Package ‘ips’

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R topics documented:

ips-package ........................................... 2
aliscore .............................................. 3
code.simple.gaps .................................... 5
collapseUnsupportedEdges ........................... 5
del.miss ............................................. 6
deleteEmptyCells ................................... 7
delGaps ............................................ 8
descendants ........................................ 9
DNAbin2index ....................................... 10
eoi ................................................... 10
**ips-package**  
*Interfaces to Phylogenetic Software*

<table>
<thead>
<tr>
<th>Function</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>fixNodes</td>
<td>12</td>
</tr>
<tr>
<td>forceEqualTipHeights</td>
<td>13</td>
</tr>
<tr>
<td>gblocks</td>
<td>14</td>
</tr>
<tr>
<td>index2DNABin</td>
<td>16</td>
</tr>
<tr>
<td>ips.16S</td>
<td>16</td>
</tr>
<tr>
<td>ips.28S</td>
<td>17</td>
</tr>
<tr>
<td>ips.cox1</td>
<td>17</td>
</tr>
<tr>
<td>ips.tree</td>
<td>18</td>
</tr>
<tr>
<td>mafft</td>
<td>19</td>
</tr>
<tr>
<td>mafft.merge</td>
<td>21</td>
</tr>
<tr>
<td>mrbayes</td>
<td>21</td>
</tr>
<tr>
<td>mrbayes.lset</td>
<td>23</td>
</tr>
<tr>
<td>mrbayes.mcmc</td>
<td>24</td>
</tr>
<tr>
<td>mrbayes.preset</td>
<td>26</td>
</tr>
<tr>
<td>multistate</td>
<td>28</td>
</tr>
<tr>
<td>neighboringPairs</td>
<td>29</td>
</tr>
<tr>
<td>ntip</td>
<td>30</td>
</tr>
<tr>
<td>partitionfinder</td>
<td>30</td>
</tr>
<tr>
<td>pathd8</td>
<td>31</td>
</tr>
<tr>
<td>phylo2mafft</td>
<td>32</td>
</tr>
<tr>
<td>phylo2mst</td>
<td>33</td>
</tr>
<tr>
<td>pis</td>
<td>33</td>
</tr>
<tr>
<td>prank</td>
<td>34</td>
</tr>
<tr>
<td>raxml</td>
<td>35</td>
</tr>
<tr>
<td>raxml.partitions</td>
<td>38</td>
</tr>
<tr>
<td>rbeauti</td>
<td>39</td>
</tr>
<tr>
<td>read</td>
<td>40</td>
</tr>
<tr>
<td>read.beast</td>
<td>41</td>
</tr>
<tr>
<td>read.beast.table</td>
<td>42</td>
</tr>
<tr>
<td>sister</td>
<td>43</td>
</tr>
<tr>
<td>splitIntoClades</td>
<td>44</td>
</tr>
<tr>
<td>terminalSisters</td>
<td>44</td>
</tr>
<tr>
<td>tipHeights</td>
<td>45</td>
</tr>
<tr>
<td>traitRate</td>
<td>46</td>
</tr>
<tr>
<td>trimEnds</td>
<td>47</td>
</tr>
<tr>
<td>unlistFirstLevel</td>
<td>48</td>
</tr>
<tr>
<td>write.fas</td>
<td>48</td>
</tr>
</tbody>
</table>

**Index**

**Description**

This package presents a set of functions that were formerly included in the *phyloch* package and which wrap popular phylogenetic software for sequence alignment, masking of sequence alignments, and estimation of phylogenies and ancestral character states.
**Details**

Package:    ips  
Type:       Package  
Version:    0.0.11  
Date:       2019-06-20  
License:    GPL (>= 2)

There are several functions for reading and writing DNA sequences in FASTA, PHYLIP, and NEXUS format: `read.fas`, `read.phy`, `read.nex`, `write.fas`, `write.phy`, and `write.nex`. Some functions are available for integrating BEAST with R. XML input files for BEAST can be generated with `rbeauti`. Two functions are designed to read TreeAnnotator output: `read.beast` will render an object of class `phylo` with additional node statistics appended as list elements. These additional node statistics will be lost by the subsequent use of `ladderize` or `rotate` (or similar functions that change the ordering of internal nodes). `read.beast.table` also parses the TreeAnnotator output, but returns a matrix of node statistics. This package itself does not implement techniques for phylogenetic analyses, but provides a series of wrappers for commonly used software packages. Sequence alignment can be done with the `mafft` and `prank`; cleaning of sequences with `gblocks` and `aliscore`. The function `raxml` and `mrbayes` are intended for phylogenetic tree search. Running `mrbayes` with argument `run = FALSE` can be used to create MrBayes-executable NEXUS files. Finally, wrappers is provided for Multistate in the BayesTraits package (see `multistateML` and `multistateMCMC`). Several plotting functions (`HPDbars`, `clade.bars`, `box.clades`, `box.tips`, `tip.color`, `edge.color` have been moved to the `viper` package.

**Author(s)**

Natalie Cusimano, Christoph Heibl, Franz-Sebastian Krah, Maintainer: Christoph Heibl (<christoph.heibl@gmx.net>)

**See Also**

`ape`
Arguments

- **x**
  - DNA sequences of class `DNAbin`.

- **gaps**
  - A vector of mode "character" indicating how gaps shall be treated: as "5state" or as "ambiguous".

- **w**
  - An integer giving the size of the sliding window.

- **r**
  - An integer giving the number of random pairwise sequence comparisons; defaults to $4 \times N$.

- **t**
  - *Not yet implemented.*

- **l**
  - *Not yet implemented.*

- **s**
  - *Not yet implemented.*

- **o**
  - A vector of mode "character" containing outgroup taxon names.

- **exec**
  - A character string, giving the path to the Aliscore script.

Value

A matrix of class "DNAbin".

Note

This function was developed with ALISCORE version 2.

References


Aliscore website: https://www.zfmk.de/en/research/research-centres-and-groups/aliscore

See Also

- `mafft` and `prank` for multiple sequence alignment; `gblocks` for another alignment masking algorithm.

Examples

```r
data(ips.28S)
## Not run: aliscore(ips.28S)
```
**Description**

code.simple.gaps takes an aligned DNA sequence matrix and codes the simple gaps, i.e. gaps that do not overlap with other gaps. The gapped positions are excluded from the matrix and the coded gap characters are appended to the matrix.

**Usage**

code.simple.gaps(x, append = TRUE)

**Arguments**

- **x**
  - An object of class `DNAbin`.
- **append**
  - Logical.

**Value**

An object of class `DNAbin`.

**Author(s)**

Christoph Heibl

**References**


**See Also**

deleteGaps, deleteEmptyCells, trimEnds

collapseUnsupportedEdges

**Description**

Given a set of node support values (e.g., bootstrap proportions, posterior probabilities) and a certain threshold, all edges receiving less support than the threshold will be collapsed.
Usage

collapseUnsupportedEdges(phy, value = "node.label", cutoff)

Arguments

phy An object of class phylo.
value A character string giving the name of the list element that contains the support values; default is "node.label".
cutoff A numeric value giving the threshold below which edges will be collapsed.

Value

An object of class phylo.

Examples

## phylogeny of bark beetles
data(ips.tree)
## non-parametric bootstrap proportions (BP)
ips.tree$node.label
## collapse clades with < 70 BP
tr <- collapseUnsupportedEdges(ips.tree, "node.label", 70)
## show new topology
par(mfrow = c(1, 2))
plot(ips.tree, no.margin = TRUE)
modelabels(ips.tree$node.label, cex = .5, frame = "n", adj = c(0, .5))
plot(tr, no.margin = TRUE)

---

del.miss  Delete Missing Data from DNA Sequences

Description

Remove gaps ("-") and/or missing and ambiguous data ("N", "?") from a sample of DNA sequences.

