Package 'treestructure'

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Type Package
Title Detect Population Structure Within Phylogenetic Trees
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Description Algorithms for detecting population structure from the history of coalescent events recorded in phylogenetic trees. This method classifies each tip and internal node of a tree into disjoint sets characterized by similar coalescent patterns. The methods are described in Volz, E., Wiuf, C., Grad, Y., Frost, S., Dennis, A., & Didelot, X. (2020) <doi:10.1093 syaa009="" sysbio="">.</doi:10.1093>
License GPL (>= 2)
Suggests ggtree,ggplot2,knitr
Imports ape (>= 5.0)
LinkingTo Rcpp
VignetteBuilder knitr
RoxygenNote 6.1.0
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R topics documented:
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Plot TreeStructure tree with cluster and partition variables

Description

Plot TreeStructure tree with cluster and partition variables

Usage

```
## S3 method for class 'TreeStructure'
plot(x, use_ggtree = TRUE, ...)
```

Arguments

x A TreeStructure object

use_ggtree Toggle ggtree or ape plotting behaviour

... Additional arguments passed to ggtree or ape::plot.phylo

trestruct

Detect cryptic population structure in time trees

Description

Detect cryptic population structure in time trees

Usage

```
trestruct(tre, minCladeSize = 25, minOverlap = -Inf, nsim = 1000,
  level = 0.01, ncpu = 1, verbosity = 1, debugLevel = 0)
```

Arguments

tre A tree of type ape::phylo. Must be rooted and binary.

minCladeSize All clusters within parititon must have at least this many tips.

minOverlap Threshold time overlap required to find splits in a clade

nsim Number of simulations for computing null distribution of test statistics

level Significance level for finding new split within a set of tips
ncpu If >1 will compute statistics in parallel using multiple CPUs
verbosity If > 0 will print information about progress of the algorithm

debugLevel If > 0 will produce additional data in return value

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Details

Estimates a partition of a time-scaled tree by contrasting coalescent patterns. The algorithm is premised on a Kingman coalescent null hypothesis and a test statistic is formulated based on the rank sum of node times in the tree.

Value

A TreeStructure object which includes cluster and partitition assignment for each tip of the tree.

References

E.M. Volz, Wiuf, C., Grad, Y., Frost, S., Dennis, A., Didelot, X.D. (2020) Identification of hidden population structure in time-scaled phylogenies.

Author(s)

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Examples

```
tree <- ape::rcoal(50)
struct <- trestruct( tree )</pre>
```

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