Package 'TreeSearch'

March 22, 2022

```
Title Phylogenetic Analysis with Morphological Data
Version 1.1.1
License GPL (>= 3)
Copyright Incorporates C/C++ code from
      Morphy Phylogenetic Library by Martin Brazeau
      <a href="https://github.com/mbrazeau/MorphyLib">https://github.com/mbrazeau/MorphyLib</a> (GPL3)
Description Reconstruct phylogenetic trees from discrete data.
      Inapplicable character states are handled using the algorithm of Brazeau,
      Guillerme and Smith (2019) <doi:10.1093/sysbio/syy083> with the ``Morphy"
      library, under equal or implied step weights.
      Contains a ``shiny" user interface for interactive tree search and exploration
      of results, including character visualization, rogue taxon detection,
      tree space mapping, and cluster consensus trees (Smith 2022a, b)
      <doi:10.1093/sysbio/syab099>, <doi:10.1093/sysbio/syab100>.
      Profile Parsimony (Faith and Trueman, 2001) <doi:10.1080/10635150118627>,
      Successive Approximations (Farris, 1969) <doi:10.2307/2412182>
      and custom optimality criteria are implemented.
URL https://ms609.github.io/TreeSearch/, https://github.com/ms609/TreeSearch/
BugReports https://github.com/ms609/TreeSearch/issues/
Depends R (>= 3.5.0)
Imports ape (>= 5.6),
      cli (>= 3.0),
      cluster,
      fastmatch (>= 1.1.3),
      future.
      promises,
      protoclust,
      Rcpp,
      Rdpack (>= 0.7),
      Rogue (> 2.0.0),
      shiny (>= 1.6.0),
      shinyjs,
      stats,
      TreeDist (>= 2.3.0),
      TreeTools (>= 1.7.0),
Suggests knitr,
      phangorn (>= 2.2.1),
```

R topics documented:

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Description

Generates a starting tree by adding each taxon in turn to the most parsimonious location.

Usage

AdditionTree(dataset, concavity = Inf, constraint, sequence)

Arguments

dataset	A phylogenetic data matrix of phangorn class phyDat, whose names correspond to the labels of any accompanying tree.
concavity	Numeric specifying concavity constant for implied step weighting. The most appropriate value will depend on the dataset, but values around 10–15 often perform well (Goloboff et al. 2018; Smith 2019). The character string "profile" employs an approximation of profile parsimony (Faith and Trueman 2001). Set as Inf for equal step weights, which underperforms step weighting approaches (Goloboff et al. 2008; Goloboff et al. 2018; Goloboff and Arias 2019; Smith 2019).
constraint	An object of class phyDat; returned trees will be perfectly compatible with each character in constraint. See ImposeConstraint() and vignette for further examples.
sequence	Character or numeric vector listing sequence in which to add taxa. Randomized if not provided.

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Value

AdditionTree() returns a tree of class phylo, rooted on sequence[1].

Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

See Also

```
Impose a constraint: TreeTools::ImposeConstraint()
Neighbour-joining trees: TreeTools::NJTree(); TreeTools::ConstrainedNJ()
Other tree generation functions: RandomMorphyTree()
```

Examples

```
data('Lobo', package = 'TreeTools')
AdditionTree(Lobo.phy, concavity = 10)
```

AllSPR

All SPR trees

Description

All SPR trees

Usage

```
AllSPR(parent, child, nEdge, notDuplicateRoot, edgeToBreak)
```

Arguments

parent Integer vector corresponding to the first column of the edge matrix of a tree of

class phylo, i.e. tree\$edge[,1].

child Integer vector corresponding to the second column of the edge matrix of a tree

of class phylo, i.e. tree\$edge[,2].

nEdge integer specifying the number of edges of a tree of class phylo, i.e. dim(tree\$edge)[1]

notDuplicateRoot

edgeToBreak

logical vector of length nEdge, specifying for each edge whether it is the second edge leading to the root (in which case its breaking will be equivalent to breaking the other root edge... except insofar as it moves the position of the root.)

the other root edge... except hisoral as it moves the position of the root.)

(optional) integer specifying the index of an edge to bisect/prune, generated randomly if not specified. Alternatively, set to -1 to return a complete list of all

trees one step from the input tree.

Value

AllSPR() returns a list of edge matrices for all trees one SPR rearrangement from the starting tree

Author(s)

Martin R. Smith

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Carter1

Number of trees with m steps

Description

Calculate the number of trees in which Fitch parsimony will reconstruct m steps, where a leaves are labelled with one state, and b leaves are labelled with a second state.

Usage

```
Carter1(m, a, b)
Log2Carter1(m, a, b)
LogCarter1(m, a, b)
```

Arguments

m Number of steps.

a, b Number of leaves labelled 0 and 1.

Details

Implementation of theorem 1 from Carter et al. (1990)

Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

References

Carter M, Hendy M, Penny D, Székely LA, Wormald NC (1990). "On the distribution of lengths of evolutionary trees." *SIAM Journal on Discrete Mathematics*, **3**(1), 38–47.

See also:

Steel MA (1993). "Distributions on bicoloured binary trees arising from the principle of parsimony." *Discrete Applied Mathematics*, **41**(3), 245–261.

```
\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.1016/0166-218X(90)90058-K")}
```

```
,https://doi.org/10.1016/0166-218X(90)90058-K.
```

Steel M, Charleston M (1995). "Five surprising properties of parsimoniously colored trees." *Bulletin of Mathematical Biology*, **57**(2), 367–375.

```
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1016/0092-8240(94)00051-D")}
```

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```
, https://doi.org/10.1016/0092-8240(94)00051-D.

(Steel M, Goldstein L, Waterman MS (1996). "A central limit theorem for the parsimony length of trees." Advances in Applied Probability, 28(4), 1051-1071.

\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.2307/1428164")}
.)
```

See Also

Other profile parsimony functions: PrepareDataProfile(), StepInformation(), WithOneExtraStep(), profiles

Examples

```
# The character `0 0 0 1 1 1`
Carter1(1, 3, 3) # Exactly one step
Carter1(2, 3, 3) # Two steps (one extra step)

# Number of trees that the character can map onto with exactly _m_ steps
# if non-parsimonious reconstructions are permitted:
cumsum(sapply(1:3, Carter1, 3, 3))
# Three steps allow the character to map onto any of the 105 six-leaf trees.
```

CharacterLength

Character length

Description

Homoplasy length of each character in a dataset on a specified tree.

Usage

```
CharacterLength(tree, dataset, compress = FALSE)
FitchSteps(tree, dataset)
FastCharacterLength(tree, dataset)
```

Arguments

tree A tree of class phylo.

dataset A phylogenetic data matrix of phangorn class phyDat, whose names corre-

spond to the labels of any accompanying tree.

compress Logical specifying whether to retain the compression of a phyDat object or to

return a vector specifying to each individual character, decompressed using the

dataset's index attribute.

ClusterStrings 7

Value

CharacterLength() returns a vector listing the contribution of each character to tree score, according to the algorithm of Brazeau et al. (2019).

Functions

• FastCharacterLength: Do not perform checks. Use with care: may cause erroneous results or software crash if variables are in the incorrect format.

Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

References

Brazeau MD, Guillerme T, Smith MR (2019). "An algorithm for morphological phylogenetic analysis with inapplicable data." *Systematic Biology*, **68**(4), 619–631.

```
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1093/sysbio/syy083")}
```

.

See Also

```
Other tree scoring: IWScore(), MinimumLength(), MorphyTreeLength()
```

Examples

```
data('inapplicable.datasets')
dataset <- inapplicable.phyData[[12]]
tree <- TreeTools::NJTree(dataset)
CharacterLength(tree, dataset)
CharacterLength(tree, dataset, compress = TRUE)</pre>
```

ClusterStrings

Cluster similar strings

Description

Calculate string similarity using the Levenshtein distance and return clusters of similar strings.

Usage

```
ClusterStrings(x, maxCluster = 12)
```

Arguments

Character vector.

maxCluster Integer specifying maximum number of clusters to consider.

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Value

NameClusters() returns an integer assigning each element of x to a cluster, with an attribute med specifying the median string in each cluster, and silhouette reporting the silhouette coefficient of the optimal clustering. Coefficients < 0.5 indicate weak structure, and no clusters are returned. If the number of unique elements of x is less than maxCluster, all occurrences of each entry are assigned to an individual cluster.

Author(s)

```
Martin R. Smith (martin.smith@durham.ac.uk)
```

Examples

ConcordantInformation Evaluate the concordance of information between a tree and a dataset

Description

Details the amount of information in a phylogenetic dataset that is consistent with a specified phylogenetic tree, and the signal:noise ratio of the character matrix implied if the tree is true.

Usage

```
ConcordantInformation(tree, dataset)
Evaluate(tree, dataset)
ConcordantInfo(tree, dataset)
```

Arguments

tree A tree of class phylo.

dataset A phylogenetic data matrix of phangorn class phyDat, whose names corre-

spond to the labels of any accompanying tree.

Details

Presently restricted to datasets whose characters contain a maximum of two parsimony-informative states.

Value

ConcordantInformation() returns a named vector with elements:

- informationContent: cladistic information content of dataset
- signal, noise: amount of cladistic information that represents phylogenetic signal and noise, according to tree

- signalToNoise: the implied signal:noise ratio of dataset
- treeInformation: the cladistic information content of a bifurcating tree on dataset; this is the minimum amount of information necessary to resolve a bifurcating tree, assuming no duplicate information or noise
- matrixToTree: the ratio of the cladistic information content of the matrix to the cladistic information content of the tree, a measure of the redundancy of the matrix
- ignored: information content of characters whose signal and noise could not be calculated (too many states) and so are not included in the totals above.

Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

Examples

```
data(congreveLamsdellMatrices)
myMatrix <- congreveLamsdellMatrices[[10]]
ConcordantInformation(TreeTools::NJTree(myMatrix), myMatrix)</pre>
```

congreveLamsdellMatrices

100 simulated data matrices

Description

Contains the 100 simulated matrices generated by (Congreve and Lamsdell 2016) using a heterogeneous Markov-k model, generated from the referenceTree topology, with all branches sharing an equal length.

Usage

congreveLamsdellMatrices

Format

A list with 100 entries, each comprising a phyDat object of 55 characters for 22 taxa

Source

```
doi: 10.5061/dryad.7dq0j
```

References

Congreve CR, Lamsdell JC (2016). "Implied weighting and its utility in palaeontological datasets: a study using modelled phylogenetic matrices." *Palaeontology*, **59**(3), 447–465.

