

Package ‘data.tree’

August 25, 2025

Type Package

Title General Purpose Hierarchical Data Structure

Version 1.2.0

Language en-US

Date 2025-08-06

VignetteBuilder knitr, rmarkdown

Imports R6, stringi, methods

Suggests Formula, graphics, testthat, knitr, rmarkdown, ape, yaml,
networkD3, jsonlite, treemap, party, partykit, doParallel,
foreach, htmlwidgets, DiagrammeR (>= 1.0.0), mockery, rpart

Enhances igraph

Description Create tree structures from hierarchical data, and traverse the
tree in various orders. Aggregate, cumulate, print, plot, convert to and from
data.frame and more. Useful for decision trees, machine learning, finance,
conversion from and to JSON, and many other applications.

License GPL (>= 2)

URL <https://github.com/gluc/data.tree>

BugReports <https://github.com/gluc/data.tree/issues>

Depends R (>= 3.5)

RoxygenNote 7.3.2

Encoding UTF-8

NeedsCompilation no

Author Russ Hyde [ctb] (improve dependencies),
Chris Hammill [ctb] (improve getting),
Facundo Munoz [ctb] (improve list conversion),
Markus Wamser [ctb] (fixed some typos),
Pierre Formont [ctb] (additional features),
Kent Russel [ctb] (documentation),
Noam Ross [ctb] (fixes),
Duncan Garmonsway [ctb] (fixes),
Christoph Glur [aut, cre] (R interface)

Maintainer Christoph Glur <christoph.glur@powerpartners.pro>

Repository CRAN

Date/Publication 2025-08-25 11:00:02 UTC

Contents

acme	3
Aggregate	3
AreNamesUnique	5
as.data.frame.Node	5
as.dendrogram.Node	8
as.igraph.Node	9
as.list.Node	10
as.Node	11
as.Node.BinaryTree	12
as.Node.data.frame	13
as.Node.dendrogram	15
as.Node.list	16
as.Node.party	18
as.Node.phylo	20
as.Node.rpart	21
as.phylo.Node	22
averageBranchingFactor	23
CheckNameReservedWord	23
Climb	24
Clone	25
CreateRandomTree	26
CreateRegularTree	26
Cumulate	27
DefaultPlotHeight	27
Distance	28
Do	28
FindNode	29
FormatFixedDecimal	30
FormatPercent	31
Get	32
GetAttribute	33
GetPhyloNr	34
isLeaf	35
isNotLeaf	36
isNotRoot	36
isRoot	37
mushroom	37
Navigate	38
Node	38
NODE_RESERVED_NAMES_CONST	52
plot.Node	53

<i>acme</i>	3
print.Node	55
Prune	57
Revert	58
Set	58
SetFormat	59
Sort	60
ToNewick	61
Traverse	62
Index	64

<i>acme</i>	<i>Sample Data: A Simple Company with Departments</i>
-------------	---

Description

acme’s tree representation is accessed through its root, acme.

Usage

data(acme)

Format

A data.tree root Node

Details

- cost, only available for leaf nodes. Cost of the project.
- p probability that a project will be undertaken.

<i>Aggregate</i>	<i>Aggregate child values of a Node, recursively.</i>
------------------	---

Description

The Aggregate method lets you fetch an attribute from a Node’s children, and then aggregate them using aggFun. For example, you can aggregate cost by summing costs of child Nodes. This is especially useful in the context of tree traversal, when using post-order traversal mode.

Usage

Aggregate(node, attribute, aggFun, ...)

Arguments

node	the Node on which to aggregate
attribute	determines what is collected. The attribute can be • a.) the name of a field or a property/active of each Node in the tree, e.g. <code>acme\$Get("p")</code> or <code>acme\$Get("position")</code> • b.) the name of a method of each Node in the tree, e.g. <code>acme\$Get("levelZeroBased")</code>, where e.g. <code>acme\$levelZeroBased <- function() acme\$level - 1</code> • c.) a function, whose first argument must be a Node e.g. <code>acme\$Get(function(node) node\$cost * node\$p)</code>
aggFun	the aggregation function to be applied to the children's attributes
...	any arguments to be passed on to attribute (in case it's a function)

Details

As with [Get](#), the attribute can be a field, a method or a function. If the attribute on a child is NULL, Aggregate is called recursively on its children.