Usage

del.miss(x)

Arguments

x A matrix, a list, or a vector of class DNAbin containing the DNA sequences.

Value

A list or a vector of class DNAbin.
deleteEmptyCells  

*Delete Spurious Rows and Columns from DNA Alignments*

**Description**

After subsetting (see e.g. DNAbin), DNA sequence alignments can contain rows and columns that consist entirely of missing and/or ambiguous character states. deleteEmptyCells will delete all such rows (taxa) and columns (characters) from a DNA sequence alignment.

**Usage**

```r
deleteEmptyCells(DNAbin, margin = c(1, 2), nset = c("-", "n", "?"), quiet = FALSE)
```

**Arguments**

- **DNAbin**: An object of class DNAbin.
- **margin**: A vector giving the subscripts the function will be applied over: 1 indicates rows, 2 indicates columns, and c(1, 2) indicates rows and columns.
- **nset**: A vector of mode character; rows or columns that consist only of the characters given in nset will be deleted from the alignment. Allowed are "-", "?", "n", "b", "d", "h", "v", "r", "y", "s", "w", "k", and "m".
- **quiet**: Logical: if set to TRUE, screen output will be suppressed.

**Details**

For faster execution, deleteEmptyCells handles sequences in ape's bit-level coding scheme.

**Value**

An object of class DNAbin.

**References**


**See Also**

trimEnds, deleteGaps
Examples

```r
# COX1 sequences of bark beetles
data(ips.cox1)
# introduce completely ambiguous rows and columns
x <- as.character(ips.cox1[1:6, 1:60])
x[3, ] <- rep("n", 60)
x[, 20:24] <- rep("-", 6)
x <- as.DNAbin(x)
image(x)
# delete those rows and columns
x <- deleteEmptyCells(x)
image(x)
```

---

**deleteGaps**

*Remove Gap Positions From DNA Sequences*

### Description

Remove indel positions (or gaps) from a DNA sequence alignment. For faster execution, `deleteGaps` handles sequences in *ape*’s bit-level coding scheme.

### Usage

```r
deleteGaps(x, gap.max = nrow(x) - 4)
```

### Arguments

- `x` An object of class *DNAbin*.
- `gap.max` An integer, which gives the maximum number of gap characters ("-") that will be tolerated at any given alignment position (column). Only values between 0 and `nrow(x) - 4` make sense phylogenetically.

### Details

The default, `nmax = nrow(x) - 4`, removes all those positions from the alignment, which contain at least four non-gap characters, which is the minimum number of sequences needed to produce a non-trivial unrooted topology. All gaps will be excluded by selecting `nmax = 0` and half of all gaps with `nmax = nrow(x) / 2`.

In contrast, `del.gaps` removes all gap characters from the alignment, so most probably the result will not be a set of sequences of equal length and the matrix will be coerced to a list.

### Value

An object of class *DNAbin*.

### See Also

- `code.simple.gaps` for coding of simple gaps, `del.gaps` for removal of all gap symbols from an alignment, `gblocks` and `aliscore` for more sophisticated methods of cleaning/masking alignments.
descendants

Descendants of an Internal Node in a Phylogeny

Description
For any given internal node of a phylogeny, the function returns a vector containing the node numbers descending from that node.

Usage
descendants(phy, node, type = "t", ignore.tip = TRUE, labels = FALSE)

Arguments
- **phy**: an object of class `phylo`.
- **node**: an integer giving the number of the internal node.
- **type**: a character string, may be "daughter", "internal", "terminal", "all", or any unambiguous abbreviation of these.
- **ignore.tip**: logical, if `ignore.tip = FALSE`, the function will issue an error when node is not internal, otherwise the number of the corresponding terminal node will be returned.
- **labels**: logical, determines if node labels are returned instead of node number, currently ignored unless `type = "t"`.

Value
A vector containing terminal node numbers or tip labels.

Author(s)
Christoph Heibl

See Also
- `sister`, `noi`

Examples
```r
# generate a random tree with 12 terminal and 11 internal nodes:
tree <- rtree(12)

# get the descendants of internal node 15:
x <- descendants(tree, 15)
```
**DNAbin2index**  
*Conversion of DNAbin to Index*

**Description**
Extract the indices of non-empty positions in a sample of DNA sequences to

**Usage**
```
DNAbin2index(x)
```

**Arguments**
- `x` A matrix of class `DNAbin`.

**See Also**
`index2DNAbin`

**eoi**  
*Identification of Stem-Lineage-Edges and MRCAs*

**Description**
`noi` (node of interest) identifies the most recent common ancestor (MRCA) and `eoi` (edge of interest) its subtending stem-lineage edge of one or more sets of taxa/tips.

**Usage**
```
eoi(phy, node, group, regex = FALSE, stem = FALSE, monophyletic = FALSE)
noi(phy, group, regex = FALSE, stem = FALSE, monophyletic = FALSE)
```

**Arguments**
- `phy` An object of class `phylo`.
- `node` A vector of mode "numeric" giving the nodes numbers of the nodes whose subtending stem-lineages will be identified.
- `group` A vector or list of vectors of mode character specifying the taxon set(s). Will be ignored if `node` is given.
- `regex` A logical, if `regex = TRUE`, taxon sets are matched to the tip labels as regular expressions of the form "taxon1|taxon2"; otherwise strings will be matched exactly (see `which`).
- `stem` Logical, ...
- `monophyletic` Logical, ...
Value

A vector of mode "numeric" containing node numbers.

See Also

`mrca`, `descendants` for the contrary operation to `noi`.

Examples

```r
# molecular phylogeny of Ips bark beetles
# ---------------------------------------
data(ips.tree)
ips.tree <- ladderize(ips.tree)
ips.tree <- fixNodes(ips.tree)
clade1 <- descendants(ips.tree, 44, labels = TRUE)
mrca <- noi(ips.tree, clade1)
stem_lineage <- eoi(ips.tree, mrca)
ecol <- rep("black", Nedge(ips.tree))
ecol[stem_lineage] <- "red"
plot(ips.tree, no.margin = TRUE, edge.color = ecol)
modelabels(node = mrca, pch = 22, col = "blue")
#gen <- sapply(viperidae$tip.label, function(x) unlist(strsplit(x, "."))[1])
#tax <- data.frame(genus = gen, species = viperidae$tip.label, row.names = NULL)

# group can be a list
# -------------------
#myclades <- split(tax$species, tax$genus)
#nds <- noi(viperidae, myclades)
#plot(viperidae)
#nodeInfo(nds)

# group might contain tip numbers
# -------------------------------
#group <- list(c(17, 22), c(13, 1))
#plot(viperidae)
#append2tips(phy, tips = unlist(group), pch = 19)
#nds <- noi(viperidae, myclades)
#nodeInfo(nds)

# the 'group' argument can also take regular expressions
# ------------------------------------------------------
#rex <- "aspis"
#node <- noi(viperidae, rex, regex = TRUE)
#plot.phylo(viperidae, tip.color = 0, edge.color = 0)
#box.clades(viperidae, nodes = node, col = "#D2A6A7", align = "all")
#plot.phylo.upon(viperidae)
#nodeLabels(node = node, pch = 21, cex = 1.2, col = "red", bg = "#D2A6A7")

# if the 'group' argument is a list of elements of length 2,
# n = length(group) nodes of interest will be returned
# -----------------------------------------------
#group <- list(
# c("Vipera_berus", "Vipera_ursinii"),
# c("Vipera_aspis_ssp._aspis", "Vipera_latastei"),
# c("Vipera_ammodytes_ssp._ammodytes",
#   "Vipera_ammodytes_ssp._montandoni"),
# c("Macrovipera_lebetina", "Vipera_wagneri")
#)
#clades <- noi(viperidae, group)
#plot.phylo(viperidae, tip.color = 0, edge.color = 0)
#box.clades(viperidae, nodes = clades, col = c("FFFFA5", "#D2A6A7",
#   "#A7D2A5", "#A5A6D2"), align = "all")
#plot.phylo.upon(viperidae)

define the function to establish the standard numbering of terminal and internal nodes in phylogenies represented as objects of class phylo.

**Usage**

fixNodes(phy)

**Arguments**

phy An object of class phylo.

**Details**

When reading phylogenetic trees from a NEXUS file that contains a translate section, it can happen that the terminal nodes (tips, leaves) of the corresponding phylo object are not numbered consecutively, which can be a problem in some downstream applications. You can use fixNodes to get the correct order of terminal node numbers.

fixNodes is also intended to re-establish the standard numbering of internal nodes and reorder all node value elements (e.g. node.label, posterior, ...) if a phylo object has been modified by either root, ladderize, or rotate.

**Value**

An object of class phylo.

**Note**

fixNodes has been completely rewritten for ips version 1.0-0. It should now run absolutely stable and is much quicker. Nevertheless, I recommend checking carefully the results of fixNodes, until the function has been tested by a number of users. Then this comment will be removed.
**forceEqualTipHeights**

**Author(s)**

Christoph Heibl

**See Also**

*read.tree, read.nexus, read.beast* for reading trees in NEWICK and NEXUS format; *ladderize* and *rotate* for tree manipulation; *node.support* for plotting node support values has been moved to package *viper*.

---

### forceEqualTipHeights  Equal Tip Heights

**Description**

Modify terminal edge lengths to create "exactly" (see Details) equal tip heights (sum of edge lengths from root to tip)

**Usage**

