

Package ‘oncomodel’

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Type Package

Title Maximum likelihood tree models for oncogenesis

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Description Computing probabilistic tree models for oncogenesis based on genetic data using maximum likelihood.

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oncomodel-package *Maximum likelihood tree models for oncogenesis*

Description

Computing probabilistic tree models for oncogenesis based on genetic data using maximum likelihood.

Details

Package: oncomodel
Type: Package
Version: 1.0
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License: GPL version 2 or newer

Author(s)

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References

von Heydebreck A, Gunawan B, Fuezesi L. 2004. Maximum likelihood estimation of oncogenetic tree models. *Biostatistics* 5:545-556.

Examples

```
## NOT RUN
## The example needs longer run time.
#data(kidney)
## Maximum likelihood tree model
#y <- MLtopology(kidney$x)

## Graphical presentation
#y.phyl <- newick2phylog(y$newick)
#plot.phylog(y.phyl, cnodes =1, clabel.n=0.6, f=0.75, sub="Oncogenetic tree of given aberrations")

## Bootstrap confidence values (in percent) and the splits occurring in
## more than 10 percent of the bootstrap data sets
#boot.conf.values(kidney$x, nrep=2)

## Probability for aberration -3|-3p
#leafset.prob(c("-3|-3p", "+5|+5q"), kidney$res)
```

```
## Probability for aberration -3|-3p
#leafset.prob2(c("-3|-3p", "+5|+5q"), kidney$res)
## END(NOT RUN)
```

boot.conf.values *Bootstrap Confidence Values*

Description

Compute the bootstrap confidence values (in percent) for the inner edges and display the splits occurring in > 10 percent of the bootstrap data sets (the splits are characterized by one of the two subsets of leaves).

Usage

```
boot.conf.values(data, random.seed = 12345, nrep = 500)
```

Arguments

data	a binary matrix with rows representing tumors and columns representing genetic alterations.
random.seed	a single value, interpreted as an integer (containing the random number generator (RNG) state for random number generation in R).
nrep	number of replications.

Details

We use the nonparametric bootstrap (Felsenstein, 1985) to assess the uncertainty of properties of the estimated tree model. The proposed tree structure has to be interpreted with caution. Nevertheless we think that the model can at least serve an exploratory purpose, allowing us to formulate hypotheses about the evolution of karyotypes in the data set.

Value

frequencies	the splits occurring in more than 10 percent of the bootstrap data sets.
confidence values	bootstrap confidence values (in percent) for the inner edges.

References

von Heydebreck A, Gunawan B, Fuezesi L. 2004. Maximum likelihood estimation of oncogenetic tree models. *Biostatistics* 5:545-556. Felsenstein, J. 1985. Confidence limits on phylogenies: an approach using the bootstrap. *Evolution* 39:783-791.

Examples

```
## NOT RUN
## The calculation of bootstrap confidence values needs longer run time.
#data(kidney)
#boot.conf.values(kidney$x, nrep=2)
## END(NOT RUN)
```

comp.freq *Compare Model Probabilities to Frequencies*

Description

Compares the model probabilities of single alterations and pairs to the observed frequencies and shows scatterplots for the comparisons.

Usage

```
comp.freq(x, tree, p)
```

Arguments

x a binary data matrix with rows representing tumors and columns representing genetic alterations.

tree the tree in matrix format.

p a vector of edge parameters (model probabilities).

Examples

```
data(kidney)
comp.freq(kidney$x, kidney$res$tree, kidney$res$p)
```

is.tree *Compare with Tree Format*

Description

Tests whether a 2 x n matrix represents a rooted tree in the format accepted by the oncomodel package.

Usage

```
is.tree(tree)
```

Arguments

tree a 2 x n matrix.

Details

In the accepted format, the columns of the integer matrix represent the edges of the tree, with the entry in the first row being closer to the root. The leaves have to be the smallest integers of the matrix. If an edge has a smaller column index than a second one, it may not be on the path from the second edge to the root (the order of the columns has to be compatible with the partial order of the edges of the tree).

Examples

```
data(kidney)
is.tree(kidney$res$tree)
```

kidney

Cytogenetic Data from Kidney Carcinoma

Description

This data set contains cytogenetic data from 173 cases of renal clear cell carcinoma, covering 7 frequent chromosomal aberrations, as well as the corresponding maximum likelihood tree model.