See Also

[Node](#)

Examples

```
data(acme)

#Aggregate on a field
Aggregate(acme, "cost", sum)

#This is the same as:
HomeRolledAggregate <- function(node) {
  sum(sapply(node$children, function(child) {
    if (!is.null(child$cost)) child$cost
    else HomeRolledAggregate(child)
  }))
}
HomeRolledAggregate(acme)

#Aggregate using Get
print(acme, "cost", minCost = acme$Get(Aggregate, "cost", min))

#use Aggregate with a function:
Aggregate(acme, function(x) x$cost * x$p, sum)

#cache values along the way
acme$Do(function(x) x$cost <- Aggregate(x, "cost", sum), traversal = "post-order")
acme$IT$cost
```

AreNamesUnique	<i>Test whether all node names are unique.</i>
----------------	--

Description

This can be useful for some conversions.

Usage

```
AreNamesUnique(node)
```

Arguments

node	The root Node of the data.tree structure to test
------	--

Value

TRUE if all Node\$name == TRUE for all nodes in the tree

See Also

as.igraph.Node

Examples

```
data(acme)
AreNamesUnique(acme)
acme$name <- "IT"
AreNamesUnique(acme)
```

as.data.frame.Node	<i>Convert a data.tree structure to a data.frame</i>
--------------------	--

Description

If a node field contains data of length > 1, then that is converted into a string in the data.frame.

Usage

```
## S3 method for class 'Node'
as.data.frame(
  x,
  row.names = NULL,
  optional = FALSE,
  ...,
  traversal = c("pre-order", "post-order", "in-order", "level", "ancestor"),
  pruneFun = NULL,
  filterFun = NULL,
  format = FALSE,
  inheritFromAncestors = FALSE
)

ToDataFrameTree(x, ..., pruneFun = NULL)

ToDataFrameTable(x, ..., pruneFun = NULL)

ToDataFrameNetwork(
  x,
  ...,
  direction = c("climb", "descend"),
  pruneFun = NULL,
  format = FALSE,
  inheritFromAncestors = FALSE
)

ToDataFrameTypeCol(x, ..., type = "level", prefix = type, pruneFun = NULL)
```

Arguments

x	The root Node of the tree or sub-tree to be convert to a data.frame
row.names	NULL or a character vector giving the row names for the data frame. Missing values are not allowed.
optional	logical. If TRUE, setting row names and converting column names (to syntactic names: see <code>make.names</code>) is optional.
...	the attributes to be added as columns of the data.frame. See Get for details. If a specific Node does not contain the attribute, NA is added to the data.frame.
traversal	any of 'pre-order' (the default), 'post-order', 'in-order', 'level', or 'ancestor'. See Traverse for details.
pruneFun	allows providing a prune criteria, i.e. a function taking a Node as an input, and returning TRUE or FALSE. If the pruneFun returns FALSE for a Node, then the Node and its entire sub-tree will not be considered.
filterFun	a function taking a Node as an argument. See Traverse for details.
format	if FALSE (the default), then no formatting will be applied. If TRUE, then the first formatter (if any) along the ancestor path is used for formatting.

inheritFromAncestors	if FALSE, and if the attribute is a field or a method, then only a Node itself is searched for the field/method. If TRUE, and if the Node does not contain the attribute, then ancestors are also searched.
direction	when converting to a network, should the edges point from root to children ("climb") or from child to parent ("descend")?
type	when converting type columns, the type is the discriminator, i.e. an attribute (e.g. field name) of each node
prefix	when converting type columns, the prefix used for the column names. Can be NULL to omit prefixes.

