Package ‘treenomial’

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Type Package

Title Comparison of Trees using a Tree Defining Polynomial

Version 1.1.3

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Description Provides functionality for creation and comparison of polynomials that uniquely describe trees as introduced in Liu (2019, <arXiv:1904.03332>). The core method converts rooted unlabeled phylo objects from 'ape' to the tree defining polynomials described with coefficient matrices. Additionally, a conversion for rooted binary trees with binary trait labels is also provided. Once the polynomials of trees are calculated there are functions to calculate distances, distance matrices and plot different distance trees from a target tree. Manipulation and conversion to the tree defining polynomials is implemented in C++ with 'Rcpp' and 'RcppArmadillo'. Furthermore, parallel programming with 'RcppThread' is used to improve performance converting to polynomials and calculating distances.

Depends R (>= 3.5.0)

License GPL (>= 2)

URL https://github.com/mattgould/treenomial

Encoding UTF-8

Imports Rcpp (>= 1.0.1), ape, methods

LinkingTo Rcpp, RcppArmadillo, RcppThread

RoxygenNote 7.1.1

Suggests knitr, rmarkdown, testthat

NeedsCompilation yes

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alignPoly

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Description
Align various types of coefficient matrices

Usage
alignPoly(coefficientMatrices)

Arguments

  coefficientMatrices
    a list of coefficient matrices of various sizes

Details
Alignment depends on the type of coefficient matrix:

  "real"  the smaller matrices columns are prepended with zero columns to align with
         the max number of columns and the rows are appended with zero rows to match
         the max number of rows

  "yEvaluated"  the smaller vectors are appended with zeroes to match the max length vector

  "tipLabel"  the smaller matrices are appended with zeroes to match the max number of rows and
              columns

Value
the aligned list of coefficient matrices
allTrees

Examples

library(treenomial)
library(ape)
differentSizeTrees <- c(rtree(2), rmree(10, 10))
coeffs <- treeToPoly(differentSizeTrees, numThreads = 0)
alinedCoeffs <- alignPoly(coeffs)

Description

Return normal coefficient matrices, substituted y coefficient vectors, or phylo objects for all possible unordered full m-ary trees up to n tips. For binary trees (m = 2), the number of trees at each number of tips follows the Wedderburn-Etherington numbers.

Usage

allTrees(n, m = 2, type = c("default", "yEvaluated", "phylo"), y)

Arguments

n
max number of tips

m
max number of children for each node

type
one of:

"real" tree distinguishing polynomials in two variables x (columns) and y (rows)

"yEvaluated" tree distinguishing polynomials with y evaluated at a specified argument

"phylo" phylo objects

y
the y value to evaluate the polynomial at when type is "yEvaluated", ignored otherwise

Value

list of lists containing all the trees in type format for each number of tips

Note

only m = 2 is currently supported
**Examples**

```r
library(treenomial)
library(ape)

# generate coefficient matrices describing the polynomials of all possible
# unordered full binary trees up to 10 tips
allBinTenRealCoeff <- allTrees(10, type = "phylo")

# number of trees at each number of tips follows Wedderburn-Etherington numbers
lengths(allBinTenRealCoeff)

# phylo type example, plot all 6 tip unordered full binary trees

# backup par options
oldpar <- par(no.readonly = TRUE)

allBinSixPhylo <- allTrees(6, type = "phylo")[[6]]
par(mfrow = c(1, 6))
plots <- lapply(allBinSixPhylo, function(t){
  plot.phylo(ladderize(t), direction = "downwards", show.tip.label = FALSE)
})

# restore par options
par(oldpar)
```

---

**plotExtremeTrees**

*Plot the min/max distance trees from a target tree*

**Description**

Plot the min/max distance trees from a target tree

**Usage**

```r
plotExtremeTrees(
  target, 
  trees, 
  n, 
  comparison = "min", 
  method = c("fraction", "logDiff", "wLogDiff", "pa", "ap"), 
  type = c("default", "yEvaluated", "tipLabel"), 
  y, 
  numThreads = -1
)
```
Arguments