```r
forceEqualTipHeights(phy, baseline = "mean")
```

**Arguments**

- `phy`  
  An object of class *phylo*.
- `baseline`  
  A character string giving a function to calculate the baseline tip height, e.g. "min", "max" or "mean".

**Details**

What is "exactly" equal depends on the precision of the system (.Machine); in any case the resulting phylogeny will pass *is.ultrametric* with default arguments.

**Value**

An object of class *phylo* with changed terminal edge lengths.

**Note**

*forceEqualTipHeights* is only intended to correct small rounding errors in edge lengths, not to make an additive phylogeny ultrametric. For the latter, see e.g. *chronos*.

**See Also**

*tipHeights*
Description

Provides a wrapper to Gblocks, a computer program written in ANSI C language that eliminates poorly aligned positions and divergent regions of an alignment of DNA or protein sequences. Gblocks selects conserved blocks from a multiple alignment according to a set of features of the alignment positions.

Usage

```
gblocks(x, b1 = 0.5, b2 = b1, b3 = ncol(x), b4 = 2, b5 = "a",
        target = "alignment", exec)
```

Arguments

- **x**: A matrix of DNA sequences of classes `DNAbin`.
- **b1**: A real number, the **minimum number of sequences for a conserved position** given as a fraction. Values between 0.5 and 1.0 are allowed. **Larger** values will **decrease** the number of selected positions, i.e. are more **conservative**. Defaults to 0.5
- **b2**: A real number, the **minimum number of sequences for a flank position** given as a fraction. Values must be equal or larger than b1. **Larger** values will **decrease** the number of selected positions, i.e. are more **conservative**. Defaults to 0.5
- **b3**: An integer, the **maximum number of contiguous nonconserved positions**; any integer is allowed. **Larger** values will **increase** the number of selected position, i.e. are **less conservative**. Defaults to the number of positions in the alignment.
- **b4**: An integer, the **minimum length of a block**, any integer equal to or bigger than 2 is allowed. **Larger** values will **decrease** the number of selected positions, i.e. are more conservative. Defaults to 2.
- **b5**: A character string indicating the **treatment of gap positions**. Three choices are possible. 1. "n": **No** gap positions are allowed in the final alignment. All positions with a single gap or more are treated as a gap position for the block selection procedure, and they and the adjacent nonconserved positions are eliminated. 2. "h": **Only** positions where **50% or more** of the sequences have a gap are treated as a gap position. Thus, positions with a gap in less than 50% of the sequences can be selected in the final alignment if they are within an appropriate block. 3. "a": **All** gap positions can be selected. Positions with gaps are not treated differently from other positions (default).
- **target**: A vector of mode "character" giving the output format: "alignment" will return the alignment with only the selected positions, "index" will return the indices of the selected position, and "score" will provide a score for every position in the original alignment (0 for excluded, 1 for included).
- **exec**: A character string indicating the path to the GBLOCKS executable.
Details

Explanation of the routine taken from the Online Documentation: First, the degree of conservation of every positions of the multiple alignment is evaluated and classified as nonconserved, conserved, or highly conserved. All stretches of contiguous nonconserved positions bigger than a certain value (b3) are rejected. In such stretches, alignments are normally ambiguous and, even when in some cases a unique alignment could be given, multiple hidden substitutions make them inadequate for phylogenetic analysis. In the remaining blocks, flanks are examined and positions are removed until blocks are surrounded by highly conserved positions at both flanks. This way, selected blocks are anchored by positions that can be aligned with high confidence. Then, all gap positions -that can be defined in three different ways (b5)- are removed. Furthermore, nonconserved positions adjacent to a gap position are also eliminated until a conserved position is reached, because regions adjacent to a gap are the most difficult to align. Finally, small blocks (falling below the limit of b4) remaining after gap cleaning are also removed.

Value

A matrix of class "DNAbin"

Note

gblocks was last updated and tested to work with Gblocks 0.91b. If you have problems getting the function to work with a newer version of Gblocks, please contact the package maintainer.

References


Gblocks website: http://molevol.cmima.csic.es/castresana/Gblocks.html

See Also

mafft and prank for multiple sequence alignment; aliscore for another alignment masking algorithm.

Examples

data(ips.28S)
## Not run: gblocks(ips.28S)
**index2DNAbin  
*Conversion of Index to DNAbin*

**Description**

Use indices of non-empty positions to convert a list of DNA sequences into a matrix.

**Usage**

```r
index2DNAbin(DNAbin, index)
```

**Arguments**

- `DNAbin` A list of class `DNAbin`.
- `index` A list of integers containing the indices of base positions.

**See Also**

`DNAbin2index`

---

**ips.16S  
*Bark Beetle 16S Sequences* 

**Description**

This DNA alignment contains 376 positions of 42 sequences of 16S ribosomal DNA of the bark beetle genera *Ips*, *Orthotomicus*, and *Pityogenes* (Scolytinae, Curculionidae, Coleoptera).

**Usage**

```r
data(ips.16S)
```

**Format**

The sequences are stored in binary format (see `DNAbin`).

**Source**

The sequences were downloaded and assembled from the Nucleotide repository at GenBank on February 8, 2014.

**References**


**Examples**

```r
data(ips.16S)
```
ips.28S  

*Bark Beetle 28S Sequences*

**Description**

This DNA alignment contains 562 positions of 28 sequences of 28S ribosomal DNA of the bark beetle genus *Ips* (Scolytinae, Curculionidae, Coleoptera).

**Usage**

data(ips.28S)

**Format**

The sequences are stored in binary format (see *DNAbin*).

**Source**

The sequences were downloaded and assembled from the Nucleotide repository at GenBank on February 8, 2014.

**References**


**Examples**

data(ips.28S)

ips.cox1  

*Bark Beetle COX1 Sequences*

**Description**

This DNA alignment contains 770 positions of 26 sequences of cox1 of the bark beetle genera *Ips*, *Orthotomicus*, and *Pityogenes* (Scolytinae, Curculionidae, Coleoptera).

**Usage**

data(ips.cox1)

**Format**

The sequences are stored in binary format (see *DNAbin*).
Source

The sequences were downloaded and assembled from the Nucleotide repository at GenBank on February 8, 2014.

References


Examples

data(ips.cox1)

ips.tree

Ips Phylogeny

Description

Phylogentic tree of bark beetles (genus *Ips*).

Usage

data(ips.tree)

Format

The format is: List of 5 $ edge : int [1:72, 1:2] 38 39 39 40 41 42 42 43 44 45 ... $ Nnode : int 36 $ tip.label : chr [1:37] "Ips_acuminatus" "Ips_duplicatus" "Ips_integer" "Ips_plastographus" ... $ edge.length: num [1:72] 0.2806 0.0727 0.0295 0.0097 0.021 ... $ node.label : chr [1:36] "" "100" "21" "12" ... - attr(*, "class")= chr "phylo" - attr(*, "order")= chr "cladewise"

Examples

data(ips.tree)
plot(ips.tree)
**mafft**  

*Sequence Alignment with MAFFT*

---

**Description**

This function is a wrapper for MAFFT and can be used for (profile) aligning of DNA and amino acid sequences.

**Usage**