Value

ToDataFrameTree: a data.frame, where each row represents a Node in the tree or sub-tree spanned by x, possibly pruned according to pruneFun.

ToDataFrameTable: a data.frame, where each row represents a leaf Node in the tree or sub-tree spanned by x, possibly pruned according to pruneFun.

ToDataFrameNetwork: a data.frame, where each row represents a Node in the tree or sub-tree spanned by x, possibly pruned according to pruneFun. The first column is called 'from', while the second is called 'to', describing the parent to child edge (for direction "climb") or the child to parent edge (for direction "descend"). If [AreNamesUnique](#) is TRUE, then the Network is based on the Node\$name, otherwise on the Node\$pathString

ToDataFrameTypeCol: a data.frame in table format (i.e. where each row represents a leaf in the tree or sub-tree spanned by x), possibly pruned according to pruneFun. In addition to ..., each distinct type is output to a column.

Examples

```
data(acme)
acme$attributesAll
as.data.frame(acme, row.names = NULL, optional = FALSE, "cost", "p")

ToDataFrameTree(acme, "cost", "p")
ToDataFrameNetwork(acme, "cost", "p", direction = "climb")
ToDataFrameTable(acme, "cost", "p")
ToDataFrameTypeCol(acme)

#use the pruneFun:
acme$Do(function(x) x$totalCost <- Aggregate(x, "cost", sum), traversal = "post-order")
ToDataFrameTree(acme, "totalCost", pruneFun = function(x) x$totalCost > 300000)

#inherit
acme$Set(floor = c(1, 2, 3), filterFun = function(x) x$level == 2)
as.data.frame(acme, row.names = NULL, optional = FALSE, "floor", inheritFromAncestors = FALSE)
as.data.frame(acme, row.names = NULL, optional = FALSE, "floor", inheritFromAncestors = TRUE)

#using a function as an attribute:
acme$Accounting$Head <- "Mrs. Numright"
acme$Research$Head <- "Mr. Stein"
```

```

acme$IT$Head <- "Mr. Squarehead"
ToDataFrameTable(acme, department = function(x) x$parent$name, "name", "Head", "cost")

#complex TypeCol
acme$IT$Outsource$AddChild("India")
acme$IT$Outsource$AddChild("Poland")
acme$Set(type = c('company', 'department', 'project', 'project', 'department',
                  'project', 'project', 'department', 'program', 'project',
                  'project', 'project', 'project'
                  )
          )
print(acme, 'type')
ToDataFrameTypeCol(acme, type = 'type')

```

as.dendrogram.Node	<i>Convert a Node to a dendrogram</i>
--------------------	---------------------------------------

Description

Convert a data.tree structure to a [dendrogram](#)

Usage

```

## S3 method for class 'Node'
as.dendrogram(
  object,
  heightAttribute = DefaultPlotHeight,
  edgetext = FALSE,
  ...
)

```

Arguments

object	The Node to convert
heightAttribute	The attribute (field name or function) storing the height
edgetext	If TRUE, then the for non-leaf nodes the node name is stored as the dendrogram's edge text.
...	Additional parameters

Value

An object of class dendrogram

See Also

Other Conversions from Node: [ToNewick\(\)](#)

Examples

```

data(acme)
acmed <- as.dendrogram(acme)
plot(acmed, center = TRUE)

#you can take an attribute for the height:
acme$Do( function(x) x$myPlotHeight <- (10 - x$level))
acmed <- as.dendrogram(acme, heightAttribute = "myPlotHeight")
plot(acmed, center = TRUE)

#or directly a function
acmed <- as.dendrogram(acme, heightAttribute = function(x) 10 - x$level)
plot(acmed)

```

as.igraph.Node	<i>Convert a data.tree structure to an igraph network</i>
----------------	---

Description

This requires the igraph package to be installed. Also, this requires the names of the Nodes to be unique within the data.tree structure.