```
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1111/pala.12236")}
```

.

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Examples

```
data('referenceTree')
data('congreveLamsdellMatrices')
TreeLength(referenceTree, congreveLamsdellMatrices[[17]], 'profile')
```

Consistency

Consistency / retention 'indices'

Description

Consistency() calculates the so-called consistency and retention 'indices' for each character in a dataset, given a bifurcating tree. Although there is not a straightforward interpretation of these indices, they are sometimes taken as an indicator of the fit of a character to a tree. Values correlate with the number of species sampled and the distribution of taxa between character states, so are not strictly comparable between characters in which these factors differ.

Usage

```
Consistency(dataset, tree, compress = FALSE)
```

Arguments

dataset A phylogenetic data matrix of **phangorn** class phyDat, whose names corre-

spond to the labels of any accompanying tree.

tree A tree of class phylo.

compress Logical specifying whether to retain the compression of a phyDat object or to

return a vector specifying to each individual character, decompressed using the

dataset's index attribute.

Details

#TODO: Retention index not yet implemented.

Value

Consistency() returns a named vector specifying the consistency index (ci), retention index (ri), and rescaled consistency index (rc).

Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

Examples

```
data(inapplicable.datasets)
dataset <- inapplicable.phyData[[4]]
Consistency(dataset, TreeTools::NJTree(dataset))</pre>
```

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cSPR

cSPR() expects a tree rooted on a single tip.

Description

cSPR() expects a tree rooted on a single tip.

Usage

```
cSPR(tree, whichMove = NULL)
```

Arguments

tree A tree of class phylo.

whichMove Integer specifying which SPR move index to perform.

Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

Examples

```
tree <- TreeTools::BalancedTree(8)

# Tree must be rooted on leaf
tree <- TreeTools::RootTree(tree, 1)

# Random rearrangement
cSPR(tree)

# Specific rearrangement
cSPR(tree, 9)</pre>
```

GapHandler

Read how a Morphy Object handles the inapplicable token

Description

Gaps represented by the inapplicable token can be treated as 'missing data', i.e. as equivalent to the ambiguous token ?; as an extra state, equivalent to other states such as \emptyset or 1; or as 'inapplicable data' using the algorithm of Brazeau, Guillerme and Smith (2019).

Usage

```
GapHandler(morphyObj)
```

Arguments

morphy0bj Object of class morphy, perhaps created with PhyDat2Morphy().

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Value

GapHandler() returns a character string stating how gaps are handled by morphy0bj.

Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

See Also

```
Other Morphy API functions: MorphyErrorCheck(), MorphyWeights(), PhyDat2Morphy(), SingleCharMorphy(), UnloadMorphy(), is.morphyPtr(), mpl_apply_tipdata(), mpl_attach_rawdata(), mpl_attach_symbols(), mpl_delete_Morphy(), mpl_delete_rawdata(), mpl_first_down_recon(), mpl_first_up_recon(), mpl_get_charac_weight(), mpl_get_gaphandl(), mpl_get_num_charac(), mpl_get_num_internal_nodes(), mpl_get_numtaxa(), mpl_get_symbols(), mpl_init_Morphy(), mpl_new_Morphy(), mpl_second_down_recon(), mpl_second_up_recon(), mpl_set_charac_weight(), mpl_set_num_internal_nodes(), mpl_set_parsim_t(), mpl_translate_error(), mpl_update_lower_root(), mpl_update_tip(), summary.morphyPtr()
```

Examples

```
morphyObj <- SingleCharMorphy('-0-0', 'Extra')
GapHandler(morphyObj)
morphyObj <- UnloadMorphy(morphyObj)</pre>
```

inapplicable.datasets Thirty datasets with inapplicable data

Description

These are the datasets used to evaluate the behaviour of the inapplicable algorithm in Brazeau et al. (2019). The name of each item corresponds to the datasets listed below. Datasets are sorted into two subsets, each sorted alphabetically; the first subset comprise simpler datasets with faster processing times. inapplicable.datasets provide the data in the matrix format generated by read.nexus.data(); inapplicable.phyData are in phyDat format. inapplicable.trees lists for each dataset a sample of up to 50 trees obtained by tree search under each inapplicable treatment, named accordingly. inapplicable.citations is a named character vector specifying the source of each dataset.

Usage

```
inapplicable.datasets
inapplicable.phyData
inapplicable.trees
inapplicable.citations
```

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Format

An object of class list of length 30.

An object of class list of length 30.

An object of class list of length 31.

An object of class character of length 30.

Source

Subset one (faster processing):

- **Agnarsson2004** AGNARSSON, I. 2004. Morphological phylogeny of cobweb spiders and their relatives (Araneae, Araneoidea, Theridiidae). Zoological Journal of the Linnean Society, 141, 447–626.
- Capa2011 CAPA, M., HUTCHINGS, P., AGUADO, M. T. and BOTT, N. J. 2011. Phylogeny of Sabellidae (Annelida) and relationships with other taxa inferred from morphology and multiple genes. Cladistics, 27, 449–469.
- **DeAssis2011** DE ASSIS, J. E. and CHRISTOFFERSEN, M. L. 2011. Phylogenetic relationships within Maldanidae (Capitellida, Annelida), based on morphological characters. Systematics and Biodiversity, 9, 233–245.
- **OLeary1999** O'LEARY, M. A. and GEISLER, J. H. 1999. The position of Cetacea within Mammalia: phylogenetic analysis of morphological data from extinct and extant taxa. Systematic Biology, 48, 455–490.
- **Rousset2004** ROUSSET, V., ROUSE, G. W., SIDDALL, M. E., TILLIER, A. and PLEIJEL, F. 2004. The phylogenetic position of Siboglinidae (Annelida) inferred from 18S rRNA, 28S rRNA and morphological data. Cladistics, 20, 518–533.
- **Sano2011** SANO, M. and AKIMOTO, S.-I. 2011. Morphological phylogeny of gall-forming aphids of the tribe Eriosomatini (Aphididae: Eriosomatinae). Systematic Entomology, 36, 607–627.
- **Sansom2010** SANSOM, R. S., FREEDMAN, K., GABBOTT, S. E., ALDRIDGE, R. J. and PURNELL, M. A. 2010. Taphonomy and affinity of an enigmatic Silurian vertebrate, *Jamoytius kerwoodi* White. Palaeontology, 53, 1393–1409.
- **Schulze2007** SCHULZE, A., CUTLER, E. B. and GIRIBET, G. 2007. Phylogeny of sipunculan worms: A combined analysis of four gene regions and morphology. Molecular Phylogenetics and Evolution, 42, 171–92.
- **Shultz2007** SHULTZ, J. W. 2007. A phylogenetic analysis of the arachnid orders based on morphological characters. Zoological Journal of the Linnean Society, 150, 221–265.
- **Wetterer2000** WETTERER, A. L., ROCKKMAN, M. V. and SIMMONS, N. B. 2000. Phylogeny of phyllostomid bats (Mammalia: Chiroptera): data from diverse morphological systems, sex chromosomes, and restriction sites. Bulletin of the American Museum of Natural History, 248, 1–200.
- Wills2012 WILLS, M. A., GERBER, S., RUTA, M. and HUGHES, M. 2012. The disparity of priapulid, archaeopriapulid and palaeoscolecid worms in the light of new data. Journal of Evolutionary Biology, 25, 2056–2076.

Subset two (longer processing times):

Aguado2009 AGUADO, M. T. and SAN MARTIN, G. 2009. Phylogeny of Syllidae (Polychaeta) based on morphological data. Zoologica Scripta, 38, 379–402.

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Aria2015 ARIA, C., CARON, J. B. and GAINES, R. 2015. A large new leanchoiliid from the Burgess Shale and the influence of inapplicable states on stem arthropod phylogeny. Palaeontology, 58, 629–660.

- **Asher2005** ASHER, R. J. and HOFREITER, M. 2006. Tenrec phylogeny and the noninvasive extraction of nuclear DNA. Systematic biology, 55, 181–94.
- **Baker2009** BAKER, W. J., SAVOLAINEN, V., ASMUSSEN-LANGE, C. B., CHASE, M. W., DRANSFIELD, J., FOREST, F., HARLEY, M. M., UHL, N. W. and WILKINSON, M. 2009. Complete generic-level phylogenetic analyses of palms (Arecaceae) with comparisons of supertree and supermatrix approaches. Systematic Biology, 58, 240–256.
- **Bouchenak2010** BOUCHENAK-KHELLADI, Y., VERBOOM, G. A., SAVOLAINEN, V. and HODKINSON, T. R. 2010. Biogeography of the grasses (Poaceae): a phylogenetic approach to reveal evolutionary history in geographical space and geological time. Botanical Journal of the Linnean Society, 162, 543–557.
- **Conrad2008** CONRAD, J. L. 2008. Phylogeny And Systematics Of Squamata (Reptilia) Based On Morphology. Bulletin of the American Museum of Natural History, 310, 1–182.
- **Dikow2009** DIKOW, T. 2009. A phylogenetic hypothesis for Asilidae based on a total evidence analysis of morphological and DNA sequence data (Insecta: Diptera: Brachycera: Asiloidea). Organisms Diversity and Evolution, 9, 165–188.
- **Eklund2004** EKLUND, H., DOYLE, J. A. and HERENDEEN, P. S. 2004. Morphological phylogenetic analysis of living and fossil Chloranthaceae. International Journal of Plant Sciences, 165, 107–151.
- **Geisler 2001** GEISLER, J. H. 2001. New morphological evidence for the phylogeny of Artiodactyla, Cetacea, and Mesonychidae. American Museum Novitates, 3344, 53.
- **Giles2015** GILES, S., FRIEDMAN, M. and BRAZEAU, M. D. 2015. Osteichthyan-like cranial conditions in an Early Devonian stem gnathostome. Nature, 520, 82–85.
- **Griswold1999** GRISWOLD, C. E., CODDINGTON, J. A., PLATNICK, N. I. and FORSTER, R. R. 1999. Towards a phylogeny of entelegyne spiders (Araneae, Araneomorphae, Entelegynae). Journal of Arachnology, 27, 53–63.
- **Liljeblad2008** LILJEBLAD, J., RONQUIST, F., NIEVES-ALDREY, J. L., FONTAL-CAZALLA, F., ROS-FARRE, P., GAITROS, D. and PUJADE-VILLAR, J. 2008. A fully web-illustrated morphological phylogenetic study of relationships among oak gall wasps and their closest relatives (Hymenoptera: Cynipidae).
- **Loconte1991** LOCONTE, H. and STEVENSON, D. W. 1991. Cladistics of the Magnoliidae. Cladistics, 7, 267–296.
- **Longrich2010** LONGRICH, N. R., SANKEY, J. and TANKE, D. 2010. *Texacephale langstoni*, a new genus of pachycephalosaurid (Dinosauria: Ornithischia) from the upper Campanian Aguja Formation, southern Texas, USA. Cretaceous Research, 31, 274–284.
- **OMeara2014** O'MEARA, R. N. and THOMPSON, R. S. 2014. Were There Miocene Meridiolestidans? Assessing the phylogenetic placement of *Necrolestes patagonensis* and the presence of a 40 million year Meridiolestidan ghost lineage. Journal of Mammalian Evolution, 21, 271–284.
- **Rougier2012** ROUGIER, G. W., WIBLE, J. R., BECK, R. M. D. and APESTEGUIA, S. 2012. The Miocene mammal *Necrolestes* demonstrates the survival of a Mesozoic nontherian lineage into the late Cenozoic of South America. Proceedings of the National Academy of Sciences, 109, 20053–8.
- Sharkey2011 SHARKEY, M. J., CARPENTER, J. M., VILHELMSEN, L., HERATY, J., LILJEBLAD, J., DOWLING, A. P. G., SCHULMEISTER, S., MURRAY, D., DEANS, A. R., RONQUIST, F., KROGMANN, L. and WHEELER, W. C. 2012. Phylogenetic relationships among superfamilies of Hymenoptera. Cladistics, 28, 80–112.

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Sundue2010 SUNDUE, M. A., ISLAM, M. B. and RANKER, T. A. 2010. Systematics of Grammitid Ferns (Polypodiaceae): Using Morphology and Plastid Sequence Data to Resolve the Circumscriptions of Melpomene and the Polyphyletic Genera *Lellingeria* and *Terpsichore*. Systematic Botany, 35, 701–715.

- Vinther2008 VINTHER, J., VAN ROY, P. and BRIGGS, D. E. G. 2008. Machaeridians are Palaeozoic armoured annelids. Nature, 451, 185–188.
- **Wilson2003** WILSON, G. D. F. and EDGECOMBE, G. D. 2003. The Triassic isopod *Protam-phisopus wianamattensis* (Chilton) and comparison by extant taxa (Crustacea, Phreatoicidea). Journal of Paleontology, 77, 454–470.
- **Wortley2006** WORTLEY, A. H. and SCOTLAND, R. W. 2006. The effect of combining molecular and morphological data in published phylogenetic analyses. Systematic Biology, 55, 677–685.
- **Zanol2014** ZANOL, J., HALANYCH, K. M. and FAUCHALD, K. 2014. Reconciling taxonomy and phylogeny in the bristleworm family Eunicidae (Polychaete, Annelida). Zoologica Scripta, 43, 79–100.
- **Zhu2013** ZHU, M., YU, X., AHLBERG, P. E., CHOO, B., LU, J., QIAO, T., QU, Q., ZHAO, W., JIA, L., BLOM, H. and ZHU, Y. 2013. A Silurian placoderm with osteichthyan-like marginal jaw bones. Nature, 502, 188–193.

References

Brazeau MD, Guillerme T, Smith MR (2019). "An algorithm for morphological phylogenetic analysis with inapplicable data." *Systematic Biology*, **68**(4), 619–631.

