# Package: treestructure (via r-universe)

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Type Package
Title Detect Population Structure Within Phylogenetic Trees
Version 0.1.0
<b>Date</b> 2020-01-20
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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
<b>Imports</b> ape (>= 5.0)
LinkingTo Rcpp
VignetteBuilder knitr
RoxygenNote 6.1.0
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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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