Usage

```

as.igraph.Node(
  x,
  vertexAttributes = character(),
  edgeAttributes = character(),
  directed = FALSE,
  direction = c("climb", "descend"),
  ...
)

```

Arguments

x	The root Node to convert
vertexAttributes	A vector of strings, representing the attributes in the data.tree structure to add as attributes to the vertices of the igraph
edgeAttributes	A vector of strings, representing the attributes in the data.tree structure to add as edge attributes of the igraph
directed	Logical scalar, whether or not to create a directed graph.
direction	when converting to a network, should the edges point from root to children ("climb") or from child to parent ("descend")?
...	Currently unused.

Value

an igraph object

See Also

AreNamesUnique

Examples

```
data(acme)
library(igraph)
ig <- as.igraph(acme, "p", c("level", "isLeaf"))
plot(ig)
```

as.list.Node

Convert a data.tree structure to a list-of-list structure

Description

Convert a data.tree structure to a list-of-list structure

Usage

```
## S3 method for class 'Node'
as.list(
  x,
  mode = c("simple", "explicit"),
  unname = FALSE,
  nameName = ifelse(unname, "name", ""),
  childrenName = "children",
  rootName = "",
  keepOnly = NULL,
  pruneFun = NULL,
  ...
)

ToListSimple(x, nameName = "name", pruneFun = NULL, ...)

ToListExplicit(
  x,
  unname = FALSE,
  nameName = ifelse(unname, "name", ""),
  childrenName = "children",
  pruneFun = NULL,
  ...
)
```

Arguments

x	The Node to convert
mode	How the list is structured. "simple" (the default) will add children directly as nested lists. "explicit" puts children in a separate nested list called childrenName
unname	If TRUE, and if mode is "explicit", then the nested children list will not have named arguments. This can be useful e.g. in the context of conversion to JSON, if you prefer the children to be an array rather than named objects.
nameName	The name that should be given to the name element
childrenName	The name that should be given to the children nested list
rootName	The name of the node. If provided, this overrides Node\$name
keepOnly	A character vector of attributes to include in the result. If NULL (the default), all attributes are kept.
pruneFun	allows providing a prune criteria, i.e. a function taking a Node as an input, and returning TRUE or FALSE. If the pruneFun returns FALSE for a Node, then the Node and its entire sub-tree will not be considered.
...	Additional parameters passed to as.list.Node

Examples

```
data(acme)

str(ToListSimple(acme))
str(ToListSimple(acme, keepOnly = "cost"))

str(ToListExplicit(acme))
str(ToListExplicit(acme, unname = TRUE))
str(ToListExplicit(acme, unname = TRUE, nameName = "id", childrenName = "descendants"))
```

as.Node

Convert an object to a data.tree data structure

Description

Convert an object to a data.tree data structure

Usage

```
as.Node(x, ...)
```

Arguments

x	The object to be converted
...	Additional arguments

See Also

Other as.Node: [as.Node.data.frame\(\)](#), [as.Node.dendrogram\(\)](#), [as.Node.list\(\)](#), [as.Node.phylo\(\)](#), [as.Node.rpart\(\)](#)

as.Node.BinaryTree	<i>Convert a a SplitNode from the party package to a data.tree structure.</i>
--------------------	---

Description

Convert a a SplitNode from the party package to a data.tree structure.

Usage

```
## S3 method for class 'BinaryTree'
as.Node(x, ...)
```

Arguments

x	The BinaryTree
...	additional arguments (unused)

Examples

```
library(party)
airq <- subset(airquality, !is.na(Ozone))
airct <- ctree(Ozone ~ ., data = airq,
               controls = ctree_control(maxsurrogate = 3))

tree <- as.Node(airct)
tree

print(tree,
      "label",
      criterion = function(x) round(x$criterion$maxcriterion, 3),
      statistic = function(x) round(max(x$criterion$statistic), 3)
)

FindNode(tree, 6)$path
```

as.Node.data.frame *Convert a data.frame to a data.tree structure*

Description

Convert a data.frame to a data.tree structure

Usage

```
## S3 method for class 'data.frame'
as.Node(
  x,
  ...,
  mode = c("table", "network"),
  pathName = "pathString",
  pathDelimiter = "/",
  colLevels = NULL,
  na.rm = TRUE
)

FromDataFrameTable(
  table,
  pathName = "pathString",
  pathDelimiter = "/",
  colLevels = NULL,
  na.rm = TRUE,
  check = c("check", "no-warn", "no-check"),
  suffix = "_attr"
)

FromDataFrameNetwork(network, check = c("check", "no-warn", "no-check"))
```