```
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1093/sysbio/syy083")}
```

Examples

```
data('inapplicable.datasets', package = 'TreeSearch')
names(inapplicable.datasets)
```

is.morphyPtr

Is an object a valid Morphy object?

Description

Is an object a valid Morphy object?

Usage

```
is.morphyPtr(morphyObj)
```

Arguments

morphy0bj Object of class morphy, perhaps created with PhyDat2Morphy().

Value

is.morphyPtr() returns TRUE if morphyObj is a valid morphy pointer, FALSE otherwise.

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Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

See Also

```
Other Morphy API functions: GapHandler(), MorphyErrorCheck(), MorphyWeights(), PhyDat2Morphy(), SingleCharMorphy(), UnloadMorphy(), mpl_apply_tipdata(), mpl_attach_rawdata(), mpl_attach_symbols(), mpl_delete_Morphy(), mpl_delete_rawdata(), mpl_first_down_recon(), mpl_first_up_recon(), mpl_get_charac_weight(), mpl_get_gaphandl(), mpl_get_num_charac(), mpl_get_num_internal_nodes(), mpl_get_numtaxa(), mpl_get_symbols(), mpl_init_Morphy(), mpl_new_Morphy(), mpl_second_down_recon(), mpl_second_up_recon(), mpl_set_charac_weight(), mpl_set_num_internal_nodes(), mpl_set_parsim_t(), mpl_translate_error(), mpl_update_lower_root(), mpl_update_tip(), summary.morphyPtr()
```

IWScore

Calculate the parsimony score of a tree given a dataset

Description

TreeLength() uses the Morphy library (Brazeau et al. 2017) to calculate a parsimony score for a tree, handling inapplicable data according to the algorithm of Brazeau et al. (2019). Trees may be scored using equal weights, implied weights (Goloboff 1993), or profile parsimony (Faith and Trueman 2001).

Usage

```
IWScore(tree, dataset, concavity = 10L, ...)
TreeLength(tree, dataset, concavity = Inf)
## S3 method for class 'phylo'
TreeLength(tree, dataset, concavity = Inf)
## S3 method for class 'numeric'
TreeLength(tree, dataset, concavity = Inf)
## S3 method for class 'list'
TreeLength(tree, dataset, concavity = Inf)
## S3 method for class 'multiPhylo'
TreeLength(tree, dataset, concavity = Inf)
Fitch(tree, dataset)
```

Arguments

tree A tree of class phylo, a list thereof (optionally of class multiPhylo), or an

integer – in which case tree random trees will be uniformly sampled.

dataset A phylogenetic data matrix of **phangorn** class phyDat, whose names corre-

spond to the labels of any accompanying tree.

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concavity

Determines the degree to which extra steps beyond the first are penalized. Specify a numeric value to use implied weighting (Goloboff 1993); concavity specifies k in k / e + k. A value of 10 is recommended; TNT sets a default of 3, but this is too low in some circumstances (Goloboff et al. 2018; Smith 2019). Better still explore the sensitivity of results under a range of concavity values, e.g. $k = 2 ^ (1:7)$. Specify Inf to weight each additional step equally. Specify 'profile' to employ profile parsimony (Faith and Trueman 2001).

unused; allows additional parameters specified within ... to be received by the function without throwing an error.

Value

TreeLength() returns a numeric vector containing the score for each tree in tree.

Author(s)

Martin R. Smith (using Morphy C library, by Martin Brazeau)

References

Brazeau MD, Guillerme T, Smith MR (2019). "An algorithm for morphological phylogenetic analysis with inapplicable data." *Systematic Biology*, **68**(4), 619–631.

.

Brazeau MD, Smith MR, Guillerme T (2017). "MorphyLib: a library for phylogenetic analysis of categorical trait data with inapplicability."

```
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.5281/zenodo.815372")}
```

.

Faith DP, Trueman JWH (2001). "Towards an inclusive philosophy for phylogenetic inference." *Systematic Biology*, **50**(3), 331–350.

```
\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.1080/10635150118627")}
```

•

Goloboff PA (1993). "Estimating character weights during tree search." Cladistics, 9(1), 83–91.

```
\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.1111/j.1096-0031.1993.tb00209.x")}
```

.

Goloboff PA, Torres A, Arias JS (2018). "Weighted parsimony outperforms other methods of phylogenetic inference under models appropriate for morphology." *Cladistics*, **34**(4), 407–437.

```
\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.1111/cla.12205")}
```

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.

Smith MR (2019). "Bayesian and parsimony approaches reconstruct informative trees from simulated morphological datasets." *Biology Letters*, **15**(2), 20180632.

```
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1098/rsbl.2018.0632")}
```

See Also

- Conduct tree search using MaximizeParsimony() (command line), EasyTrees() (graphical user interface), or TreeSearch() (custom optimality criteria).
- See score for each character: CharacterLength().

Other tree scoring: CharacterLength(), MinimumLength(), MorphyTreeLength()

Examples

```
data("inapplicable.datasets")
tree <- TreeTools::BalancedTree(inapplicable.phyData[[1]])
TreeLength(tree, inapplicable.phyData[[1]])
TreeLength(tree, inapplicable.phyData[[1]], concavity = 10)
TreeLength(tree, inapplicable.phyData[[1]], concavity = "profile")
TreeLength(5, inapplicable.phyData[[1]])</pre>
```

Jackknife

Jackknife resampling

Description

Resample trees using Jackknife resampling, i.e. removing a subset of characters.

Usage

```
Jackknife(
   tree,
   dataset,
   resampleFreq = 2/3,
   InitializeData = PhyDat2Morphy,
   CleanUpData = UnloadMorphy,
   TreeScorer = MorphyLength,
   EdgeSwapper = TBRSwap,
   jackIter = 5000L,
   searchIter = 4000L,
   searchHits = 42L,
   verbosity = 1L,
   ...
)
```

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Arguments

dataset a dataset in the format required by TreeScorer().	
resampleFreq Double between 0 and 1 stating proportion of characters to resample.	
InitializeData Function that sets up data object to prepare for tree search. The function will be passed the dataset parameter. Its return value will be passed to TreeScorer(and CleanUpData().	
CleanUpData Function to destroy data object on function exit. The function will be passed the value returned by InitializeData().	ıe
TreeScorer function to score a given tree. The function will be passed three parameter corresponding to the parent and child entries of a tree's edge list, and a dataset	
EdgeSwapper a function that rearranges a parent and child vector, and returns a list with modified vectors; for example SPRSwap().	1-
jackIter Integer specifying number of jackknife iterations to conduct.	
searchIter Integer specifying maximum rearrangements to perform on each bootstrap of ratchet iteration. To override this value for a single swapper function, set e. attr(SwapperFunction, 'searchIter') <-99	
searchHits Integer specifying maximum times to hit best score before terminating a tre search within a ratchet iteration. To override this value for a single swapper function, set e.g. attr(SwapperFunction, 'searchHits') <-99	
verbosity Numeric specifying level of detail to display in console: larger numbers provide more verbose feedback to the user.	le
further arguments to pass to TreeScorer(), e.g. dataset = .	

Details

The function assumes that InitializeData() will return a morphy object; if this doesn't hold for you, post a GitHub issue or e-mail the maintainer.

Value

Jackknife() returns a list of trees recovered after jackknife iterations.

Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

See Also

• JackLabels(): Label nodes of a tree with jackknife supports.

 $Other\ split\ support\ functions:\ JackLabels(), Maximize Parsimony(),\ Site Concordance$

Other custom search functions: EdgeListSearch(), MorphyBootstrap(), SuccessiveApproximations()

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JackLabels 5 4 1

Label nodes with jackknife support values

Description

Label nodes with jackknife support values

Usage

```
JackLabels(
   tree,
   jackTrees,
   plot = TRUE,
   add = FALSE,
   adj = 0,
   col = NULL,
   frame = "none",
   pos = 2L,
   ...
)
```

Arguments

```
tree A tree of class phylo.

jackTrees A list or multiPhylo object containing trees generated by Jackknife().

plot Logical specifying whether to plot results; if FALSE, returns blank labels for nodes near the root that do not correspond to a unique split.

add Logical specifying whether to add the labels to an existing plot.

adj, col, frame, pos, ...