- **target**: the phylo object of the tree to calculate the distances to
- **trees**: a list of phylo objects to compare with the *target*
- **n**: the number of trees to find and plot
- **comparison**: whether to find the “min” or the “max” distance trees from the *target*
- **method**: method to use when calculating coefficient distances:
  - “fraction” for two coefficient matrices A and B returns \( \text{sum}(|A-B|)/(A+B) \), excluding elements where both A and B are zero
  - “logDiff” for two coefficient matrices A and B returns \( \text{sum}(|\log(1+|A-B|)) \)
  - “wLogDiff” performs the “logDiff” method with weights on the rows
  - “pa” total pairs where the coefficient is present in one matrix and absent in the other (presence-absence)
  - “ap” opposite comparison of pa (absence-presence)
- **type**: one of:
  - “real” tree distinguishing polynomials in two variables x (columns) and y (rows)
  - “yEvaluated” tree distinguishing polynomials with y evaluated at a specified argument
  - “tipLabel” complex coefficient polynomial that utilize binary trait tip labels on the phylo objects
- **y**: the y value to evaluate the polynomial at when type is “yEvaluated”, ignored otherwise
- **numThreads**: number of threads to be used, the default (-1) will use the number of cores in the machine and numThreads = 0 will only use the main thread

Value

- a list of lists containing the *n* min/max distance trees and their distances to *target*

Note

- the substituted y coefficient vector only supports the “logDiff” method and the “fraction” method
- “pa” and “ap” force symmetry in the output distance matrix

Examples

```r
library(treenomial)
library(ape)
trees <- c(rmtree(1000, 50), rmtree(10, 9))
target <- rtree(50)
minTrees <- plotExtremeTrees(target, trees, 2, comparison = "min", numThreads = 0)
```
polyDist  
*Calculates the distance between coefficient matrices*

**Description**

Calculates the distance between two coefficient matrices or a coefficient matrix and a list of coefficient matrices.

**Usage**

```r
polyDist(
  x, 
  Y, 
  method = c("fraction", "logDiff", "wLogDiff", "pa", "ap"),
  numThreads = -1
)
```

**Arguments**

- `x`: single coefficient matrix to find distances to
- `Y`: a list of coefficient matrices
- `method`: method to use when calculating coefficient distances:
  - "fraction": for two coefficient matrices A and B returns \( \text{sum}(\text{abs}(A-B)/(A+B)) \), excluding elements where both A and B are zero
  - "logDiff": for two coefficient matrices A and B returns \( \text{sum}(\log(1+\text{abs}(A-B))) \)
  - "wLogDiff": performs the "logDiff" method with weights on the rows
  - "pa": total pairs where the coefficient is present in one matrix and absent in the other (presence-absence)
  - "ap": opposite comparison of pa (absence-presence)
- `numThreads`: number of threads to be used, the default (-1) will use the number of cores in the machine and numThreads = 0 will only use the main thread

**Value**

vector of distances

**Note**

- the substituted y coefficient vector only supports the “logDiff” method and the “fraction” method
- “pa” and “ap” force symmetry in the output distance matrix
Examples

```r
library(treenomial)
library(ape)

# distance between coefficient matrices of one 10 tip tree
# and 100 trees with 30 tips using
# create the coefficient matrices
tenTipTree <- rtree(10)
tenTipTreeCoeff <- treeToPoly(tenTipTree, numThreads = 0)

thirtyTipList <- rmrree(100, 30)
thirtyTipCoeffs <- treeToPoly(thirtyTipList, numThreads = 0)

# find the distance
polyDist(tenTipTreeCoeff, thirtyTipCoeffs, numThreads = 0)
```

polyToDistMat

Calculates the distance matrix from a list coefficient matrices

Description

Calculates the distance matrix from a list coefficient matrices

Usage

```r
polyToDistMat(
  coefficientMatrices,
  method = c("fraction", "logDiff", "wLogDiff", "pa", "ap"),
  numThreads = -1
)
```

