

clstutils

October 25, 2011

`classifyPlacements` *Taxonomic classification by phylogenetic placement.*

Description

Given taxonomic information from a reference package and inter-node distances from a reference tree, perform classification of one or more placements provided by `pplacer`.

Usage

```
classifyPlacements(taxdata, treedists, placetab, ...,  
                   verbose = FALSE, debug = FALSE)
```

Arguments

| | |
|------------------------|--|
| <code>taxdata</code> | <code>data.frame</code> , output of <code>taxonomyFromRefpkg</code> |
| <code>treedists</code> | output of <code>treeDists</code> |
| <code>placetab</code> | a <code>data.frame</code> with columns <code>at</code> , <code>edge</code> , and <code>branch</code> |
| <code>...</code> | extra arguments passed to <code>classifyIter</code> |
| <code>verbose</code> | writes progress messages to terminal if TRUE |
| <code>debug</code> | be very verbose if TRUE |

Value

The output is a `data.frame` describing the taxonomic assignment, along with a description of the confidence of the classification. See the man page for `classify` for details on the output.

Author(s)

Noah Hoffman

See Also

`treeDists`, `taxonomyFromRefpkg`

Examples

```
placefile <- system.file('extdata', 'merged.json', package='clstutils')
distfile <- system.file('extdata', 'merged.distmat.bz2', package='clstutils')
refpkgz <- system.file('extdata', 'vaginal_16s.refpkg.tar.bz2', package='clstutils')

tmpdir <- tempdir()

orig.dir <- getwd()
setwd(tmpdir)
system(paste("tar --no-same-owner -xjf", refpkgz))
setwd(orig.dir)

refpkg <- file.path(tmpdir, "vaginal_16s.refpkg")

treedists <- treeDists(distfile=distfile, placefile=placefile)
taxdata <- taxonomyFromRefpkg(refpkg, seqnames=rownames(treedists$dmat), lowest_rank="spe")
placetab <- data.frame(at=49, edge=5.14909e-07, branch=5.14909e-07)
result <- classifyPlacements(taxdata, treedists, placetab)
result
```

clstutils-package *Sequence based classification and selection of reference sequences.*

Description

Tools for performing taxonomic assignment based on phylogeny using pplacer and clst.

Details

Package: clstutils
Type: Package
Author: Noah Hoffman <ngh2@uw.edu>
License: GPL3

Index:

Author(s)

Noah Hoffman

Maintainer: <ngh2@uw.edu>

See Also

[clst](#)

Examples

```
library(clstutils)
packageDescription("clstutils")
```

| | |
|--------------|---|
| findOutliers | <i>Identify outlier objects given a square distance matrix.</i> |
|--------------|---|

Description

Outliers are defined as elements with edge length to the centermost element > cutoff. The distance threshold (cutoff) can be either specified, or calculated as a quantile of all pairwise distances in the matrix.

Usage

```
findOutliers(mat, quant, cutoff)
```

Arguments

| | |
|--------|--|
| mat | square matrix of distances |
| quant | given all pairwise distances x, calculate distance threshold as quantile(x, quant). Values closer to 0 are more stringent. |
| cutoff | an absolute cutoff overriding quant |

Value

Returns a boolean vector corresponding to margin of mat; outliers have a value of TRUE.

Author(s)

Noah Hoffman

Examples

```
library(ape)
data(seqs)
data(seqdat)
dmat <- ape::dist.dna(seqs[seqdat$tax_name == 'Enterococcus faecium',],
  pairwise.deletion=TRUE, as.matrix=TRUE, model='raw')
summary(dmat[lower.tri(dmat)])
outliers <- findOutliers(dmat, cutoff=0.015)
table(outliers)
```

`maxDists`*Select a maximally diverse set of items given a distance matrix.*

Description

Given a square matrix of pairwise distances, return indices of N objects with a maximal sum of pairwise distances.

Usage

```
maxDists(mat, idx = NA, N = 1,
         exclude = rep(FALSE, nrow(mat)),
         include.center = TRUE)
```

Arguments

| | |
|-----------------------------|--|
| <code>mat</code> | square distance matrix |
| <code>idx</code> | starting indices; if missing, starts with the object with the maximum median distance to all other objects. |
| <code>N</code> | total number of selections; length of <code>idx</code> is subtracted. |
| <code>exclude</code> | boolean vector indicating elements to exclude from the calculation. |
| <code>include.center</code> | includes the "most central" element (ie, the one with the smallest median of pairwise distances to all other elements) if TRUE |

Value

A vector of indices corresponding to the margin of `mat`.

Note

Note that it is important to evaluate if the candidate sequences contain outliers (for example, mislabeled sequences), because these will assuredly be included in a maximally diverse set of elements!