```r
mafft(x, y, add, method = "auto", maxiterate = 0, op = 1.53, ep = 0, gt, options, thread = -1, exec, quiet, file)
```

**Arguments**

- **x**: An object of class `DNAbin` or `AAbin`.
- **y**: An object of class `DNAbin` or `AAbin`, if given both `x` and `y` are preserved and aligned to each other ("profile alignment").
- **add**: A character string giving the method used for adding `y` to `x`: "add", "addprofile" (default), or any unambiguous abbreviation of these.
- **method**: A character string giving the alignment method. Available accuracy-oriented methods for less than 200 sequences are "localpair", "globalpair", and "genafpair"; "retree 1" and "retree 2" are for speed-oriented alignment. The default is "auto", which lets MAFFT choose an appropriate alignment method.
- **maxiterate**: An integer giving the number of cycles of iterative refinement to perform. Possible choices are 0: progressive method, no iterative refinement (default); 2: two cycles of iterative refinement; 1000: at most 1000 cycles of iterative refinement.
- **op**: A numeric giving the gap opening penalty at group-to-group alignment; default 1.53.
- **ep**: A numeric giving the offset value, which works like gap extension penalty, for group-to-group alignment; default 0.0, but 0.123 is recommended if no long indels are expected.
- **gt**: An object of class `phylo` that is to be used as a guide tree during alignment.
- **options**: A vector of mode character specifying additional arguments to MAFFT, that are not included in `mafft` such as, e.g., --adjustdirection.
- **thread**: Integer giving the number of physical cores MAFFT should use; with `thread = -1` the number of cores is determined automatically.
- **exec**: A character string giving the path to the MAFFT executable including its name, e.g. something like `/user/local/bin/mafft` under UNIX-alikes.
- **quiet**: Logical, if set to `TRUE`, mafft progress is printed out on the screen.
- **file**: A character string indicating the filename of the output FASTA file; if this is missing the the alignment will be returned as matrix of class `DNAbin` or `AAbin`. 

---
Details

"localpair" selects the L-INS-i algorithm, probably most accurate; recommended for <200 sequences; iterative refinement method incorporating local pairwise alignment information.

"globalpair" selects the G-INS-i algorithm suitable for sequences of similar lengths; recommended for <200 sequences; iterative refinement method incorporating global pairwise alignment information.

"genafpair" selects the E-INS-i algorithm suitable for sequences containing large unalignable regions; recommended for <200 sequences.

"retree 1" selects the FFT-NS-1 algorithm, the simplest progressive option in MAFFT; recommended for >200 sequences.

"retree 2" selects the FFT-NS-2 algorithm that uses a second iteration of alignment based on a guide tree computed from an FFT-NS-1 alignment; this is the default in MAFFT; recommended for >200 sequences.

Value

A matrix of class "DNAbin" or "AAbin".

Note

mafft was last updated and tested to work with MAFFT 7.205. If you have problems getting the function to work with a newer version of MAFFT, please contact the package maintainer.

References


http://mafft.cbrc.jp/alignment/software/index.html

See Also

read.fas to import DNA sequences; prank for another alignment algorithm; gblocks and aliscore for alignment cleaning.
mafft.merge

Profile Alignment with MAFFT

Description

Merge two or more DNA or amino acid sequence alignments by profile alignment with MAFFT.

Usage

mafft.merge(subMSA, method = "auto", gt, thread = -1, exec, quiet = TRUE)

Arguments

- subMSA: A list of objects of class "DNAbin" or "AAbin".
- method: A character string giving the alignment method. Available accuracy-oriented methods for less than 200 sequences are "localpair", "globalpair", and "genafpair"; "retree 1" and "retree 2" are for speed-oriented alignment. The default is "auto", which lets MAFFT choose an appropriate alignment method.
- gt: An object of class phylo that is to be used as a guide tree during alignment.
- thread: Integer giving the number of physical cores MAFFT should use; with thread = -1 the number of cores is determined automatically.
- exec: A character string giving the path to the MAFFT executable including its name, e.g. something like /user/local/bin/mafft under UNIX-alikes.
- quiet: Logical, if set to TRUE, mafft progress is printed out on the screen.

Value

An object of class "DNAbin" or "AAbin".

mrbayes

Bayesian MCMC Tree Search with MrBayes

Description

Provides a wrapper for Bayesian phylogenetic tree search through MrBayes (Huelsenbeck & Ronquist, 2001; Ronquist & Huelsenbeck, 2003).

Usage

mrbayes(x, file = "", lset, prset, mcmc, unlink, constraint, burnin = 10, contype = "allcompat", exec, run = FALSE)
Arguments

x  An object of class DNAbin in the case of mrbayes.

file  A character string, giving the name of the MrBayes input file.

lset  A list as returned by mrbayes.lset containing the parameter settings of the model of molecular evolution.

prset  A list as returned by mrbayes.prset containing the parameter setting for the prior distributions.

mcmc  A list as returned by mrbayes.mcmc containing the parameter setting for the Markov chain Monte Carlo (MCMC).

unlink  constraint

burnin  An integer; the number of samples from the MCMC to be discarded prior to further analysis.

contype  A character string; the type of consensus tree calculated from the posterior distribution of trees: either "halfcompat" (majority-rule consensus tree) or "allcombat" (strict consensus tree).

exec  A character string giving the full path of the MrBayes program.

run  Logical; run = FALSE will only print the NEXUS file, run = TRUE will also start the MCMC runs, if exec is correctly specified.

Details

mrbayes was last updated and tested with MrBayes v3.2.2 under R 3.1.0 on a x86_64-apple-darwin10.8.0 (64-bit) platform. It is intended to offer a simply parameterized building block for larger scripts.

Value

None; a NEXUS file with MrBayes block is written to a file and, if run = TRUE, the MCMC runs in MrBayes are started.

Author(s)

Christoph Heibl

References


MrBayes website: http://mrbayes.sourceforge.net/

See Also

mafft and prank for sequence alignment; raxml for maximum likelihood tree search.
Examples

data(ips.cox1)
x <- ips.cox1[, 100:140] # tiny alignment
mrbayes(x, file = "", mcmc = mrbayes.mcmc(ngen = 100), run = FALSE)

## Not run:
library(phangorn)
tree <- rtree(10)
Y1 <- simSeq(tree, l = 20)
Y2 <- simSeq(tree, l = 20, type = "USER", levels=c("0", "1"))
Y <- cbind(as.character(Y1), as.character(Y2))
mrbayes(Y, file = "", run = FALSE)

## End(Not run)

---

mrbayes.lset  Model Settings for MrBayes

Description

Set model parameters for mrbayes.

Usage

mrbayes.lset(..., partition)

Arguments

... arguments in tag = value form, or a list of tagged values. The tags must come from the names of model parameters described in the ‘Model Parameters’ section.

partition

Value

a list containing a subset (including the empty and the full set) of model parameters.

Model Parameters

nucmodel  "4by4", "doublet", "codon", or "protein".
nst  1, 2, 6, or "mixed".
code  "universal", "vertmt", "mycoplasma", "yeast", "ciliates", or "metmt".
ploidy  "haploid", "diploid", or "zlinked".
rates  "equal", "gamma", "propinv", "invgamma", or "adgamma".
### mrbayes.mcmc

**Description**

Set Markov chain Monte Carlo (MCMC) parameters for `mrbayes`.

**Usage**

```
mrbayes.mcmc(...)```

**Examples**

```r
## F81
mrbayes.lset(nst = 2)

## GTR + Gamma
mrbayes.lset(nst = 6, rates = "gamma")

## GTR + Gamma + I
mrbayes.lset(nst = 6, rates = "invgamma")
```

**Author(s)**

Christoph Heibl

**References**


---

**ngammacat** 1-24

**nbetacat** 1-24

**omegavar** "equal", "ny98", or "m3".

**covarion** "no" or "yes".

**coding** "all", "variable", "noabsencesites", or "nopresencesites".

**parsmodel** "no" or "yes".
Arguments

... arguments in tag = value form, or a list of tagged values. The tags must come from the names of MCMC parameters described in the ‘MCMC Parameters’ section.