Arguments

- `coefficientMatrices` list of coefficient matrices
- `method` method to use when calculating coefficient distances:
  - "fraction" for two coefficient matrices A and B returns \( \frac{\sum|A-B|}{A+B} \), excluding elements where both A and B are zero
  - "logDiff" for two coefficient matrices A and B returns \( \sum \log(1+|A-B|) \)
  - "wLogDiff" performs the "logDiff" method with weights on the rows
  - "pa" total pairs where the coefficient is present in one matrix and absent in the other (presence-absence)
  - "ap" opposite comparison of pa (absence-presence)
- `numThreads` number of threads to be used, the default (-1) will use the number of cores in the machine and numThreads = 0 will only use the main thread
Value

distance matrix calculated from argument coefficient matrices

Note

• the substituted y coefficient vector only supports the “logDiff” method and the “fraction”
  method
• “pa” and “ap” force symmetry in the output distance matrix

Examples

library(treemomial)
library(ape)

# coefficient matrices for ten trees of 20 tips
coeffs <- treeToPoly(rmtree(10, 20), numThreads = 0)

# distance matrix from the list of coefficient matrices
d <- polyToDistMat(coeffs, method = "logDiff", numThreads = 0)

# using the absence-presence method
d <- polyToDistMat(coeffs, method = "ap", numThreads = 0)

Description

Calculates the distance between trees.

Usage

treeDist(
  x,
  Y,
  type = c("default", "yEvaluated", "tipLabel"),
  method = c("fraction", "logDiff", "wLogDiff", "pa", "ap"),
  y,
  numThreads = -1
)

Arguments

x single phylo object
Y a list of phylo objects
type one of:
treeDist

“real” tree distinguishing polynomials in two variables x (columns) and y (rows)
“yEvaluated” tree distinguishing polynomials with y evaluated at a specified argument
“tipLabel” complex coefficient polynomial that utilize binary trait tip labels on the phylo objects

method method to use when calculating coefficient distances:
“fraction” for two coefficient matrices A and B returns sum(abs(A-B)/(A+B)), excluding elements where both A and B are zero
“logDiff” for two coefficient matrices A and B returns sum(log(1+abs(A-B))
“wLogDiff” performs the “logDiff” method with weights on the rows
“pa” total pairs where the coefficient is present in one matrix and absent in the other (presence-absence)
“ap” opposite comparison of pa (absence-presence)

y the y value to evaluate the polynomial at when type is “yEvaluated”, ignored otherwise
numThreads number of threads to be used, the default (-1) will use the number of cores in the machine and numThreads = 0 will only use the main thread

Value

vector of distances

Note

• the substituted y coefficient vector only supports the “logDiff” method and the “fraction” method
• “pa” and “ap” force symmetry in the output distance matrix

Examples

library(treenomial)
library(ape)

# distance between one 10 tip tree and 100 trees with 30 tips

# generate the trees
tenTipTree <- rtree(10)
thirtyTipList <- rmimage(100, 30)

# find the distance
treeDist(tenTipTree, thirtyTipList, numThreads = 0)
treeJuliaSet

Plots a Julia Set for a tree

Description

 Finds the Julia Set for the y evaluated polynomial of a tree and plots in a square image.

Usage

```r
treeJuliaSet(
  tree,
  pixelLength = 700,
  center = 0,
  maxZ = 2,
  maxIter = 100,
  col = c("white", colorRampPalette(c("dodgerblue4", "lightblue"))(98), "black"),
  y
)
```

Arguments

- **tree**: phylo object
- **pixelLength**: number of pixels on one side of the image
- **center**: complex number giving the center of the image on the complex plane
- **maxZ**: the max value for the real and imaginary axis
- **maxIter**: maximum count for iterations
- **col**: colours to be used for the image
- **y**: the y value to evaluate the polynomial at

Examples

```r
library(treenomial)
library(ape)
treeJuliaSet(stree(5,type = "right"), y = 1+1i)
```
treeToDistMat

Calculates the distance matrix from a list of phylo objects

Description

Calculates the distance matrix from a list of phylo objects

Usage

treeToDistMat(
  trees,
  method = c("fraction", "logDiff", "wLogDiff", "pa", "ap"),
  type = c("default", "yEvaluated", "tipLabel"),
  y,
  numThreads = -1
)