Arguments

x	The data.frame in the required format.
...	Any other argument implementations of this might need
mode	Either "table" (if x is a data.frame in tree or table format) or "network"
pathName	The name of the column in x containing the path of the row
pathDelimiter	The delimiter used to separate nodes in pathName
colLevels	Nested list of column names, determining on what node levels the attributes are written to.
na.rm	If TRUE, then NA's are treated as NULL and values will not be set on nodes
table	a data.frame in table or tree format, i.e. having a row for each leaf (and optionally for additional nodes). There should be a column called pathName, separated by pathDelimiter, describing the path of each row.

check	Either • "check": if the name conformance should be checked and warnings should be printed in case of non-conformance (the default) • "no-warn": if the name conformance should be checked, but no warnings should be printed in case of non-conformance (if you expect non-conformance) • "no-check" or FALSE: if the name conformance should not be checked; use this if performance is critical. However, in case of non-conformance, expect cryptic follow-up errors
suffix	optional suffix added to the column name in case the column name is the same as a path element. Defaults to '_attr'
network	A data.frame in network format, i.e. it must adhere to the following requirements: • It must contain as many rows as there are nodes (excluding the root, there is no row for the root) • Its first and second columns contain the network relationships. This can be either climbing (from parent to children) or descending (from child to parent) • Its subsequent columns contain the attributes to be set on the nodes • It must contain a single root • There are no cycles in the network

Value

The root Node of the data.tree structure

See Also

[as.data.frame.Node](#)

Other as.Node: [as.Node.dendrogram\(\)](#), [as.Node.list\(\)](#), [as.Node.phylo\(\)](#), [as.Node.rpart\(\)](#), [as.Node\(\)](#)

Examples

```
data(acme)

#Tree
x <- ToDataFrameTree(acme, "pathString", "p", "cost")
x
xN <- as.Node(x)
print(xN, "p", "cost")

#Table
x <- ToDataFrameTable(acme, "pathString", "p", "cost")
x
xN <- FromDataFrameTable(x)
print(xN, "p", "cost")
```

```

#More complex Table structure, using colLevels
acme$Set(floor = c(1, 2, 3), filterFun = function(x) x$level == 2)
x <- ToDataFrameTable(acme, "pathString", "floor", "p", "cost")
x
xN <- FromDataFrameTable(x, colLevels = list(NULL, "floor", c("p", "cost")), na.rm = TRUE)
print(xN, "floor", "p", "cost")

#Network
x <- ToDataFrameNetwork(acme, "p", "cost", direction = "climb")
x
xN <- FromDataFrameNetwork(x)
print(xN, "p", "cost")

```

as.Node.dendrogram	Convert a dendrogram to a <i>data.tree</i> Node
--------------------	---

Description

Convert a [dendrogram](#) to a *data.tree* Node

Usage

```

## S3 method for class 'dendrogram'
as.Node(
  x,
  name = "Root",
  heightName = "plotHeight",
  check = c("check", "no-warn", "no-check"),
  ...
)

```

Arguments

x	The dendrogram
name	The name of the root Node
heightName	The name under which the dendrogram's height is stored
check	Either "check": if the name conformance should be checked and warnings should be printed in case of non-conformance (the default) "no-warn": if the name conformance should be checked, but no warnings should be printed in case of non-conformance (if you expect non-conformance) "no-check" or FALSE: if the name conformance should not be checked; use this if performance is critical. However, in case of non-conformance, expect cryptic follow-up errors
...	Additional parameters