Value

a list containing a subset (including the empty and the full set) of model parameters.

MCMC Parameters

ngen "NUMERIC"
nruns "NUMERIC"
nchains "NUMERIC"
temp "NUMERIC"
swapfreq "NUMERIC"
nswaps "NUMERIC"
samplefreq "NUMERIC"
printfreq "NUMERIC"
printall "yes" or "no"
printmax "NUMERIC"
mcmcdiagn "yes" or "no"
diagnfreq "NUMERIC"
diagnstat "avgsddev" or "maxstddev"
minpartfreq "NUMERIC"
allchains "yes" or "no"
allcomps "yes" or "no"
relburnin "yes" or "no"
burnin "NUMERIC"
burninfrac "NUMERIC"
stoprule "yes" or "no"
stopval "NUMERIC"
savetrees "yes" or "no"
checkpoint "yes" or "no"
checkfreq "NUMERIC"
startparams "current" or "reset"
starttree "current", "random", or "parsimony"
nperts "NUMERIC"
data "yes" or "no"
ordertaxa "yes" or "no"
append "yes" or "no"
autotune "yes" or "no"
tunefreq "NUMERIC"
Note

The parameters reweight and filename cannot be set via mrbayes.mcmc.

Author(s)

Christoph Heibl

References


See Also

mrbayes.lset to set model parameters, mrbayes.prset to set prior distributions, and mrbayes to run MrBayes locally or prepare input files for a computer cluster.

Examples

mrbayes.mcmc()

mrbayes.prset

Set Priors for MrBayes

Description

Set prior distributions for mrbayes.

Usage

mrbayes.prset(...)

Arguments

... arguments in tag = value form, or a list of tagged values. The tags must come from the names of prior distribution parameters described in the ‘Prior Distribution Parameters’ section.

Value

a list of length zero (see ‘Note’).
Prior Distribution Parameters

traitiopr
revmatpr
aamodelpr
aarevmatpr
omegapr
ny98omega1pr
ny98omega3pr
m3omegapr
codoncatfreqs
statefreqpr
shapepr
ratecorrpr
pinvarpr
covswitchpr
symdirihyperpr
topologypr
brlenspr
clockvarpr
igrvarpr

Note
This function currently returns an empty set of prior distribution parameters, i.e., you cannot change the MrBayes default parameters.

Author(s)
Christoph Heibl

References


See Also

*mrbayes.lset* to set model parameters, *mrbayes.mcmc* to set parameters of the Markov chain Monte Carlo (MCMC), and *mrbayes* to run MrBayes locally or prepare input files for a computer cluster.
Examples

mrbayes.prset()

Description

These functions provide wrappers to BayesMultiState in the BayesTraits package written by Mark Pagel and Andrew Meade.

Usage

multistateML(phy, traits, model = "ARD", anc.states = TRUE,
             path = "/Applications/BayesTraits", dir = NULL)

multistateMCMC(phy, traits, model = "ARD", anc.states = TRUE,
                rd = 2, rjhp = NULL, fixNodes = NULL, it = 1e+05, bi = 10000,
                sa = 1000, path = "/Applications/BayesTraits", dir = NULL)

Arguments

phy an object of class phylo.
traits a data.frame with two columns. The first column contains the taxon labels, the second column contains the character states.
model
anc.states either logical or a list, the latter containing the tip labels of those internal nodes, for which the likelihood of ancestral character states should be estimated.
rd a real number, giving the RateDev parameter, i.e., the deviation of the normal distribution, that changes to the rates are drawn from. Should be set such that acceptance of the rate parameters is about 0.2.
rjhp a character string giving the details of priors and hyperpriors for the reversible jump MCMC (rjMCMC). If left NULL, a conventional MCMC is used. In order to use the rjMCMC, you must specify the distribution of the prior and the interval of the uniform hyperprior distribution that seeds it. For example, exp 0 30 specifies an exponential distribution seeded from a uniform distribution on the interval 0 to 30, and gamma 0 10 0 10 specifies a gamma prior with its mean and standard deviation seeded from uniform distributions on the interval 0 to 10.
fixNodes a list giving fixed character states of certain internal nodes. This argument corresponds to the fossil command in the MultiState manual.
it numeric, sets the number of iterations to run the MCMC for.
bi numeric, sets the number of iterations of the MCMC that will be discarded as burn-in.
sa numeric, sets the the sample period in the MCMC.
path
a character string giving the path to executables in the BayesTraits package.

dir
a character string giving a directory name where the input and output files will be stored. The directory will be created by \texttt{multistateML} and must not exist already. If \texttt{dir = NULL} (default) input and output is written to the working directory (thereby overwriting existing output).

Author(s)

Christoph Heibl

References

The BayesTraits manual: \url{http://www.evolution.rdg.ac.uk/Files/BayesTraits-V1.0-Manual.pdf}.


See Also

ace

\begin{verbatim}
neighborsPairs  Neighboring Nodes in a Minimum Spanning Tree
\end{verbatim}

Description

Finds all pairs of adjacent nodes, i.e. nodes separated by only one edge, in a minimum spanning tree

Usage

\texttt{neighborsPairs(mst)}

Arguments

\begin{verbatim}
  mst  An object of class \texttt{mst}.
\end{verbatim}
ntip

Numbers of Tips of (Sub)trees

Description
Counts the number of tips of a given clade of a phylogenetic tree.

Usage
ttip(phy, node)

Arguments
phy An object of class phylo.
node An integer given the number of an internal node.

Value
An integer giving the number of tips.

Examples
set.seed(1234)
tr <- rtree(12)
plot(tr); nodelabels()
ntip(tr, 16)

partitionfinder

PartitionFinder

Description
Provides a wrapper to the PartitionFinder software.

Usage
partitionfinder(alignment, user.tree, branchlengths = "linked",
models = "all", model.selection = "BIC", search = "greedy",
exec = "/Applications/PartitionFinderV1.1.1_Mac/PartitionFinder.py")
pathd8

Arguments

alignment A
user.tree A
branchlengths A
models A
model.selection A
search A
exec A character string giving the path to the executable (python script).

References


Description

This function is a wrapper for PATHd8 and can be used for phylogenetic dating, especially of large trees.

Usage

pathd8(phy, exec = "/Applications/PATHd8/PATHd8", seql, calibration)

Arguments

phy An object of class phylo.
exec A character string giving the path to the PATHd8 program.
seql sequence length of alignment
 calibration A data frame with 4 columns and as many rows as calibration points. Columns are: taxon 1; taxon 2; one of c("minage", "maxage", "fixage"); age.

Value

tree list of ultrametric trees returned from PATHd8 of class phylo. First tree is PATHd8 chronogram, which is a calibrated ultrametric tree. Second is a PATH tree, which is a ultrametric tree without calibration.
phylo2mafft

Convert Trees for MAFFT

Description

Converts a phylogenetic tree of class "phylo" to a format usable as a guide tree by MAFFT. This function is called internally by mafft.