Parameters to pass to nodelabels().
```

Value

A named vector specifying the proportion of jackknife trees consistent with each node in tree, as plotted. If plot = FALSE, blank entries are included corresponding to nodes that do not require labelling; the return value is in the value required by phylo\$node.label.

Author(s)

```
Martin R. Smith (martin.smith@durham.ac.uk)
```

See Also

```
Jackknife(): Generate trees by jackknife resampling
Other split support functions: Jackknife(), MaximizeParsimony(), SiteConcordance
```

Examples

```
library('TreeTools', quietly = TRUE) # for as.phylo

# jackTrees will usually be generated with Jackknife(), but for simplicity:
jackTrees <- as.phylo(1:100, 8)

tree <- as.phylo(0, 8)
JackLabels(tree, jackTrees)

tree$node.label <- JackLabels(tree, jackTrees, plot = FALSE)</pre>
```

MaximizeParsimony

Find most parsimonious trees

Description

Search for most parsimonious trees using the parsimony ratchet and TBR rearrangements, treating inapplicable data as such using the algorithm of Brazeau et al. (2019).

Tree search will be conducted from a specified or automatically-generated starting tree in order to find a tree with an optimal parsimony score, under implied or equal weights, treating inapplicable characters as such in order to avoid the artefacts of the standard Fitch algorithm (see Maddison 1993; Brazeau et al. 2019). Tree length is calculated using the MorphyLib C library (Brazeau et al. 2017).

Usage

```
MaximizeParsimony(
  dataset,
  tree,
  ratchIter = 6L,
  tbrIter = 2L,
  startIter = 2L,
  finalIter = 1L,
  maxHits = NTip(dataset) * 1.8,
  maxTime = 60,
  quickHits = 1/3,
  concavity = Inf,
  tolerance = sqrt(.Machine$double.eps),
  constraint,
  verbosity = 3L
)
Resample(
  dataset,
  tree,
  method = "jack",
  proportion = 2/3,
  ratchIter = 1L,
  tbrIter = 8L,
  finalIter = 3L,
  maxHits = 12L,
```

```
concavity = Inf,
tolerance = sqrt(.Machine$double.eps),
constraint,
verbosity = 2L,
...
)

EasyTrees()

EasyTreesy()
```

Arguments

dataset A phylogenetic data matrix of phangorn class phyDat, whose names corre-

spond to the labels of any accompanying tree.

tree (optional) A bifurcating tree of class phylo, containing only the tips listed in

dataset, from which the search should begin. If unspecified, an addition tree will be generated from dataset, respecting any supplied constraint. Edge $\frac{1}{2}$

lengths are not supported and will be deleted.

ratchIter Numeric specifying number of iterations of the parsimony ratchet (Nixon 1999)

to conduct.

tbrIter Numeric specifying the maximum number of TBR break points to evaluate be-

fore concluding each search. The counter is reset to zero each time tree score improves. The counter is reset to zero each time tree score improves. One 'iteration' comprises breaking a single branch and evaluating all possible recon-

nections.

startIter Numeric: an initial round of tree search with startIter × tbrIter TBR break

points is conducted in order to locate a local optimum before beginning ratchet

searches.

finalIter Numeric: a final round of tree search will evaluate finalIter × tbrIter TBR

break points, in order to sample the final optimal neighbourhood more intensely.

maxHits Numeric specifying the maximum times that an optimal parsimony score may

be hit before concluding a ratchet iteration or final search concluded.

maxTime Numeric: after maxTime minutes, stop tree search at the next opportunity.

quickHits Numeric: iterations on subsampled datasets will retain quickHits × maxHits

trees with the best score.

concavity Numeric specifying concavity constant for implied step weighting. The most

appropriate value will depend on the dataset, but values around 10–15 often perform well (Goloboff et al. 2018; Smith 2019). The character string "profile" employs an approximation of profile parsimony (Faith and Trueman 2001). Set as Inf for equal step weights, which underperforms step weighting approaches (Goloboff et al. 2008; Goloboff et al. 2018; Goloboff and Arias 2019; Smith

2019).

tolerance Numeric specifying degree of suboptimality to tolerate before rejecting a tree. The default, sqrt(.Machine\$double.eps), retains trees that may be equally

parsimonious but for rounding errors. Setting to larger values will include trees suboptimal by up to tolerance in search results, which may improve the accu-

racy of the consensus tree (at the expense of resolution) (Smith 2019).

constraint	An object of class phyDat; returned trees will be perfectly compatible with each character in constraint. See ImposeConstraint() and vignette for further examples.
verbosity	Integer specifying level of messaging; higher values give more detailed commentary on search progress. Set to \emptyset to run silently.
method	Unambiguous abbreviation of jackknife or bootstrap specifying how to resample characters. Note that jackknife is considered to give more meaningful results.
proportion	Numeric between 0 and 1 specifying what proportion of characters to retain under jackknife resampling.
	Additional parameters to MaximizeParsimony().

Details

Tree search commences with ratchIter iterations of the parsimony ratchet (Nixon 1999), which bootstraps the input dataset in order to escape local optima. A final round of tree bisection and reconnection (TBR) is conducted to broaden the sampling of trees.

This function can be called using the R command line / terminal, or through the 'shiny' graphical user interface app (type EasyTrees() to launch).

For detailed documentation of the 'TreeSearch' package, including full instructions for loading phylogenetic data into R and initiating and configuring tree search, see the package documentation.

Value

MaximizeParsimony() returns a list of trees with class multiPhylo. This lists all trees found during each search step that are within tolerance of the optimal score, listed in the sequence that they were first visited, and named according to the step in which they were first found; it may contain more than maxHits elements. Note that the default search parameters may need to be increased in order for these trees to be the globally optimal trees; examine the messages printed during tree search to evaluate whether the optimal score has stabilized.

The return value has the attribute firstHit, a named integer vector listing the number of optimal trees visited for the first time in each stage of the tree search. Stages are named:

- seed: starting trees;
- start: Initial TBR search;
- ratchN: Ratchet iteration N;
- final: Final TBR search. The first tree hit for the first time in ratchet iteration three is named ratch3_1.

Resample() returns a multiPhylo object containing a list of trees obtained by tree search using a resampled version of dataset.

Resampling

Note that bootstrap support is a measure of the amount of data supporting a split, rather than the amount of confidence that should be afforded the grouping. "Bootstrap support of 100% is not enough, the tree must also be correct" (Phillips et al. 2004). See discussion in Egan (2006); Wagele et al. (2009); (Simmons and Freudenstein 2011); Kumar et al. (2012).

For a discussion of suitable search parameters in resampling estimates, see Muller (2005). The user should decide whether to start each resampling from the optimal tree (which may be quicker, but

result in overestimated support values as searches get stuck in local optima close to the optimal tree) or a random tree (which may take longer as more rearrangements are necessary to find an optimal tree on each iteration).

For other ways to estimate clade concordance, see SiteConcordance().

Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

References

Brazeau MD, Guillerme T, Smith MR (2019). "An algorithm for morphological phylogenetic analysis with inapplicable data." *Systematic Biology*, **68**(4), 619–631.

```
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1093/sysbio/syy083")}
```

.

Brazeau MD, Smith MR, Guillerme T (2017). "MorphyLib: a library for phylogenetic analysis of categorical trait data with inapplicability."

```
\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.5281/zenodo.815372")}
```

.

Egan MG (2006). "Support versus corroboration." *Journal of Biomedical Informatics*, **39**(1), 72–85.

```
\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.1016/j.jbi.2005.11.007")}
```

•

Faith DP, Trueman JWH (2001). "Towards an inclusive philosophy for phylogenetic inference." *Systematic Biology*, **50**(3), 331–350.

```
\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.1080/10635150118627")}
```

.

Goloboff PA, Arias JS (2019). "Likelihood approximations of implied weights parsimony can be selected over the Mk model by the Akaike information criterion." *Cladistics*, **35**(6), 695–716.

```
\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.1111/cla.12380")}
```

.

Goloboff PA, Carpenter JM, Arias JS, Esquivel DRM (2008). "Weighting against homoplasy improves phylogenetic analysis of morphological data sets." *Cladistics*, **24**(5), 758–773.

```
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1111/j.1096-0031.2008.00209.x")}
```

.

Goloboff PA, Torres A, Arias JS (2018). "Weighted parsimony outperforms other methods of phylogenetic inference under models appropriate for morphology." *Cladistics*, **34**(4), 407–437.