Author(s)

Noah Hoffman

See Also

[findOutliers](#)

Examples

```
library(ape)
library(clstutils)
data(seqs)
data(seqdat)
efaecium <- seqdat$tax_name == 'Enterococcus faecium'
seqdat <- subset(seqdat, efaecium)
seqs <- seqs[efaecium,]
dmat <- ape::dist.dna(seqs, pairwise.deletion=TRUE, as.matrix=TRUE, model='raw')
```

```
## find a maximally diverse set without first identifying outliers
picked <- maxDists(dmat, N=10)
picked
prettyTree(nj(dmat), groups=ifelse(1:nrow(dmat) %in% picked, 'picked', 'not picked'))

## restrict selected elements to non-outliers
outliers <- findOutliers(dmat, cutoff=0.015)
picked <- maxDists(dmat, N=10, exclude=outliers)
picked
prettyTree(nj(dmat), groups=ifelse(1:nrow(dmat) %in% picked, 'picked', 'not picked'),
X = outliers)
```

```
prettyTree          Draw an annotated phylogenetic tree.
```

Description

Extends `plot.phylo` to draw a phylogenetic tree with additional annotation.

Usage

```
prettyTree(x, groups, fill,
           X, O, indices, labels,
           show = rep(TRUE, length(x)),
           largs = list(cex = 0.75, bty = "n", yjust = 0.5),
           parargs = list(mar = c(bottom = 5, left = 2, top = 2,
                                   right = ifelse(is.null(largs), 2, 8))),
           xpd = NA),
           pointargs = list(), glyphs,
           shuffleGlyphs = NA, ...)
```

Arguments

| | |
|------------------------|---|
| <code>x</code> | an object of class <code>phylo</code> , eg <code>x <- nj(ddist)</code> |
| <code>groups</code> | a factor (or object coercible) to a factor assigning group identity to leaf nodes in <code>x</code> |
| <code>fill</code> | vector (logical or indices) of points to fill |
| <code>X</code> | vector of points to mark with an X |
| <code>O</code> | vector of points to mark with a circle |
| <code>indices</code> | label points with indices (all points if 'yes', or a subset indicated by a vector) |
| <code>labels</code> | character vector of tip labels in the same order as <code>x\$tip.label</code> |
| <code>show</code> | boolean vector of points to show |
| <code>largs</code> | arguments controlling appearance of the legend or NULL for no legend |
| <code>parargs</code> | arguments to pass <code>par()</code> |
| <code>pointargs</code> | arguments to pass <code>points()</code> (other than <code>pch</code> , <code>col</code> , <code>bg</code>) |

`glyphs` a `data.frame` with columns named 'col' and 'pch' corresponding to elements of `unique(groups)`
`shuffleGlyphs` NA or an integer (argument to `set.seed`)
`...` passed to `plot.phylo`

Details

`prettyTree` adds to a plot drawn by `plot.phylo`

Vectors specifying annotation should be in the order of row or column labels of the distance matrix used to generate `x`.

Value

Plots to the active device; no return value.

Note

See package vignette for additional examples.

Author(s)

Noah Hoffman

See Also

`plot.phylo`

Examples

```
library(ape)
data(seqs)
data(seqdat)
prettyTree(nj(dist.dna(seqs)), groups=seqdat$tax_name)
```

`refpkgContents` *Read the contents of a collection of reference sequences ("refpkg").*

Description

Read the manifest file from a repackage and return a list containing the package contents.

Usage

```
refpkgContents(path, manifest = "CONTENTS.json")
```

Arguments

`path` path to a repackage directory
`manifest` name of the manifest file

Value

Returns a list of lists. Run `example(refpkgContents)` for details.

Author(s)

Noah Hoffman

References

The description and specification for a reference package can be found in the project repository in github: <https://github.com/fhcr/taxtastic>

Scripts and tools for creating reference packages are provided in the python package `taxonomy`, also available from the `taxtastic` project site.

See Also

[taxonomyFromRefpkg](#)

Examples

```
archive <- 'vaginal_16s.refpkg.tar.bz2'
destdir <- tempdir()
system(gettextf('tar -xjf %s --directory="%s"',
               system.file('extdata', archive, package='clstutils'),
               destdir))
refpkg <- file.path(destdir, sub('.tar.bz2','',archive))
contents <- refpkgContents(refpkg)
str(refpkg)
```

seqdat

Annotation for the Enterococcus sequence data set.

Description

Provides annotation for `link{seqs}`, an aligned 16S rRNA sequences representing three *Enterococcus* species.

Usage

```
data(seqdat)
```

Format

A data frame with 200 observations on the following 5 variables.

`seqname` a character vector

`accession` a character vector containing GenBank accession numbers.

`tax_id` a character vector

`tax_name` a character vector

`isType` a logical vector indicating if the sequence is from a type strain.

Source

These sequences were downloaded from the Ribosomal Database Project website <http://rdp.cme.msu.edu/>

Examples

```
data(seqdat)
with(seqdat, {
  table(tax_name, isType)
})
```

| | |
|------|--|
| seqs | <i>Enterococcus sequence data set.</i> |
|------|--|

Description

Aligned 16S rRNA sequences representing three *Enterococcus* species.