Value

The root Node of a data.tree

See Also

Other as.Node: [as.Node.data.frame\(\)](#), [as.Node.list\(\)](#), [as.Node.phylo\(\)](#), [as.Node.rpart\(\)](#), [as.Node\(\)](#)

Examples

```
hc <- hclust(dist(USArrests), "ave")
dend1 <- as.dendrogram(hc)
tree1 <- as.Node(dend1)
tree1$attributesAll
tree1$totalCount
tree1$leafCount
tree1$height
```

as.Node.list

Convert a nested list structure to a data.tree structure

Description

Convert a nested list structure to a data.tree structure

Usage

```
## S3 method for class 'list'
as.Node(
  x,
  mode = c("simple", "explicit"),
  nameName = "name",
  childrenName = "children",
  nodeName = NULL,
  interpretNullAsList = FALSE,
  check = c("check", "no-warn", "no-check"),
  ...
)

FromListExplicit(
  explicitList,
  nameName = "name",
  childrenName = "children",
  nodeName = NULL,
  check = c("check", "no-warn", "no-check")
)
```

```

FromListSimple(
  simpleList,
  nameName = "name",
  nodeName = NULL,
  interpretNullAsList = FALSE,
  check = c("check", "no-warn", "no-check")
)

```

Arguments

x	The list to be converted.
mode	How the list is structured. "simple" (the default) will interpret any list to be a child. "explicit" assumes that children are in a nested list called childrenName
nameName	The name of the element in the list that should be used as the name, can be NULL if mode = explicit and the children lists are named, or if an automatic name (running number) should be assigned
childrenName	The name of the element that contains the child list (applies to mode 'explicit' only).
nodeName	A name suggestion for x, if the name cannot be deferred otherwise. This is for example the case for the root with mode explicit and named lists.
interpretNullAsList	If TRUE, then NULL-valued lists are interpreted as child nodes. Else, they are interpreted as attributes. This has only an effect if mode is "simple".
check	Either • "check": if the name conformance should be checked and warnings should be printed in case of non-conformance (the default) • "no-warn": if the name conformance should be checked, but no warnings should be printed in case of non-conformance (if you expect non-conformance) • "no-check" or FALSE: if the name conformance should not be checked; use this if performance is critical. However, in case of non-conformance, expect cryptic follow-up errors
...	Any other argument to be passed to generic sub implementations
explicitList	A list in which children are in a separate nested list called childrenName.
simplelist	A list in which children are stored as nested list alongside other attributes. Any list is interpreted as a child Node

See Also

Other as.Node: [as.Node.data.frame\(\)](#), [as.Node.dendrogram\(\)](#), [as.Node.phylo\(\)](#), [as.Node.rpart\(\)](#), [as.Node\(\)](#)

Examples

```

kingJosephs <- list(name = "Joseph I",
  spouse = "Mary",

```

```

      born = "1818-02-23",
      died = "1839-08-29",
      children = list(
        list(name = "Joseph II",
              spouse = "Kathryn",
              born = "1839-03-28",
              died = "1865-12-19"),
        list(name = "Helen",
              born = "1840-17-08",
              died = "1845-01-01")
      )
    )
  )
  FromListExplicit(kingJosephs)

kingJosephs <- list(head = "Joseph I",
                    spouse = "Mary",
                    born = "1818-02-23",
                    died = "1839-08-29",
                    list(head = "Joseph II",
                          spouse = "Kathryn",
                          born = "1839-03-28",
                          died = "1865-12-19"),
                    list(head = "Helen",
                          born = "1840-17-08",
                          died = "1845-01-01")
                    )
  )
  FromListSimple(kingJosephs, nameName = "head")

kingJosephs <- list(spouse = "Mary",
                    born = "1818-02-23",
                    died = "1839-08-29",
                    `Joseph II` = list(spouse = "Kathryn",
                                         born = "1839-03-28",
                                         died = "1865-12-19"),
                    Helen = list(born = "1840-17-08",
                                  died = "1845-01-01")
                    )
  )
  FromListSimple(kingJosephs, nodeName = "Joseph I")

```

as.Node.party

Convert a a party from the partykit package to a data.tree structure.