```
\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.1111/cla.12205")}
Kumar S, Filipski AJ, Battistuzzi FU, Kosakovsky Pond SL, Tamura K (2012). "Statistics and
truth in phylogenomics." Molecular Biology and Evolution, 29(2), 457-472.
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1093/molbev/msr202")}
Maddison WP (1993). "Missing data versus missing characters in phylogenetic analysis." Sys-
tematic Biology, 42(4), 576-581.
\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.1093/sysbio/42.4.576")}
Muller KF (2005). "The efficiency of different search strategies in estimating parsimony jackknife,
bootstrap, and Bremer support." BMC Evolutionary Biology, 5(1), 58.
\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.1186/1471-2148-5-58")}
Nixon KC (1999). "The Parsimony Ratchet, a new method for rapid parsimony analysis." Cladis-
tics, 15(4), 407-414. ISSN 0748-3007,
\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.1111/j.1096-0031.1999.tb00277.x")}
Phillips MJ, Delsuc F, Penny D (2004). "Genome-scale phylogeny and the detection of system-
atic biases." Molecular biology and evolution, 21(7), 1455-8.
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1093/molbev/msh137")}
Simmons MP, Freudenstein JV (2011). "Spurious 99% bootstrap and jackknife support for un-
supported clades." Molecular Phylogenetics and Evolution, 61(1), 177–191.
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1016/j.ympev.2011.06.003")}
Smith MR (2019). "Bayesian and parsimony approaches reconstruct informative trees from simu-
lated morphological datasets." Biology Letters, 15(2), 20180632.
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1098/rsbl.2018.0632")}
Wagele JW, Letsch H, Klussmann-Kolb A, Mayer C, Misof B, Wagele H (2009). "Phylogenetic
```

Wagele JW, Letsch H, Klussmann-Kolb A, Mayer C, Misof B, Wagele H (2009). "Phylogenetic support values are not necessarily informative: the case of the Serialia hypothesis (a mollusk phylogeny)." *Frontiers in Zoology*, **6**(1), 12–29.

```
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1186/1742-9994-6-12")}
```

See Also

```
Tree search via graphical user interface: EasyTrees()
Other split support functions: JackLabels(), Jackknife(), SiteConcordance
```

Examples

```
## Only run examples in interactive R sessions
if (interactive()) {
  # launch 'shiny' point-and-click interface
  EasyTrees()
  # Here too, use the "continue search" function to ensure that tree score
  # has stabilized and a global optimum has been found
# Load data for analysis in R
library('TreeTools')
data('congreveLamsdellMatrices', package = 'TreeSearch')
dataset <- congreveLamsdellMatrices[[42]]</pre>
# A very quick run for demonstration purposes
trees <- MaximizeParsimony(dataset, ratchIter = 0, startIter = 0,</pre>
                           tbrIter = 1, maxHits = 4, maxTime = 1/100,
                           concavity = 10, verbosity = 4)
names(trees)
# In actual use, be sure to check that the score has converged on a global
# optimum, conducting additional iterations and runs as necessary.
if (interactive()) {
# Jackknife resampling
nReplicates <- 10
jackTrees <- replicate(nReplicates,</pre>
  #c() ensures that each replicate returns a list of trees
  c(Resample(dataset, trees, ratchIter = 0, tbrIter = 2, startIter = 1,
             maxHits = 5, maxTime = 1 / 10,
             concavity = 10, verbosity = 0))
 )
# In a serious analysis, more replicates would be conducted, and each
# search would undergo more iterations.
\# Now we must decide what to do with the multiple optimal trees from
# each replicate.
# Treat each tree equally
JackLabels(ape::consensus(trees), unlist(jackTrees, recursive = FALSE))
# Take the strict consensus of all trees for each replicate
JackLabels(ape::consensus(trees), lapply(jackTrees, ape::consensus))
```

MinimumLength 27

MinimumLength

Minimum length

Description

The smallest length that a character can obtain on any tree.

Usage

```
MinimumLength(x, compress = FALSE)
## S3 method for class 'phyDat'
MinimumLength(x, compress = FALSE)
## S3 method for class 'numeric'
MinimumLength(x, compress = NA)
## S3 method for class 'character'
MinimumLength(x, compress = TRUE)
MinimumSteps(x)
```

Arguments

Χ

An object of class phyDat; or a string to be coerced to a phyDat object via TreeTools::StringToPhyDat(); or an integer vector listing the tokens that may be present at each tip along a single character, with each token represented as a binary digit; e.g. a value of $11 (= 2^0 + 2^1 + 2^3)$ means that the tip may have tokens 0, 1 or 3.

Inapplicable tokens should be denoted with the integer \emptyset (not 2^0).

compress

Logical specifying whether to retain the compression of a phyDat object or to return a vector specifying to each individual character, decompressed using the dataset's index attribute.

Value

MinimumLength() returns a vector of integers specifying the minimum number of steps that each character must contain.

Author(s)

```
Martin R. Smith (martin.smith@durham.ac.uk)
```

See Also

```
Other tree scoring: CharacterLength(), IWScore(), MorphyTreeLength()
```

Examples

```
data('inapplicable.datasets')
myPhyDat <- inapplicable.phyData[[4]]

# load your own data with
# my.PhyDat <- as.phyDat(read.nexus.data('filepath'))
# or Windows users can select a file interactively using:
# my.PhyDat <- as.phyDat(read.nexus.data(choose.files()))

class(myPhyDat) # phyDat object

# Minimum length of each character in turn
MinimumLength(myPhyDat)

# Collapse duplicate characters, per phyDat compression
MinimumLength(myPhyDat, compress = TRUE)

# Calculate length of a single character from its textual representation
MinimumLength('-{-1}{-2}{-3}2233')</pre>
```

MorphyBootstrap

Parsimony Ratchet

Description

Ratchet() uses the parsimony ratchet (Nixon 1999) to search for a more parsimonious tree using custom optimality criteria.

Usage

```
MorphyBootstrap(
  edgeList,
  morphyObj,
  EdgeSwapper = NNISwap,
  maxIter,
  maxHits,
  verbosity = 1L,
  stopAtPeak = FALSE,
  stopAtPlateau = 0L,
   ...
)
Ratchet(
  tree,
```

```
dataset,
  InitializeData = PhyDat2Morphy,
  CleanUpData = UnloadMorphy,
  TreeScorer = MorphyLength,
  Bootstrapper = MorphyBootstrap,
  swappers = list(TBRSwap, SPRSwap, NNISwap),
 BootstrapSwapper = if (is.list(swappers)) swappers[[length(swappers)]] else swappers,
  returnAll = FALSE,
  stopAtScore = NULL,
  stopAtPeak = FALSE,
  stopAtPlateau = 0L,
  ratchIter = 100,
  ratchHits = 10,
  searchIter = 4000,
  searchHits = 42,
  bootstrapIter = searchIter,
  bootstrapHits = searchHits,
  verbosity = 1L,
  suboptimal = sqrt(.Machine$double.eps),
)
MultiRatchet(
  tree,
  dataset,
  ratchHits = 10,
  searchIter = 500,
  searchHits = 20,
  verbosity = 0L,
  swappers = list(RootedNNISwap),
  nSearch = 10,
  stopAtScore = NULL,
)
RatchetConsensus(
  tree,
  dataset,
  ratchHits = 10,
  searchIter = 500,
  searchHits = 20,
  verbosity = 0L,
  swappers = list(RootedNNISwap),
  nSearch = 10,
  stopAtScore = NULL,
```

Arguments

 ${\tt edgeList}$

a list containing the following: - vector of integers corresponding to the parent of each edge in turn - vector of integers corresponding to the child of each edge in turn - (optionally) score of the tree - (optionally, if score provided) number of

times this score has been hit

morphy0bj Object of class morphy, perhaps created with PhyDat2Morphy().

EdgeSwapper a function that rearranges a parent and child vector, and returns a list with mod-

ified vectors; for example SPRSwap().

maxIter Numeric specifying maximum number of iterations to perform in tree search.

Numeric specifying maximum number of hits to accomplish in tree search.

verbosity Numeric specifying level of detail to display in console: larger numbers provide

more verbose feedback to the user.

stopAtPeak Logical specifying whether to terminate search once a subsequent iteration re-

covers a sub-optimal score. Will be overridden if a passed function has an attribute stopAtPeak set by attr(FunctionName, 'stopAtPeak') <-TRUE.

stopAtPlateau Integer. If > 0, tree search will terminate if the score has not improved after

stopAtPlateau iterations. Will be overridden if a passed function has an attribute stopAtPlateau set by attr(FunctionName, 'stopAtPlateau') <-TRUE.

... further arguments to pass to TreeScorer(), e.g. dataset = .

tree A tree of class phylo.

dataset a dataset in the format required by TreeScorer().

InitializeData Function that sets up data object to prepare for tree search. The function will be

passed the dataset parameter. Its return value will be passed to TreeScorer()

and CleanUpData().

CleanUpData Function to destroy data object on function exit. The function will be passed the

value returned by InitializeData().

TreeScorer function to score a given tree. The function will be passed three parameters,

corresponding to the parent and child entries of a tree's edge list, and a dataset.

Bootstrapper Function to perform bootstrapped rearrangements of tree. First arguments will

be an edgeList and a dataset, initialized using InitializeData(). Should

return a rearranged edgeList.

swappers A list of functions to use to conduct edge rearrangement during tree search.

Provide functions like NNISwap to shuffle root position, or RootedTBRSwap if the position of the root should be retained. You may wish to use extreme swappers (such as TBR) early in the list, and a more subtle rearranger (such as NNI) later in the list to make incremental tinkerings once an almost-optimal tree has been

found.

BootstrapSwapper

Function such as RootedNNISwap to use to rearrange trees within Bootstrapper().

returnAll Set to TRUE to report all MPTs encountered during the search, perhaps to analyse

consensus.

stopAtScore stop search as soon as this score is hit or beaten.

ratchIter Stop when this many ratchet iterations have been performed.

ratchHits Stop when this many ratchet iterations have found the same best score.

searchIter Integer specifying maximum rearrangements to perform on each bootstrap or

ratchet iteration. To override this value for a single swapper function, set e.g.

attr(SwapperFunction, 'searchIter') <-99</pre>

searchHits Integer specifying maximum times to hit best score before terminating a tree

search within a ratchet iteration. To override this value for a single swapper

function, set e.g. attr(SwapperFunction, 'searchHits') <-99</pre>

bootstrapIter Integer specifying maximum rearrangements to perform on each bootstrap iteration (default: searchIter).

bootstrapHits Integer specifying maximum times to hit best score on each bootstrap iteration (default: searchHits).

suboptimal retain trees that are suboptimal by this score. Defaults to a small value that will

counter rounding errors.

nSearch Number of Ratchet searches to conduct (for RatchetConsensus())

Details

For usage pointers, see the vignette.

Value

MorphyBootstrap() returns a tree that is optimal under a random sampling of the original characters.

Ratchet() returns a tree modified by parsimony ratchet iterations.

MultiRatchet() returns a list of optimal trees produced by nSearch ratchet searches, from which a consensus tree can be generated using ape::consensus() or TreeTools::ConsensusWithout().

Functions

• RatchetConsensus: deprecated alias for MultiRatchet()

Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

References