Usage

phylo2mafft(phy, file)

Arguments

phy A phylogenetic tree of class phylo.

file A character string giving a filename. May be missing, in which case the results are only printed on the screen.

Value

A matrix coding the MAFFT-formatted tree, as a side effect the same matrix is written to file.
References

The MAFFT website: http://mafft.cbrc.jp/alignment/software/index.html

See Also

mafft for an interface to MAFFT.

---

**phylo2mst**  
*Conversion from PHYLO to MST Object*

**Description**

Converts a phylogenetic tree (class `phylo`) into a minimum spanning tree (class `mst`).

**Usage**

phylo2mst(phy)

**Arguments**

- `phy`  
  An object of class `phylo`.

**Details**

The current version of `phylo2mst` does not handle polytomies and does not incorporate branch length information. Note that topological information is lost during the conversion.

**Examples**

```r
phy <- rtree(12)
plot(phy)
mst <- phylo2mst(phy)
plot(mst)
```

---

**pis**  
*Number of Potentially-Informative Sites*

**Description**

This function returns the number or positions of potentially-informative (parsimony-informative, phylogenetically-informative) sites in DNA sequence alignment.

**Usage**

pis(x, what = "fraction", use.ambiguities = FALSE)
prank

Arguments

x  
An object of class DNAbin.

what  
Either of "absolute", "fraction", or "index", which will return the absolute number, the relative number or the indeces of the potentially-informative sites.

use.ambiguities  
Not yet available.

Value

Numeric (depending on what, the number, fraction, or indices of potentially-informative nucleotide sites).

Author(s)

Christoph Heibl

Examples

# example data:
# --------------
data(ips.16S)

# number of potentially-informative sites:
# ----------------------------------------
pis(ips.16S, what = "abs")

# proportion of potentially-informative sites:
# --------------------------------------------
pis(ips.16S, what = "frac")

# indeces of potentially-informative sites:
# ----------------------------------------
pis(ips.16S, what = "ind")

Description

DNA sequence Alignment Using the program PRANK.

Usage

prank(x, outfile, guidetree = NULL, gaprate = 0.025, gapext = 0.75, path)
Arguments

x  an object of class DNAbin.
outfile  a character string giving a name for the output file.
guidetree  an object of class phylo to be used as guidetree in alignment.
gaprate  numeric giving the gap opening rate; defaults to 0.025.
gapext  numeric giving the gap extension penalty; defaults to 0.75.
path  a character string indicating the path to the PRANK executable.

Value

matrix of class "DNAbin"

Note

prank was last updated and tested to work with PRANK v. 120814 on Windows XP. If you have problems getting the function to work with a newer version of PRANK, contact the package maintainer.

References

https://www.ebi.ac.uk/research/goldman/software/prank

See Also

read.fas to import DNA sequences; mafft for another alignment algorithm; gblocks and aliscore for alignment cleaning.

Description

Provides an interface to the C program RAXML (see Reference section) for maximum likelihood estimation of tree topology and/or branch lengths, rapid and conventional non-parametric bootstrapping, mapping splits onto individual topologies, and a lot more. See the RAXML manual for details, especially if you are a new user of RAXML.

Usage

raxml(DNAbin, m = "GTRCAT", f, N, p, b, x, k, weights, partitions, outgroup, backbone = NULL, file = paste0("fromR_", Sys.Date()), exec, threads)
Arguments

DNAbin  A matrix of DNA sequences of class DNAbin.

m  A vector of mode "character" defining a model of molecular evolution; currently only GTR model available.

f  A vector of mode "character" selecting an RAxML algorithm analogous to the -f flag (see Detail section and RAxML manual).

N  Either of mode "integer" or "character". Integers give the number of independent searches on different starting tree or replicates in bootstrapping. Alternatively, one of four bootstopping criteria can be chosen: "autoFC", "autoMR", "autoMRE", or "autoMRE_IGN".

p  Integer, setting a random seed for the parsimony starting trees.

b  Integer, setting a random seed for bootstrapping.

x  Integer, setting a random seed for rapid bootstrapping.

k  Logical, if TRUE, the branch lengths of bootstrapped trees are recorded.

weights  A vector of mode "numeric" giving integers to assign individual weights to each column of the alignment. (-a)

partitions  A data frame giving the partitions of the alignment.

outgroup  A vector of mode "character" containing the names of the outgroup taxa.

backbone  A phylo object representing a backbone tree.

file  A vector of mode "character" giving a name for the output files.

exec  A vector of mode "character" giving the path to the directory containing the RAxML executable. The default value will work on Mac OS X if the folder containing the executable is renamed to "RAxML-8.0.3".

threads  Integer, giving the number of parallel threads to use (PTHREADS only).

Details

There are some limitations of this wrapper compared to RAxML run directly from the command line.

1. Only DNA is allowed as data type.
2. Option f can only take a limited number of values (d, a).

RAxML needs the specification of random seeds for parsimony estimation of starting trees and for bootstrap resampling. The corresponding argument names in raxml are identical to the flags used by RAxML (-p, -b, and -x). If you choose not to give any values, raxml will generate a (different) value for each required random seed every time it is called. Be aware that set.seed will work only for p, but not for b or x.

Value

A list with a variable number of elements, depending on the analysis chosen:

"info"  RAxML log file as character string
"bestTree"  MLE of tree
RAxML is a C program and the source code is not contained in this package. This means that in order to run this function you will need to install RAxML yourself. See [http://sco.h-its.org/exelixis/web/software/raxml/](http://sco.h-its.org/exelixis/web/software/raxml/) for the most recent documentation and source code of RAxML. Depending on where you chose to install RAxML, you need to adjust the `exec` argument.

`raxml` was last tested and running fine on Mac OS X with RAxML 8.0.29. Please be aware that calling third-party software from within R is a platform-specific process and I cannot guarantee that `raxml` will behave properly on any system.

**References**

(in chronological order)


**See Also**

`raxml.partitions` to store partitioning information in a data frame suitable for input as partitions argument in `raxml`.

**Examples**

```r
## bark beetle sequences
data(ips.cox1)
data(ips.16S)
data(ips.28S)
ips <- cbind(ips.cox1, ips.16S, ips.28S, fill.with.gaps = TRUE)
exec <- "~/Applications/RAxML-code/standard-RAxML/raxmlHPC-PTHREADS-AVX"
w <- sample(1:5, ncol(ips.cox1), replace = TRUE)

## Not run:
```
# Simple tree search with GTRCAT and GTRGAMMA

```r
tr <- raxml(ips.cox1, f = "d", N = 2, p = 1234, 
              exec = exec) # -1743.528461
```

```r
tr <- raxml(ips.cox1, m = "GTRGAMMA", f = "d", N = 2, p = 1234, 
              exec = exec)
```

# Applying weights to columns

```r
tr <- raxml(ips.cox1, f = "d", N = 2, p = 1234, 
              weights = w, exec = exec) # -1743.528461
```

# Rapid bootstrap

```r
tr <- raxml(ips.cox1, m = "GTRGAMMA", 
              f = "a", N = 10, p = 1234, x = 1234, 
              exec = exec)
```

# Rapid bootstrap with automatic halt

```r
tr <- raxml(ips.cox1, m = "GTRGAMMA", 
              f = "a", N = "autoMRE", p = 1234, x = 1234, 
              exec = exec)
```

## End(Not run)

---

### raxml.partitions

**Partition scheme for RAxML**

**Description**

Given a set of DNA sequence alignments, `raxml.partitions` creates a data frame with partition boundaries that can be input into `raxml`.

**Usage**

`raxml.partitions(...)`

**Arguments**

`...` Two or more DNA sequence alignments of class `DNAbin`.

**Details**

For `raxml.partitions` to make sense, the DNA sequence alignments must be given exactly in the same order in which they are concatenated into a supermatrix (see Examples section). Without any testing, the type of sequences is supposed to be DNA.