Usage

```
data(kidney)
```

Format

A list with the following components:

- `x` the binary data matrix
- `res` the ML tree model, a list with components:
 - `tree` the tree in matrix format
 - `p` the conditional probabilities associated with the edges
 - `var.names` character vector of variable names
 - `totloglik` the log-likelihood of the tree model.
 - `newick` the tree model in Newick format

leafset.prob	<i>Probability of a Set of Leaves</i>
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Description

Computes the probability of a set of leaves (chromosomal aberrations) in an oncogenetic tree model.

Usage

```
leafset.prob(leafset, y)
```

Arguments

leafset	a character vector of one or more leaves (chromosomal aberrations).
y	the tree model as obtained from MLtopology.

Value

The probability of exactly a given set of leaves (chromosomal aberrations) in the tree.

References

von Heydebreck A, Gunawan B, Fuezesi L. 2004. Maximum likelihood estimation of oncogenetic tree models. *Biostatistics* 5:545-556.

Examples

```
data(kidney)
leafset.prob(c("-3|-3p", "+5|+5q"), kidney$res)
```

leafset.prob2	<i>Probability of a Set of Leaves</i>
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Description

Computes the probability of a set of leaves (chromosomal aberrations) in an oncogenetic tree model.

Usage

```
leafset.prob2(leafset, y)
```

Arguments

leafset	a character vector of one or more leaves (chromosomal aberrations).
y	the tree model as obtained from MLtopology.

Value

The probability of exactly the given set of leaves (chromosomal aberrations) of the tree.

References

von Heydebreck A, Gunawan B, Fuezesi L. 2004. Maximum likelihood estimation of oncogenetic tree models. *Biostatistics* 5:545-556.

Examples

```
data(kidney)
leafset.prob2(c("-3|-3p", "+5|+5q"), kidney$res)
```

MLparameters

Compute Maximum Likelihood Parameters

Description

Computes the maximum likelihood parameters for a given tree topology.

Usage

```
MLparameters(x, tree, freq = NULL)
```

Arguments

x	a binary matrix whose rows are the (preferably unique) genetic profiles.
tree	the tree in matrix format.
freq	a vector whose length equals the number of rows of x, giving the frequency of each profile in the data.

Value

p	a vector of the maximum likelihood edge parameters (model probabilities).
totloglik	the log-likelihood at the ML parameters.

Examples

```
data(kidney)
MLparameters(kidney$x, kidney$res$tree, freq = NULL)
```

`MLtopology`*Compute Maximum Likelihood Tree Topology*

Description

Tries to compute the maximum likelihood tree model for a given data set through stepwise leaf insertion and rearrangements.

Usage

```
MLtopology(x, verbose = FALSE)
```

Arguments

<code>x</code>	a binary matrix with rows representing tumors and columns representing genetic alterations.
<code>verbose</code>	a Boolean value indicating whether intermediate results of the algorithm are to be printed.

Value

A list with the following components:

<code>tree</code>	the resulting tree in matrix format.
<code>p</code>	a vector of the maximum likelihood edge parameters (model probabilities).
<code>totloglik</code>	the log-likelihood of the tree model.
<code>var.names</code>	the character vector with the names of alterations.
<code>newick</code>	the tree model in Newick format.

References

von Heydebreck A, Gunawan B, Fuezesi L. 2004. Maximum likelihood estimation of oncogenetic tree models. *Biostatistics* 5:545-556.

Examples

```
## NOT RUN
## The computation of the maximum likelihood tree model needs longer run time.
#data(kidney)
#y <- MLtopology(kidney$x)
## END(NOT RUN)
```

mrca	<i>Most Recent Common Ancestor</i>
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Description

Computes the most recent common ancestor node for a pair of nodes of a tree.

Usage

```
mrca(x, y, tree)
```

Arguments

x	a node of the tree.
y	a node of the tree.
tree	a tree in matrix format.

Value

the most recent common ancestor node of x and y.

Examples

```
data(kidney)
mrca(1,2, kidney$res$tree)
```

subtree	<i>Subtree</i>
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Description

Computes the subtree rooted at a given node.

Usage

```
subtree(node, tree)
```

Arguments

tree	a tree in matrix format.
node	a node of the tree.

Value

The edge indices of the subtree rooted at the given node.

Examples

```
data(kidney)
subtree(14, kidney$res$tree)
```

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