```
Nixon KC (1999). "The Parsimony Ratchet, a new method for rapid parsimony analysis." Cladistics, 15(4), 407–414. ISSN 0748-3007,
```

\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.1111/j.1096-0031.1999.tb00277.x")}

See Also

• Adapted from pratchet() in the **phangorn** package.

Other custom search functions: EdgeListSearch(), Jackknife(), SuccessiveApproximations()

Examples

```
data('Lobo', package = 'TreeTools')
njtree <- TreeTools::NJTree(Lobo.phy)
# Increase value of ratchIter and searchHits to do a proper search
quickResult <- Ratchet(njtree, Lobo.phy, ratchIter = 2, searchHits = 3)
# Plot result (legibly)
oldPar <- par(mar = rep(0, 4), cex = 0.75)
plot(quickResult)
par(oldPar)</pre>
```

32 MorphyWeights

MorphyWeights

Set and get the character weightings associated with a Morphy object.

Description

MorphyWeights() details the approximate and exact weights associated with characters in a Morphy object; SetMorphyWeights() edits them.

Usage

```
MorphyWeights(morphyObj)
SetMorphyWeights(weight, morphyObj, checkInput = TRUE)
```

Arguments

morphyObj Object of class morphy, perhaps created with PhyDat2Morphy(). weight A vector listing the new weights to be applied to each character

checkInput Whether to sanity-check input data before applying. Defaults to TRUE to protect

the user from crashes.

Value

MorphyWeights() returns a data frame with two named rows and one column per character pattern: row 1, approx, is a list of integers specifying the approximate (integral) weights used by MorphyLib; row 2, exact, is a list of numerics specifying the exact weights specified by the user.

SetMorphyWeights() returns the Morphy error code generated when applying weight.

Author(s)

```
Martin R. Smith (martin.smith@durham.ac.uk)
```

See Also

```
Other Morphy API functions: GapHandler(), MorphyErrorCheck(), PhyDat2Morphy(), SingleCharMorphy(), UnloadMorphy(), is.morphyPtr(), mpl_apply_tipdata(), mpl_attach_rawdata(), mpl_attach_symbols(), mpl_delete_Morphy(), mpl_delete_rawdata(), mpl_first_down_recon(), mpl_first_up_recon(), mpl_get_charac_weight(), mpl_get_gaphandl(), mpl_get_num_charac(), mpl_get_num_internal_nodes(), mpl_get_numtaxa(), mpl_get_symbols(), mpl_init_Morphy(), mpl_new_Morphy(), mpl_second_down_recon(), mpl_second_up_recon(), mpl_set_charac_weight(), mpl_set_num_internal_nodes(), mpl_set_parsim_t(), mpl_translate_error(), mpl_update_lower_root(), mpl_update_tip(), summary.morphyPtr()
```

Examples

```
tokens <- matrix(c(
  0, 0, 0, 1, 1, 2,
  0, 0, 0, 0, 0, 0), byrow = TRUE, nrow = 2L,
  dimnames = list(letters[1:2], NULL))
pd <- TreeTools::MatrixToPhyDat(tokens)
morphyObj <- PhyDat2Morphy(pd)
MorphyWeights(morphyObj)</pre>
```

NNI 33

```
if (SetMorphyWeights(c(1, 1.5, 2/3), morphyObj) != 0L) message("Errored") MorphyWeights(morphyObj) morphyObj <- UnloadMorphy(morphyObj)
```

NNI

Nearest neighbour interchange (NNI)

Description

NNI()performs a single iteration of the nearest-neighbour interchange algorithm; RootedNNI() retains the position of the root. These functions are based on equivalents in the **phangorn** package. cnnI() is an equivalent function coded in C, that runs much faster.

Usage

Arguments

tree	A tree of class phylo.
edgeToBreak	In (Rooted)NNI(), an optional integer specifying the index of an edge to bisect/prune, generated randomly if not specified. If -1, a complete list of all trees one step from the input tree will be returned. In cNNI(), an integer from zero to nEdge(tree) -nTip(tree) -2, specifying which internal edge to break.
whichSwitch	Integer from zero to one, specifying which way to re-build the broken internal edge.
parent	Integer vector corresponding to the first column of the edge matrix of a tree of class phylo, i.e. tree\$edge[,1].
child	Integer vector corresponding to the second column of the edge matrix of a tree of class phylo, i.e. tree\$edge[,2].
nTips	(optional) Number of tips.

Details

Branch lengths are not supported.

All nodes in a tree must be bifurcating; ape::collapse.singles() and ape::multi2di() may help.

NNI NNI

Value

Returns a tree with class phylo (if returnAll = FALSE) or a set of trees, with class multiPhylo (if returnAll = TRUE).

cNNI() returns a tree of class phylo, rooted on the same leaf, on which the specified rearrangement has been conducted.

NNISwap() returns a list containing two elements, corresponding in turn to the rearranged parent and child parameters.

a list containing two elements, corresponding in turn to the rearranged parent and child parameters

Functions

- NNISwap: faster version that takes and returns parent and child parameters
- RootedNNI: Perform NNI rearrangement, retaining position of root
- RootedNNISwap: faster version that takes and returns parent and child parameters

Author(s)

```
Martin R. Smith (martin.smith@durham.ac.uk)
```

References

The algorithm is summarized in Felsenstein J (2004). *Inferring phylogenies*. Sinauer Associates, Sunderland, Massachusetts.

See Also

Other tree rearrangement functions: SPR(), TBR()

Examples

```
tree <- TreeTools::BalancedTree(8)</pre>
# A random rearrangement
NNI(tree)
cNNI(tree)
# All trees one NNI rearrangement away
NNI(tree, edgeToBreak = -1)
# Manual random sampling
cNNI(tree, sample.int(14 - 8 - 1, 1), sample.int(2, 1))
# A specified rearrangement
cNNI(tree, 0, 0)
# If a tree may not be binary, collapse nodes with
tree <- TreeTools::MakeTreeBinary(tree)</pre>
# If a tree may be improperly rooted, use
tree <- TreeTools::RootTree(tree, 1)</pre>
# If a tree may exhibit unusual node ordering, this can be addressed with
tree <- TreeTools::Preorder(tree)</pre>
```

PhyDat2Morphy 35

Initialize a Morphy object from a phyDat *object*

Description

Creates a new Morphy object with the same size and characters as the phyDat object. Once finished with the object, it should be destroyed using UnloadMorphy() to free the allocated memory.

Usage

```
PhyDat2Morphy(phy, gap = "inapplicable")
```

Arguments

phy An object of **phangorn** class phyDat.

gap An unambiguous abbreviation of inapplicable, ambiguous (= missing), or

extra state, specifying how gaps will be handled.

Value

PhyDat2Morphy() returns a pointer to an initialized Morphy object.

Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

See Also

```
Other Morphy API functions: GapHandler(), MorphyErrorCheck(), MorphyWeights(), SingleCharMorphy(), UnloadMorphy(), is.morphyPtr(), mpl_apply_tipdata(), mpl_attach_rawdata(), mpl_attach_symbols(), mpl_delete_Morphy(), mpl_delete_rawdata(), mpl_first_down_recon(), mpl_first_up_recon(), mpl_get_charac_weight(), mpl_get_gaphandl(), mpl_get_num_charac(), mpl_get_num_internal_nodes(), mpl_get_numtaxa(), mpl_get_symbols(), mpl_init_Morphy(), mpl_new_Morphy(), mpl_second_down_recon(), mpl_second_up_recon(), mpl_set_charac_weight(), mpl_set_num_internal_nodes(), mpl_set_parsim_t(), mpl_translate_error(), mpl_update_lower_root(), mpl_update_tip(), summary.morphyPtr()
```

Examples

```
data('Lobo', package='TreeTools')
morphyObj <- PhyDat2Morphy(Lobo.phy)
# Set object to be destroyed at end of session or closure of function
# on.exit(morphyObj <- UnloadMorphy(morphyObj), add = TRUE)
# Do something with pointer
# ....
# Or, instead of on.exit, manually destroy morphy object and free memory:
morphyObj <- UnloadMorphy(morphyObj)</pre>
```

36 PlotCharacter

PlotCharacter

Plot the distribution of a character on a tree

Description

Reconstructs the distribution of a character on a tree topology using the modified Fitch algorithm presented in Brazeau et al. (2019).

Usage

```
PlotCharacter(
    tree,
    dataset,
    char = 1L,
    updateTips = FALSE,
    plot = TRUE,
    tokenCol = NULL,
    ambigCol = "grey",
    inappCol = "lightgrey",
    ambigLty = "dotted",
    inappLty = "dashed",
    plainLty = par("lty"),
    tipOffset = 1,
    unitEdge = FALSE,
    ...
)
```

Arguments

tree A tree of class phylo.

dataset A phylogenetic data matrix of **phangorn** class phyDat, whose names corre-

spond to the labels of any accompanying tree.

char Index of character to plot.

updateTips Logical; if FALSE, tips will be labelled with their original state in dataset.

plot Logical specifying whether to plot the output.

tokenCol Palette specifying colours to associate with each token in turn, in the sequence

listed in attr(dataset, 'levels').

ambigCol, ambigLty, inappCol, inappLty, plainLty

Colours and line types to apply to ambiguous, inapplicable and applicable tokens. See the 1ty graphical parameter for details of line styles. Overrides

tokenCol.

tipOffset Numeric: how much to offset tips from their labels.
unitEdge Logical: Should all edges be plotted with a unit length?

... Further arguments to pass to plot.phylo().

Value

PlotCharacter() returns a matrix in which each row corresponds to a numbered tip or node of tree, and each column corresponds to a token; the tokens that might parsimoniously be present at each point on a tree are denoted with TRUE.

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Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

References

Brazeau MD, Guillerme T, Smith MR (2019). "An algorithm for morphological phylogenetic analysis with inapplicable data." *Systematic Biology*, **68**(4), 619–631.

