps^0.25, max.evals = 1e+06,
  cycle.stage = 12, max.restarts = 3, user.optimal.nuc = NULL,
  output.by.restart = TRUE, output.restart.filename = "restartResult",
  user.supplied.starting.param.vals = NULL, fasta.rows.to.keep = NULL,
  recalculate.starting.brlen = TRUE, dt.threads = 1)
```

Arguments

nuc.data.path	Provides the path to the directory containing the gene specific fasta files that contains the nucleotide data.
n.partitions	The number of partitions to analyze. The order is based on the Unix order of the fasta files in the directory.
phy	The phylogenetic tree to optimize the model parameters.
edge.length	Indicates whether or not edge lengths should be optimized. By default it is set to "optimize", other option is "fixed", which user-supplied branch lengths.
edge.linked	A logical indicating whether or not edge lengths should be optimized separately for each gene. By default, a single set of each lengths is optimized for all genes.
optimal.nuc	Indicates what type of optimal.nuc should be used. At the moment there is only a single option: "majrule".
nuc.model	Indicates what type nucleotide model to use. There are three options: "JC", "GTR", or "UNREST".
set.Ne	Indicates whether Ne is to estimated or a fixed value is to be used. Either a fixed value is supplied or "optimize" is use to indicate that it is a parameter to optimize.

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diploid	d A logical indicating whether or not the organism is diploid or not	
verbose	Logical indicating whether each iteration be printed to the screen	
n.cores	The number of cores to run the analyses over.	
max.tol	Supplies the relative optimization tolerance.	
max.evals	Supplies the max number of iterations tried during optimization.	
cycle.stage	Specifies the number of cycles per restart. Default is 12.	
max.restarts	Supplies the number of random restarts.	
user.optimal.nu	IC .	

If optimal.nuc is set to "user", this option allows for the user-input optimal amino acids. Must be a list. To get the proper order of the partitions see "GetPartitionOrder" documentation.

output.by.restart

Logical indicating whether or not each restart is saved to a file. Default is TRUE.

output.restart.filename

Designates the file name for each random restart.

user.supplied.starting.param.vals

Designates user-supplied starting values for C.q.phi.Ne, Grantham alpha, and Grantham beta. Default is NULL.

fasta.rows.to.keep

Indicates which rows to remove in the input fasta files.

recalculate.starting.brlen

Whether to use given branch lengths in the starting tree or recalculate them.

Indicates how many available threads to allow data.table to use. Default is zero. dt.threads

Details

SELON stands for SELection On Nucleotides. This function takes a user supplied topology and a set of fasta formatted sequences and optimizes the parameters in the SELON model. Selection is based on selection towards an optimal nucleotide at each site, which is based simply on the majority rule of the observed data. The strength of selection is then varied along sites based on a Taylor series, which scales the substitution rates.

SelonSimulator	Simulate DNA under the SELON model

Description

Simulates nucleotide data based on parameters under the SELAC model

```
SelonSimulator(phy, pars, nuc.optim_array, nuc.model, diploid = TRUE,
  start.vals_array = NULL)
```

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Arguments

phy The phylogenetic tree with branch lengths.

pars A vector of parameters used for the simulation. They are ordered as follows: a0,

a1, a2, Ne, base.freqs for A C G, and the nucleotide rates.

nuc.optim_array

A vector of optimal nucleotide for each site to be simulated.

nuc.model Indicates what type nucleotide model to use. There are three options: "JC",

"GTR", or "UNREST".

diploid A logical indicating whether or not the organism is diploid or not.

start.vals_array

A vector of nucleotides to be used as the starting nucleotide for each site in the

simulation.

Details

Simulates a nucleotide matrix using parameters under the SELON model. Note that the output can be written to a fasta file using the write.dna() function in the ape package.

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