**Value**

A data frame with four columns (`type`, `locus`, `begin`, and `end`) and number of rows corresponding to the number of partitions.
XML Input Files for BEAST

Description

This function is intended to prepare XML files for BEAST with R. BEAST uses an MCMC approach to estimate rooted phylogenies from molecular data (Drummond & Rambaut, 2007).

Usage

rbeauti(..., file, template = "standard", taxonset)

Arguments

... one or more object(s) of class DNAbin.
file A connection, or a character string naming the file to write to. If left empty the XML tree will be printed to the screen (see Examples).
template Currently unused.
taxonset A list containing one or more taxon sets.

Details

rbeauti has been completely rewritten to work with BEAST 2. Currently rbeauti offers few options, because the idea is not to create ready-to-use XML file. That can be done conveniently with BEAUti (the BEAST package’s genuine XML generator). Instead, rbeauti is intended to make the definition of large numbers of taxon sets easy. The creation of taxon sets can be done via R scripts and the resulting XML files can be further modified with BEAUti.
Author(s)

Christoph Heibl

References

The BEAST 2 website: http://beast.bio.ed.ac.uk/BEAST_v1.5.x_XML_Reference


See Also

read.beast, read.beast.table

Examples

data(ips.16S)

```r
## define taxon sets
spec <- rownames(ips.16S)
ingroup <- spec[grep("Ips|Orthomotomicus", spec)]
outgroup <- spec[grep("Pityogenes", spec)]
ts <- list(list(id = "ingroup", taxon = ingroup),
         list(id = "outgroup", taxon = outgroup))

## print XML file to screen
rbeauti(ips.16S, taxonset = ts)
```

---

**Description**

Read DNA and amino acid sequences from FASTA, PHILIP, and NEXUS formatted files.

**Usage**

```r
read.fas(x, text)
read.nex(x)
read.phy(x)
```

**Arguments**

- `x` A character string, giving the file name.
- `text` A character string in FASTA format.
Value

An matrix (aligned sequences) or list (unaligned sequences) of class DNAbin or AAbin.

References


See Also

mafft and prank for sequence alignment, gblocks and aliscore for quality check and cleaning of sequence alignments, cbind.DNAbin for concatenation of sequence alignments.

Examples

```r
## bark beetle COX1 sequences
data(ips.cox1)
## create temporary file names
format <- c(".fas", ".phy", ".nex")
fn <- sapply(format, tempfile, pattern = "ips", tmpdir = tempdir())
## write sequences files
write.fas(ips.cox1, fn[".fas"])
write.phy(ips.cox1, fn[".phy"])
write.nex(ips.cox1, fn[".nex"])
## read sequence files
fas <- read.fas(fn[".fas"])
phy <- read.phy(fn[".phy"])
nex <- read.nex(fn[".nex"])
## remove sequence files
unlink(fn)
```

Description

These functions parse chronograms in NEXUS format as produced by TreeAnnotator or output by MrBayes.

Usage

```r
read.mrbayes(file, digits = NULL)
read.beast(file, digits = NULL)
read.starbeast(file)
```
Arguments

file  A character string giving the input file, which must be a TreeAnnotator-generated chronogram in NEXUS format.
digits  NULL or integer, if !is.null(digits) values are rounded to the given integer.

Value

An object of class phylo

Note

read.starbeast currently parses only scalars and ranges; node statistics with more than two values will be deleted and a warning message will be issued. Future version of read.starbeast will hopefully be able to append list or data frames to phylo objects. If you have any opinion or wishes regarding the question of how this exactly should be managed, send me a message.

Author(s)

Christoph Heibl

References

TreeAnnotator: http://beast.bio.ed.ac.uk/TreeAnnotator
Metacomments in NEXUS: http://code.google.com/p/beast-mcmc/wiki/NexusMetacommentFormat

See Also

read.beast.table to extract internal node data from NEXUS file, rbeauti to create XML input for BEAST. HPDbars for plotting highest posterior densities on phylogenies has been moved to package viper.

read.beast.table  Extract node data from BEAST chronogram

Description

This function reads a BEAST chronogram such as produced by TreeAnnotator and extracts time, rate, and support values for internal and external nodes. Nodes in the resulting data frame are ordered exactly like in the NEXUS file.

Usage

read.beast.table(file, digits = 2)
sister

Identification of Sister Nodes and Clades

Description

For any given internal node in a phylogeny, this function returns the sister clade.

Usage

sister(phy, node, type = "terminal", label = FALSE)

Arguments

phy An object of class phylo.
node A vector of mode "numeric" or "character" giving the number(s) or name(s) of the tiplabel(s); these must be monophyletic.
type A character string, may be "terminal", "internal", "daughter", "all", or any unambiguous abbreviation of these; "daughter" will return the MRCA of the sister clade of "node".
label Logical, determining if tip number or tip labels will be returned.

Value

A vector of mode "numeric" or "character", containing either tip numbers or labels, respectively.

See Also

descendants, noi.
Examples

# A phylogeny of bark beetles ...
data(ips.tree)
tcol <- rep("black", Ntip(ips.tree))
tcol[ips.tree$tip.label %in% c("Ips_typographus", "Ips_nitidus")]<- "blue"
tcol[ips.tree$tip.label %in% c("Ips_duplicatus")]<- "red"
plot(ips.tree, no.margin = TRUE, tip.color = tcol)
# What is the sister species of Ips typographus?
sister(ips.tree, "Ips_typographus", label = TRUE)
# Return the MRCA of the sister clade of Ips duplicatus
x <- sister(ips.tree, "Ips_duplicatus", "daughter")
nodelabels(node = x, pch = 21, bg = "red")

splitIntoClades

Find Monophyletic Subsets in Species Lists

Description

Takes a phylogeny and a subset of its tip labels and splits the list of tip labels into monophyletic groups (clades).

Usage

splitIntoClades(phy, tips)

Arguments

phy An object of class phylo.
tips A vector of mode "character" containing any subset of the tip labels in phy.

Value

A list.

terminalSisters

Find Pairs of Sister Species

Description

Finds pairs of sister species in a phylogenetic tree.

Usage

terminalSisters(phy, labels = TRUE)
tipHeights

Arguments

phy
An object of class phylo.

labels
Logical, indicating whether to return tip labels or tip numbers.

Value

A list of which each element contains the tip labels of a sister species pair.

Examples

set.seed(1234)
tr <- rtree(12)
plot(tr)
terminalSisters(tr)

tipHeights(phy)

Description

For each tip (leave, terminal node) in the phylogenetic tree the edge lengths (branch lengths) from root to tip, be it units of time or divergence, is summed up.

Usage

tipHeights(phy)

Arguments

phy
an object of class phylo.

Value

a numeric vector with distances from root to tip for each tip in the phylogenetic tree.

Author(s)

Christoph Heibl

See Also

branching.times
traitRate

Trait-Dependent Shifts in Molecular Rate

Description
Detection of trait-dependent shifts in the rate of molecular evolution with traitRate (Mayrose & Otto, 2011).

Usage
traitRate(phy, seq, x, mainType = "Optimize_Model", n, charModelParam1 = 0.5, charModelParam2 = 1, gammaParam = 0.5, seqModelParam1 = 2, exec = "/Applications/traitRate-1.1/programs/traitRate")

Arguments
phy    a ultrametric phylogenetic tree of class phylo.
seq    a multiple sequence alignment of class DNAbin.
x      data frame containing a binary character in the first column.
mainType    character string giving the type of analysis; two choices are possible: "Optimize_Model" will produce MLE of parameters and "runTraitBootstrap" will perform a parametric bootstrap analysis.
n    numeric, the number of bootstrap replicates. Will be ignored if mainType = "Optimize_Model".
charModelParam1    numeric, giving an initial value for the rate of transitions of character state 0 to 1.
charModelParam2    numeric, giving an initial value for the rate of transitions of character state 1 to 0.
gammaParam    numeric, giving an initial value for the alpha parameter of the model of sequence evolution.
seqModelParam1    numeric, giving an initial value for the kappa parameter of the model of sequence evolution.
exec    character string giving the path to the program directory.