Arguments

- **trees**: a single phylo object or a list of phylo objects
- **method**: method to use when calculating coefficient distances:
  - "fraction" for two coefficient matrices A and B returns sum(abs(A-B)/(A+B)), excluding elements where both A and B are zero
  - "logDiff" for two coefficient matrices A and B returns sum(log(1+abs(A-B))
  - "wLogDiff" performs the "logDiff" method with weights on the rows
  - "pa" total pairs where the coefficient is present in one matrix and absent in the other (presence-absence)
  - "ap" opposite comparison of pa (absence-presence)
- **type**: one of:
  - "real" tree distinguishing polynomials in two variables x (columns) and y (rows)
  - "yEvaluated" tree distinguishing polynomials with y evaluated at a specified argument
  - "tipLabel" complex coefficient polynomial that utilize binary trait tip labels on the phylo objects
- **y**: the y value to evaluate the polynomial at when type is "yEvaluated", ignored otherwise
- **numThreads**: number of threads to be used, the default (-1) will use the number of cores in the machine and numThreads = 0 will only use the main thread

Value

- a distance matrix
Note

- the substituted y coefficient vector only supports the “logDiff” method and the “fraction” method
- “pa” and “ap” force symmetry in the output distance matrix

Examples

```r
library(treenomial)
library(ape)
# distance matrix for 10 trees of 30 tips
treeToDistMat(rmtree(10, 30), method = "wLogDiff", numThreads = 0)
```

treeToPoly

Convert trees to coefficient matrices

Description

Converts rooted full binary trees to tree distinguishing polynomials described with coefficient matrices.

Usage

```r
treeToPoly(
  trees,
  type = c("default", "yEvaluated", "tipLabel"),
  y,
  varLabels = FALSE,
  numThreads = -1
)
```

Arguments

trees: a single phylo object or a list of phylo objects

type: one of:
  - "real": tree distinguishing polynomials in two variables x (columns) and y (rows)
  - "yEvaluated": tree distinguishing polynomials with y evaluated at a specified argument
  - "tipLabel": complex coefficient polynomial that utilize binary trait tip labels on the phylo objects

y: the y value to evaluate the polynomial at when type is “yEvaluated”, ignored otherwise

varLabels: boolean for whether to add row and column names corresponding to the variables in the polynomial

numThreads: number of threads to be used, the default (-1) will use the number of cores in the machine and numThreads = 0 will only use the main thread
Value

the resulting coefficient matrix or matrices of the form:

`real`  a real matrix where the ith row, jth column represents the x^(j-1)*y^(i-1) coefficient

`yEvaluated`  a vector where the kth column represents the x^(k-1) coefficient

`tipLabel`  given trees with two unique tip labels “a”, “b” a complex matrix where the ith row, jth column represents the a^(i-1)*b^(j-1) coefficient

Examples

```r
library(treenomial)
library(ape)

# generate a tree
tree <- rtree(n = 30, rooted = TRUE)

# a real coefficient matrix
treeToPoly(tree, varLabels = TRUE, numThreads = 0)

# complex coefficient vector for the tree
treeToPoly(tree, type = "yEvaluated", y = 1+1i, varLabels = TRUE, numThreads = 0)

# for a list of trees
treeToPoly(rmtree(4, 20), varLabels = TRUE, numThreads = 0)
```

wedge

Performs the wedge operation

Description

Calculates the result from the wedge operation on two real coefficient matrices, two y evaluated polynomial coefficient vectors or two phylo objects.

Usage

```r
wedge(A, B, type = c("default", "yEvaluated", "phylo"), y)
```

Arguments

- `A, B`  two real coefficient matrices, complex coefficient vectors or phylo objects
- `type`  one of:
  - `"real"`  tree distinguishing polynomials in two variables x (columns) and y (rows)
  - `"yEvaluated"`  tree distinguishing polynomials with y evaluated at a specified argument
  - `"tipLabel"`  complex coefficient polynomial that utilize binary trait tip labels on the phylo objects
- `y`  the y value to evaluate the polynomial at when type is "yEvaluated", ignored otherwise
Value

the wedge result in the same form as the arguments

Examples

library(treenomial)
library(ape)

# wedge two real coefficient matrices
leaf <- matrix(c(0,1), nrow = 1, ncol = 2)
wedge(leaf, leaf)

# wedge two complex coefficient vectors
leaf <- as.complex(c(0,1))
wedge(leaf, leaf, "yEvaluated",5)
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