```
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1093/sysbio/syy083")}
```

Examples

```
# Set up plotting area
oPar <- par(mar = rep(0, 4))

tree <- ape::read.tree(text =
   "((((((a, b), c), d), e), f), (g, (h, (i, (j, (k, 1)))));")
## A character with inapplicable data
dataset <- TreeTools::StringToPhyDat("23--1??--032", tips = tree)
PlotCharacter(tree, dataset)

# Character from a real dataset
data("Lobo", package = "TreeTools")
dataset <- Lobo.phy
tree <- TreeTools::NJTree(dataset)
PlotCharacter(tree, dataset, 14)
par(oPar)</pre>
```

PrepareDataProfile

Prepare data for Profile Parsimony

Description

Calculates profiles for each character in a dataset. Will also simplify characters, with a warning, where they are too complex for the present implementation of profile parsimony:

- inapplicable tokens will be replaced with the ambiguous token (i.e. \rightarrow ?);
- Ambiguous tokens will be treated as fully ambiguous (i.e. $\{02\} \rightarrow ?$)
- Where more than two states are informative (i.e. unambiguously present in more than one taxon), states beyond the two most informative will be ignored.

Usage

```
PrepareDataProfile(dataset)
PrepareDataIW(dataset)
```

Arguments

dataset dataset of class phyDat

38 profiles

Value

An object of class phyDat, with additional attributes. PrepareDataProfile adds the attributes:

- info.amounts: details the information represented by each character when subject to N additional steps.
- informative: logical specifying which characters contain any phylogenetic information.
- bootstrap: The character vector c('info.amounts', 'split.sizes'), indicating attributes to sample when bootstrapping the dataset (e.g. in Ratchet searches).

PrepareDataIW adds the attribute:

• min.length: The minimum number of steps that must be present in each transformation series.

Functions

• PrepareDataIW: Prepare data for implied weighting

Author(s)

```
Martin R. Smith; written with reference to phangorn:::prepareDataFitch()
```

See Also

Other profile parsimony functions: Carter1(), StepInformation(), WithOneExtraStep(), profiles

Examples

```
data('congreveLamsdellMatrices')
dataset <- congreveLamsdellMatrices[[42]]
PrepareDataProfile(dataset)</pre>
```

profiles

Empirically counted profiles for small trees

Description

The base 2 logarithm of the number of trees containing *s* steps, calculated by scoring a character on each *n*-leaf tree.

Usage

profiles

Format

A list with the structure profiles[[number of leaves]][[number of tokens]][[tokens in smallest split]] The list entry returns a named numeric vector; each entry lists log2(proportion of n-leaf trees with s or fewer steps for this character).

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See Also

```
Other profile parsimony functions: Carter1(), PrepareDataProfile(), StepInformation(), WithOneExtraStep()
```

Examples

```
data(profiles)
# Load profile for a character of the structure 0 0 0 1 1 1 1 1
profile3.5 <- profiles[[8]][[2]][[3]]
# Number of trees with _s_ or fewer steps on that character
TreeTools::NUnrooted(8) * 2 ^ profile3.5</pre>
```

 ${\tt RandomMorphyTree}$

Random postorder tree

Description

Random postorder tree

Usage

RandomMorphyTree(nTip)

Arguments

nTip

Integer specifying the number of tips to include in the tree (minimum 2).

Value

A list with three elements, each a vector of integers, respectively containing:

- The parent of each tip and node, in order
- The left child of each node
- The right child of each node.

See Also

Other tree generation functions: AdditionTree()

40 RearrangeEdges

RandomTreeScore

Parsimony score of random postorder tree

Description

Parsimony score of random postorder tree

Usage

```
RandomTreeScore(morphyObj)
```

Arguments

```
morphy0bj Object of class morphy, perhaps created with PhyDat2Morphy().
```

Value

RandomTreeScore() returns the parsimony score of a random tree for the given Morphy object.

Examples

```
tokens <- matrix(c(
    0, '-', '-', 1, 1, 2,
    0, 1, 0, 1, 2, 2,
    0, '-', '-', 0, 0, 0), byrow = TRUE, nrow = 3L,
    dimnames = list(letters[1:3], NULL))
pd <- TreeTools::MatrixToPhyDat(tokens)
morphyObj <- PhyDat2Morphy(pd)

RandomTreeScore(morphyObj)

morphyObj <- UnloadMorphy(morphyObj)</pre>
```

RearrangeEdges

Rearrange edges of a phylogenetic tree

Description

RearrangeEdges() performs the specified edge rearrangement on a matrix that corresponds to the edges of a phylogenetic tree, returning the score of the new tree. Will generally be called from within a tree search function.

Usage

```
RearrangeEdges(
  parent,
  child,
  dataset,
  TreeScorer = MorphyLength,
  EdgeSwapper,
  scoreToBeat = TreeScorer(parent, child, dataset, ...),
```

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```
iter = "?",
hits = 0L,
verbosity = 0L,
...
)
```

Arguments

parent Integer vector corresponding to the first column of the edge matrix of a tree of class phylo, i.e. tree\$edge[,1]. child Integer vector corresponding to the second column of the edge matrix of a tree of class phylo, i.e. tree\$edge[,2]. dataset Third argument to pass to TreeScorer. TreeScorer function to score a given tree. The function will be passed three parameters, corresponding to the parent and child entries of a tree's edge list, and a dataset. a function that rearranges a parent and child vector, and returns a list with mod-EdgeSwapper ified vectors; for example SPRSwap(). scoreToBeat Double giving score of input tree. iter iteration number of calling function, for reporting to user only. Integer giving number of times the input tree has already been hit. hits Numeric specifying level of detail to display in console: larger numbers provide verbosity more verbose feedback to the user. further arguments to pass to TreeScorer(), e.g. dataset = .

Details

RearrangeTree() performs one tree rearrangement of a specified type, and returns the score of the tree (with the given dataset). It also reports the number of times that this score was hit in the current function call.

Value

This function returns a list with two to four elements, corresponding to a binary tree: - 1. Integer vector listing the parent node of each edge; - 2. Integer vector listing the child node of each edge; - 3. Score of the tree; - 4. Number of times that score has been hit.

Author(s)

Martin R. Smith

```
data('Lobo', package='TreeTools')
tree <- TreeTools::NJTree(Lobo.phy)
edge <- tree$edge
parent <- edge[, 1]
child <- edge[, 2]
dataset <- PhyDat2Morphy(Lobo.phy)
RearrangeEdges(parent, child, dataset, EdgeSwapper = RootedNNISwap)
# Remember to free memory:
dataset <- UnloadMorphy(dataset)</pre>
```

42 referenceTree

referenceTree

Tree topology for matrix simulation

Description

The tree topology used to generate the matrices in congreveLamsdellMatrices

Usage

referenceTree

Format

A single phylogenetic tree saved as an object of class phylo

Source

```
Congreve & Lamsdell (2016); doi: 10.1111/pala.12236
```

References

Congreve CR, Lamsdell JC (2016). "Implied weighting and its utility in palaeontological datasets: a study using modelled phylogenetic matrices." *Palaeontology*, **59**(3), 447–465.

```
\Sexpr[results=rd, stage=build]{tools:::Rd_expr_doi("10.1111/pala.12236")}
```

. Congreve CR, Lamsdell JC (2016). "Data from: Implied weighting and its utility in palaeon-tological datasets: a study using modelled phylogenetic matrices." *Dryad Digital Repository*, doi:10.5061/dryad.7dq0j.

```
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.5061/dryad.7dq0j")}
```

```
data(referenceTree)
plot(referenceTree)
```

SingleCharMorphy 43

Description

Morphy object from single character

Usage

```
SingleCharMorphy(char, gap = "inapp")
```

Arguments

char	State of each character at each tip in turn, in a format that will be converted to a character string by paste@(char,';',collapse='').
gap	An unambiguous abbreviation of inapplicable, ambiguous (= missing), or extra state, specifying how gaps will be handled.

Value

A pointer to an object of class morphy0bj. Don't forget to unload it when you've finished with it.

Author(s)

```
Martin R. Smith (martin.smith@durham.ac.uk)
```

See Also

```
Score a tree: MorphyTreeLength()

Other Morphy API functions: GapHandler(), MorphyErrorCheck(), MorphyWeights(), PhyDat2Morphy(),
UnloadMorphy(), is.morphyPtr(), mpl_apply_tipdata(), mpl_attach_rawdata(), mpl_attach_symbols(),
mpl_delete_Morphy(), mpl_delete_rawdata(), mpl_first_down_recon(), mpl_first_up_recon(),
mpl_get_charac_weight(), mpl_get_gaphandl(), mpl_get_num_charac(), mpl_get_num_internal_nodes(),
mpl_get_numtaxa(), mpl_get_symbols(), mpl_init_Morphy(), mpl_new_Morphy(), mpl_second_down_recon(),
mpl_second_up_recon(), mpl_set_charac_weight(), mpl_set_num_internal_nodes(), mpl_set_parsim_t(),
mpl_translate_error(), mpl_update_lower_root(), mpl_update_tip(), summary.morphyPtr()
```

```
morphyObj <- SingleCharMorphy('-0-0', gap = 'Extra')
RandomTreeScore(morphyObj)
morphyObj <- UnloadMorphy(morphyObj)</pre>
```

44 SiteConcordance

SiteConcordance

Calculate site concordance factor

Description

The site concordance factor (Minh et al. 2020) is a measure of the strength of support that the dataset presents for a given split in a tree.

Usage

```
QuartetConcordance(tree, dataset = NULL)
ClusteringConcordance(tree, dataset)
PhylogeneticConcordance(tree, dataset)
MutualClusteringConcordance(tree, dataset)
SharedPhylogeneticConcordance(tree, dataset)
```

Arguments

tree A tree of class phylo.

dataset A phylogenetic data matrix of **phangorn** class phyDat, whose names corre-

spond to the labels of any accompanying tree.

Details

QuartetConcordance() is the proportion of quartets (sets of four leaves) that are decisive for a split which are also concordant with it. For example, a quartet with the characters $0\,0\,0\,1$ is not decisive, as all relationships between those leaves are equally parsimonious. But a quartet with characters $0\,0\,1\,1$ is decisive, and is concordant with any tree that groups the first two leaves together to the exclusion of the second.

NOTE: These functions are under development, and may be incompletely tested or change without notice. Complete documentation and discussion will follow soon.