Value
Currently none, but look for the output files in the 'RESULTS' subdirectory in the current working directory.

Note
This function is under development!
trimEnds

Author(s)
Christoph Heibl

References

See Also
- `read.tree` for reading phylogenetic trees, `read.fas` for reading multiple sequence alignments in FASTA format.

trimEnds

**Trim Alignment Ends**

Description
Trims both ends of a DNA sequence alignment to the first and last alignment positions that contain a minimum number of IUPAC base characters ("a", "c", "g", "t", "r", "y", "s", "w", "k", "m", "b", "d", "h", "v"). In addition, all gap characters ("-")) beyond the first and last base characters of each sequence are replaced by the character "n".

Usage
trimEnds(x, min.n.seq = 4)

Arguments
- **x**
  An object of class DNAbin.
- **min.n.seq**
  A numeric giving the required minimum number of sequences having an non-ambiguous base character (a, c, g, t) in the first and last position of the alignment; defaults to 4, which is the minimum number of sequences needed to produce a non-trivial unrooted topology. Can also be given as a fraction.

Value
An object of class DNAbin.

See Also
- `deleteEmptyCells`, `deleteGaps`
Examples

```r
# simple example alignment:
x <- structure(list(nb = 5, seq = c("acaaggtaca", "-caaggtac-",
"acaaggtaca", "aca--gtaca", "-ccaggta--"), nam = LETTERS[1:5]),
.Names = c("nb", "seq", "nam"), class = "alignment")
# convert to DNABin:
x <- as.DNAbin(x)
# fill missing nucleotides:
x <- trimEnds(x)
# show results:
as.character(x[2, ])
```

---

**unlistFirstLevel**  
*Unlist To First Level Only*

**Description**

Does the same as `unlist`, but recurses only one level.

**Usage**

```r
unlistFirstLevel(z, use.names = TRUE)
```

**Arguments**

- `z`  
  A list of lists.
- `use.names`  
  Logical, indicating if element names from the element should be preserved.

---

**write.fas**  
*Write DNA Sequences to File*

**Description**

Write DNA sequences and morphological data to FASTA, PHYLIP, or NEXUS formatted files.

**Usage**

```r
write.fas(x, file, block.width = FALSE, 
          truncate = FALSE, append = FALSE)
write.phy(x, file, block.width = FALSE, 
          strict = FALSE)
write.nex(x, file, block.width = 60, 
          taxblock = FALSE)
```
write.fas

Arguments

x  an object of class DNAbin (usually as matrix, but write.fas also accepts lists) or a list of objects of class DNAbin (only write.nex) or a data frame containing standard (morphological, etc.) data (only write.nex).

file  a character string giving the filename; a special case is file = "", which causes the file content to be written on the standard output connection (i.e. the console). If file is left unspecified (default), the file content is returned as a vector of mode "character" and can be used as a building block for more complex data files.

block.width  an integer, giving the number of characters per line.

truncate  truncation of taxon names to the number of characters given as a integer, otherwise (default) taxon names will not be changed.

append  logical, if TRUE the sequences will be appended to file (if it exists).

strict  logical, if TRUE the names of the sequences will be truncated to 10 strings.

taxblock  logical, if TRUE, a tax block will be added to the NEXUS file.

Details

write.nex can handle multiple DNA sequence alignments, which are handed over as a list of objects of class DNAbin. Correct matching of the rows in the alignments is cared for automatically, hence the individual alignments can contain different numbers of samples and samples need not be in the same order.

Value

None, except when called with file left unspecified, which causes the file content to be returned as a vector of mode "character". This is particularly useful for constructing special types of input files, e.g. for MrBayes (mrbayes).

Author(s)

Christoph Heibl

References


See Also

read.fas, read.phy, and read.nex for reading of DNA sequence files.

Examples

data(ips.cox1)
data(ips.28S)

## Examples for FASTA files
```r
## ------------------------
write.fas(ips.cox1[1:5, 1:120], block.width = 60)

## Examples for PHYLIP files
## ------------------------
write.phy(ips.cox1[1:5, 1:20], block.width = 40)

## Examples for NEXUS files
## ------------------------
x <- list(cox1 = ips.cox1[1:5, 1:10],
          rna28S = ips.28S[1:5, 1:30])
write.nex(x, block.width = 20)

# Truncation of taxonnames:
# -------------------------
rownames(ips.cox1)[1] <- "AVeeeeeeyLongName"
write.fas(ips.cox1, truncate = 10)

# If truncation leads to identical taxonnames,
# a warning will be issued:
# -------------------------
rownames(ips.cox1)[1:2] <- "AVeeeeeeyLongName"
write.fas(ips.cox1, truncate = 10)
```
## Index

* **datasets**
  - ips.16S, 16
  - ips.28S, 17
  - ips.cox1, 17
  - ips.tree, 18

* **package**
  - ips-package, 2
  - .Machine, 13

AAbin, 21
ace, 29
aliscore, 3, 3, 8, 15, 20, 35, 41
ape, 3
branching.times, 45
cbind.DNAbin, 39, 41
chronos, 13
code.simple.gaps, 5, 8
collapseUnsupportedEdges, 5
del.gaps, 8
del.miss, 6
deleteEmptyCells, 5, 7, 47
deleteGaps, 5, 7, 8, 47
descendants, 9, 11, 43
DNAbin, 5–8, 10, 14, 16, 17, 21, 22, 34, 36, 38, 39, 46, 49
DNAbin2index, 10, 16
eoi, 10
fixNodes, 12
forceEqualTipHeights, 13

gblocks, 3, 4, 8, 14, 20, 35, 41
index2DNAbin, 10, 16
ips (ips-package), 2
ips-package, 2
ips.16S, 16
ips.28S, 17
ips.cox1, 17
ips.tree, 18
is.ultrametric, 13
ladderize, 3, 12, 13
mafft, 3, 4, 15, 19, 22, 32, 33, 35, 41
mafft.merge, 21
mrbayes, 3, 21, 23, 24, 26, 27, 49
mrbayes.lset, 22, 23, 26, 27
mrbayes.mcmc, 22, 24, 26, 27
mrbayes.prset, 22, 24, 26, 26
mrca, 11
mst, 29
multistate, 28
multistateMCMC, 3
multistateMCMC (multistate), 28
multistateML, 3
multistateML (multistate), 28
neighboringPairs, 29
noi, 9, 43
noi (eoi), 10
ntip, 30
oi (eoi), 10
partitionfinder, 30
pathd8, 31
phylo, 6, 9, 10, 12, 13, 19, 21, 30–33, 36, 43–46
phylo2mafft, 32
phylo2mst, 33
pis, 33
prank, 3, 4, 15, 20, 22, 34, 41
raxml, 3, 22, 35, 38, 39
raxml.partitions, 37, 38
rbeauti, 3, 39, 42, 43
read, 40
read.beast, 3, 13, 40, 41, 43
read.beast.table, 3, 40, 42, 42
read.fas, 3, 20, 35, 47, 49
read.mrbayes (read.beast), 41
read.nex, 3, 49
read.nexus, 13
read.phy, 3, 49
read.starbeast (read.beast), 41
read.tree, 13, 47
root, 12
rotate, 3, 12, 13

set.seed, 36
sister, 9, 43
splitIntoClades, 44

terminalSisters, 44
tipHeights, 13, 45
traitRate, 46
trimEnds, 5, 7, 47

unlistFirstLevel, 48

which, 10
write.fas, 3, 48
write.nex, 3
write.nex (write.fas), 48
write.phy, 3
write.phy (write.fas), 48