Package 'clstutils'

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Type Package

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LazyLoad yes

LazyData yes

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

License GPL-3

biocViews Sequencing, Classification, Visualization, QualityControl

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clstutils-package

Description

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

Details

Package:	clstutils
Type:	Package
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License:	GPL3

Index:

Author(s)

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See Also

clst

Examples

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

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)
```

classifyPlacements

Arguments

taxdata	data.frame, output of taxonomyFromRefpkg
treedists	output of treeDists
placetab	a data.frame with columns at, edge, and branch
	extra arguments passed to classifyIter
verbose	writes progress messages to terminal if TRUE
debug	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

result

```
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.gz, package=clstutils)
tmpdir <- tempdir()
orig.dir <- getwd()
setwd(tmpdir)
system(paste("tar --no-same-owner -xzf", 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="species")
placetab <- data.frame(at=49, edge=5.14909e-07, branch=5.14909e-07)
result <- classifyPlacements(taxdata, treedists, placetab)</pre>
```

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findOutliers

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

maxDists

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

mat	square distance matrix
idx	starting indices; if missing, starts with the object with the maximum median distance to all other objects.
Ν	total number of selections; length of idx is subtracted.
exclude	boolean vector indicating elements to exclude from the calculation.
include.center	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</pre>
seqdat <- subset(seqdat, efaecium)</pre>
seqs <- seqs[efaecium,]</pre>
dmat <- ape::dist.dna(seqs, pairwise.deletion=TRUE, as.matrix=TRUE, model=raw)</pre>
## find a maximally diverse set without first identifying outliers
picked <- maxDists(dmat, N=10)</pre>
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)</pre>
picked <- maxDists(dmat, N=10, exclude=outliers)</pre>
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

Arguments

Х

an object of class phylo, eg x <- nj(ddist)

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prettyTree

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

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

refpkgContents

Description

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

Usage

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

Arguments

path	path to a refpkg directory
manifest	name of the manifest file

Value

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

Author(s)

Noah Hoffman

References

The decsription and specification for a reference package can be found in the project repository in github: https://github.com/fhcrc/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

seqdat

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

Enterococcus sequence data set.

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" "S00008133" "S000013428" "S000127028" ... ...$ : NULL
```

Source

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

Examples

data(seqs) seqs

taxonomyFromRefpkg Extract taxonomic information from a refpkg.

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 data.frame in which each row corresponds to a reference sequence and con-
	tains a tax_id followed by the corresponding lineage (columns are "root"lowest_rank)

Author(s)

Noah Hoffman

treeDists

References

The decsription and specification for a reference package can be found in the project repository in github: https://github.com/fhcrc/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

treeDists	Provide objects for determining distances among nodes of a reference
	tree.

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

treeDists

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

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

```
## 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 correspons 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
})</pre>
```

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