Usage

```
data(seqs)
```

Format

The format is: 'DNAbin' raw [1:200, 1:1848] - - - ... - attr(*, "dimnames")=List of 2 ..\$: chr [1:200] "S000001976" "S000008133" "S000013428" "S000127028"\$: NULL

Source

These sequences were downloaded from the Ribosomal Database Project website <http://rdp.cme.msu.edu/>

Examples

```
data(seqs)
seqs
```

| | |
|--------------------|---|
| taxonomyFromRefpkg | <i>Extract taxonomic information from a refpkg.</i> |
|--------------------|---|

Description

Construct a data.frame providing the lineage of each sequence represented in the reference package.

Usage

```
taxonomyFromRefpkg(path, seqnames, lowest_rank = NA)
```

Arguments

| | |
|-------------|---|
| path | path to a refpkg directory |
| seqnames | optional character vector of sequence names. If provided, determines the order of rows in \$taxTab |
| lowest_rank | name of the most specific (ie, rightmost) rank to include. Default is the name of the rightmost column in refpkg_contents\$taxonomy |

Value

A list with the following elements:

| | |
|----------|--|
| taxNames | a named character vector of taxonomic names (names are tax_ids) |
| taxTab | a <code>data.frame</code> in which each row corresponds to a reference sequence and contains a tax_id followed by the corresponding lineage (columns are "root"...lowest_rank) |

Author(s)

Noah Hoffman

References

The description and specification for a reference package can be found in the project repository in github: <https://github.com/fhcr/taxtastic>

Scripts and tools for creating reference packages are provided in the python package `taxonomy`, also available from the `taxtastic` project site.

See Also

[refpkgContents](#)

Examples

```
archive <- 'vaginal_16s.refpkg.tar.bz2'
destdir <- tempdir()
system(gettextf('tar -xjf %s --directory="%s"',
               system.file('extdata', archive, package='clstutils'),
               destdir))
refpkg <- file.path(destdir, sub('.tar.bz2', '', archive))
reftax <- taxonomyFromRefpkg(refpkg)
str(reftax)
```

treeDists

Provide objects for determining distances among nodes of a reference

Description

Provides objects (dists, paths) that can be used to calculate vectors of distances between an internal node and each leaf node. Also returns a square matrix of distances between leaf nodes.

Usage

```
treeDists(placefile, distfile)
```

Arguments

placefile path to pplacer output
 distfile path to output of guppy distmat

Details

A placement on an edge looks like this:

```

proximal
|
|   d_p
|
|---- x
|
|   d_d
|
|
distal

```

d_p is the distance from the placement x to the proximal side of the edge, and d_d the distance to the distal side.

If the distance from x to a leaf y is an S-distance Q , then the path from x to y will go through the distal side of the edge and we will need to add d_d to Q to get the distance from x to y . If the distance from x to a leaf y is a P-distance Q , then the path from x to y will go through the proximal side of the edge, and we will need to subtract off d_d from Q to get the distance from x to y . In either case, we always need to add the length of the pendant edge, which is the second column.

To review, say the values of the two leftmost columns are a and b for a given placement x , and that it is on an edge i . We are interested in the distance of x to a leaf y , which is on edge j . We look at the distance matrix, entry (i,j) , and say it is an S-distance Q . Then our distance is $Q+a+b$. If it is a P-distance Q , then the distance is $Q-a+b$.

The distances between leaves should always be P-distances, and there we need no trickery.

(thanks to Erick Matsen for this description)

Value

A list with the following elements:

`dists` rectangular matrix of distances with rows corresponding to all nodes in pplacer order, and columns corresponding to tips in the order of the corresponding `phylo{ape}` object.
`paths` rectangular matrix in the same configuration as `dists` with values of 1 or -1 if the path between nodes is serial or parallel, respectively (see Details)
`dmat` square matrix containing distances between pairs of tips.

Note

The output of this function is required for `classifyPlacements`.

Author(s)

Noah Hoffman

References

Documentation for pplacer and guppy can be found here: <http://matsen.fhcrc.org/pplacer/>

See Also

[classifyPlacements](#)

Examples

```
placefile <- system.file('extdata','merged.json', package='clstutils')
distfile <- system.file('extdata','merged.distmat.bz2', package='clstutils')
treedists <- treeDists(placefile, distfile)

## coordinates of a single placement
placetab <- data.frame(at=49, edge=5.14909e-07, branch=5.14909e-07)

## dvects is a matrix in which each row corresponds to a vector of
## distances between a single placement along the edge of the reference
## tree used to generate 'distfile', and each column corresponds to a
## reference sequence (ie, a terminal node).

dvects <- with(placetab, {
treedists$dists[at+1,,drop=FALSE] + treedists$paths[at+1,,drop=FALSE]*edge + branch
})
```

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