Author(s)

```
Martin R. Smith (martin.smith@durham.ac.uk)
```

References

Minh BQ, Hahn MW, Lanfear R (2020). "New methods to calculate concordance factors for phylogenomic datasets." *Molecular Biology and Evolution*, **37**(9), 2727–2733.

```
\Sexpr[results=rd,stage=build]{tools:::Rd_expr_doi("10.1093/molbev/msaa106")}
```

See Also

Other split support functions: JackLabels(), Jackknife(), MaximizeParsimony()

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Examples

```
data('congreveLamsdellMatrices', package = 'TreeSearch')
dataset <- congreveLamsdellMatrices[[1]][, 1:20]
tree <- referenceTree
qc <- QuartetConcordance(tree, dataset)
cc <- ClusteringConcordance(tree, dataset)
pc <- PhylogeneticConcordance(tree, dataset)
spc <- SharedPhylogeneticConcordance(tree, dataset)
mcc <- MutualClusteringConcordance(tree, dataset)

OPar <- par(mar = rep(0, 4), cex = 0.8)
plot(tree)
TreeTools::LabelSplits(tree, signif(qc, 3))
TreeTools::LabelSplits(tree, signif(cc, 3))
TreeTools::LabelSplits(tree, signif(pc, 3))
par(oPar)

pairs(cbind(qc, cc, pc, spc, mcc))</pre>
```

SPR

Subtree pruning and rearrangement (SPR)

Description

Perform one SPR rearrangement on a tree

Usage

```
SPR(tree, edgeToBreak = NULL, mergeEdge = NULL)
SPRMoves(tree, edgeToBreak = integer(0))
## S3 method for class 'phylo'
SPRMoves(tree, edgeToBreak = integer(0))
## S3 method for class 'matrix'
SPRMoves(tree, edgeToBreak = integer(0))
SPRSwap(
  parent,
  child,
  nEdge = length(parent),
  nNode = nEdge/2L,
  edgeToBreak = NULL,
  mergeEdge = NULL
)
RootedSPR(tree, edgeToBreak = NULL, mergeEdge = NULL)
RootedSPRSwap(
  parent,
  child,
```

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```
nEdge = length(parent),
nNode = nEdge/2L,
edgeToBreak = NULL,
mergeEdge = NULL
)
```

Arguments

tree A tree of class phylo.

edgeToBreak the index of an edge to bisect, generated randomly if not specified.

mergeEdge the index of an edge on which to merge the broken edge.

parent Integer vector corresponding to the first column of the edge matrix of a tree of

class phylo, i.e. tree\$edge[,1].

child Integer vector corresponding to the second column of the edge matrix of a tree

of class phylo, i.e. tree\$edge[,2].

nEdge (optional) integer specifying the number of edges of a tree of class phylo, i.e.

dim(tree\$edge)[1]

nNode (optional) Number of nodes.

Details

Equivalent to kSPR() in the **phangorn** package, but faster. Note that rearrangements that only change the position of the root WILL be returned by SPR. If the position of the root is irrelevant (as in Fitch parsimony, for example) then this function will occasionally return a functionally equivalent topology. RootIrrelevantSPR will search tree space more efficiently in these cases. Branch lengths are not (yet) supported.

All nodes in a tree must be bifurcating; ape::collapse.singles and ape::multi2di may help.

Value

This function returns a tree in phyDat format that has undergone one SPR iteration.

TBRMoves() returns a list of all trees one SPR move away from tree, with edges and nodes in preorder, rooted on the first-labelled tip.

a list containing two elements, corresponding in turn to the rearranged parent and child parameters a list containing two elements, corresponding in turn to the rearranged parent and child parameters

Functions

- SPRSwap: faster version that takes and returns parent and child parameters
- RootedSPR: Perform SPR rearrangement, retaining position of root
- RootedSPRSwap: faster version that takes and returns parent and child parameters

Author(s)

Martin R. Smith

References

The SPR algorithm is summarized in Felsenstein J (2004). *Inferring phylogenies*. Sinauer Associates, Sunderland, Massachusetts.

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See Also

• RootedSPR(): useful when the position of the root node should be retained.

Other tree rearrangement functions: NNI(), TBR()

Examples

```
{
tree <- ape::rtree(20, br=FALSE)
SPR(tree)
}</pre>
```

StepInformation

Information content of a character known to contain e steps

Description

StepInformation() calculates the phylogenetic information content of a character char when e extra steps are present, for all possible values of e.

Usage

```
StepInformation(char, ambiguousTokens = c("-", "?"))
```

Arguments

char Vector of tokens listing states for the character in question. ambiguousTokens

Vector specifying which tokens, if any, correspond to the ambiguous token (?).

Details

Calculates the number of trees consistent with the character having e extra steps, where e ranges from its minimum possible value (i.e. number of different tokens minus one) to its maximum.

Value

StepInformation() returns a numeric vector detailing the amount of phylogenetic information (in bits) associated with the character when 0, 1, 2... extra steps are present. The vector is named with the total number of steps associated with each entry in the vector: for example, a character with three observed tokens must exhibit two steps, so the first entry (zero extra steps) is named 2 (two steps observed).

Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

See Also

```
Other profile parsimony functions: Carter1(), PrepareDataProfile(), WithOneExtraStep(), profiles
```

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Examples

```
character <- rep(c(0:3, '?', '-'), c(8, 5, 1, 1, 2, 2))
StepInformation(character)
```

summary.morphyPtr

Details the attributes of a morphy object

Description

Details the attributes of a morphy object

Usage

```
## S3 method for class 'morphyPtr'
summary(object, ...)
```

Arguments

```
object A Morphy object
... any other parameters...
```

Value

A list detailing the number of taxa, internal nodes, and characters and their weights.

Author(s)

Martin R. Smith (martin.smith@durham.ac.uk)

See Also

```
Other Morphy API functions: GapHandler(), MorphyErrorCheck(), MorphyWeights(), PhyDat2Morphy(), SingleCharMorphy(), UnloadMorphy(), is.morphyPtr(), mpl_apply_tipdata(), mpl_attach_rawdata(), mpl_attach_symbols(), mpl_delete_Morphy(), mpl_delete_rawdata(), mpl_first_down_recon(), mpl_first_up_recon(), mpl_get_charac_weight(), mpl_get_gaphandl(), mpl_get_num_charac(), mpl_get_num_internal_nodes(), mpl_get_numtaxa(), mpl_get_symbols(), mpl_init_Morphy(), mpl_new_Morphy(), mpl_second_down_recon(), mpl_second_up_recon(), mpl_set_charac_weight(), mpl_set_num_internal_nodes(), mpl_set_parsim_t(), mpl_translate_error(), mpl_update_lower_root(), mpl_update_tip()
```

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TBR

Tree bisection and reconnection (TBR)

Description

TBR performs a single random TBR iteration.

Usage

```
TBR(tree, edgeToBreak = NULL, mergeEdges = NULL)
TBRMoves(tree, edgeToBreak = integer(0))
## S3 method for class 'phylo'
TBRMoves(tree, edgeToBreak = integer(0))
## S3 method for class 'matrix'
TBRMoves(tree, edgeToBreak = integer(0))
TBRSwap(
  parent,
  child,
  nEdge = length(parent),
  edgeToBreak = NULL,
  mergeEdges = NULL
RootedTBR(tree, edgeToBreak = NULL, mergeEdges = NULL)
RootedTBRSwap(
  parent,
  child,
  nEdge = length(parent),
  edgeToBreak = NULL,
  mergeEdges = NULL
```

Arguments

tree

A bifurcating tree of class phylo, with all nodes resolved; edgeToBreak (optional) integer specifying the index of an edge to bisect/prune, generated randomly if not specified. Alternatively, set to -1 to return a complete list of all

trees one step from the input tree.

(optional) vector of length 1 or 2, listing edge(s) to be joined: In SPR, this mergeEdges

is where the pruned subtree will be reconnected. In TBR, these edges will be reconnected (so must be on opposite sides of edgeToBreak); if only a single

edge is specified, the second will be chosen at random

Integer vector corresponding to the first column of the edge matrix of a tree of parent

class phylo, i.e. tree\$edge[,1].

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child Integer vector corresponding to the second column of the edge matrix of a tree of class phylo, i.e. tree\$edge[,2].

nEdge (optional) Number of edges.

Details

Branch lengths are not (yet) supported.

All nodes in a tree must be bifurcating; ape::collapse.singles and ape::multi2di may help.

Value

This function returns a tree in phyDat format that has undergone one TBR iteration.

TBRMoves() returns a multiPhylo object listing all trees one TBR move away from tree, with edges and nodes in preorder, rooted on the first-labelled tip.

a list containing two elements, corresponding in turn to the rearranged parent and child parameters

Functions

- TBRSwap: faster version that takes and returns parent and child parameters
- RootedTBR: Perform TBR rearrangement, retaining position of root
- RootedTBRSwap: faster version that takes and returns parent and child parameters

Author(s)

Martin R. Smith

References

The TBR algorithm is summarized in Felsenstein J (2004). *Inferring phylogenies*. Sinauer Associates, Sunderland, Massachusetts.

See Also

```
RootedTBR(): useful when the position of the root node should be retained.
```

Other tree rearrangement functions: NNI(), SPR()

```
{
library('ape')
tree <- rtree(20, br=NULL)
TBR(tree)
}
```

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UnloadMorphy

Destroy a Morphy object

Description

Destroys a previously-created Morphy object.

Usage

UnloadMorphy(morphyObj)

Arguments

morphy0bj

Object of class morphy, perhaps created with PhyDat2Morphy().

Details

Best practice is to call morphy0bj <-UnloadMorphy(morphy0bj) Failure to do so will cause a crash if UnloadMorphy() is called on an object that has already been destroyed

Value

Morphy error code, decipherable using mpl_translate_error

Author(s)

Martin R. Smith

See Also

```
Other Morphy API functions: GapHandler(), MorphyErrorCheck(), MorphyWeights(), PhyDat2Morphy(), SingleCharMorphy(), is.morphyPtr(), mpl_apply_tipdata(), mpl_attach_rawdata(), mpl_attach_symbols(), mpl_delete_Morphy(), mpl_delete_rawdata(), mpl_first_down_recon(), mpl_first_up_recon(), mpl_get_charac_weight(), mpl_get_gaphandl(), mpl_get_num_charac(), mpl_get_num_internal_nodes(), mpl_get_numtaxa(), mpl_get_symbols(), mpl_init_Morphy(), mpl_new_Morphy(), mpl_second_down_recon(), mpl_second_up_recon(), mpl_set_charac_weight(), mpl_set_num_internal_nodes(), mpl_set_parsim_t(), mpl_translate_error(), mpl_update_lower_root(), mpl_update_tip(), summary.morphyPtr()
```

WithOneExtraStep

Number of trees with one extra step

Description

Number of trees with one extra step

Usage

```
WithOneExtraStep(...)
```

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Arguments

... Vector or series of integers specifying the number of leaves bearing each distinct non-ambiguous token.

See Also

```
Other profile parsimony functions: Carter1(), PrepareDataProfile(), StepInformation(), profiles
```

```
WithOneExtraStep(1, 2, 3)
```

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