Description

Convert a a party from the partykit package to a data.tree structure.

Usage

```
## S3 method for class 'party'
as.Node(x, ...)
```

Arguments

```
x                The party object
...              other arguments (unused)
```

Examples

```
library(partykit)
data("WeatherPlay", package = "partykit")
### splits ###
# split in overcast, humidity, and windy
sp_o <- partysplit(1L, index = 1:3)
sp_h <- partysplit(3L, breaks = 75)
sp_w <- partysplit(4L, index = 1:2)

## query labels
character_split(sp_o)

### nodes ###
## set up partynode structure
pn <- partynode(1L, split = sp_o, kids = list(
  partynode(2L, split = sp_h, kids = list(
    partynode(3L, info = "yes"),
    partynode(4L, info = "no"))),
  partynode(5L, info = "yes"),
  partynode(6L, split = sp_w, kids = list(
    partynode(7L, info = "yes"),
    partynode(8L, info = "no")))))
pn
### tree ###
## party: associate recursive partynode structure with data
py <- party(pn, WeatherPlay)
tree <- as.Node(py)

print(tree,
  "splitname",
  count = function(node) nrow(node$data),
  "splitLevel")

SetNodeStyle(tree,
  label = function(node) paste0(node$name, ": ", node$splitname),
  tooltip = function(node) paste0(nrow(node$data), " observations"),
  fontname = "helvetica")

SetEdgeStyle(tree,
  arrowhead = "none",
  label = function(node) node$splitLevel,
  fontname = "helvetica",
```

```

        penwidth = function(node) 12 * nrow(node$data)/nrow(node$root$data),
        color = function(node) {
            paste0("grey",
                  100 - as.integer( 100 * nrow(node$data)/nrow(node$root$data))
            )
        }
    )
Do(tree$leaves,
  function(node) {
    SetNodeStyle(node,
      shape = "box",
      color = ifelse(node$splitname == "yes", "darkolivegreen4", "lightsalmon4"),
      fillcolor = ifelse(node$splitname == "yes", "darkolivegreen1", "lightsalmon"),
      style = "filled,rounded",
      penwidth = 2
    )
  }
)

plot(tree)

```

as.Node.phylo

*Convert a phylo object from the ape package to a Node***Description**

Convert a phylo object from the ape package to a Node

Usage

```

## S3 method for class 'phylo'
as.Node(
  x,
  heightName = "plotHeight",
  replaceUnderscores = TRUE,
  namesNotUnique = FALSE,
  ...
)

```

Arguments

x	The phylo object to be converted
heightName	If the phylo contains edge lengths, then they will be converted to a height and stored in a field named according to this parameter (the default is "height")
replaceUnderscores	if TRUE (the default), then underscores in names are replaced with spaces

namesNotUnique if TRUE, then the name of the Nodes will be prefixed with a unique id. This is useful if the children of a parent have non-unique names.

... any other parameter to be passed to sub-implementations

See Also

Other ape phylo conversions: [GetPhyloNr\(\)](#), [as.phylo.Node\(\)](#)

Other as.Node: [as.Node.data.frame\(\)](#), [as.Node.dendrogram\(\)](#), [as.Node.list\(\)](#), [as.Node.rpart\(\)](#), [as.Node\(\)](#)

Examples

```
#which bird familes have the max height?
library(ape)
data(bird.families)
bf <- as.Node(bird.families)
height <- bf$height
t <- Traverse(bf, filterFun = function(x) x$level == 25)
Get(t, "name")
```

as.Node.rpart	<i>Convert an rpart object to a data.tree structure</i>
---------------	---

Description

Convert an [rpart](#) object to a data.tree structure

Usage

```
## S3 method for class 'rpart'
as.Node(x, digits = getOption("digits") - 3, use.n = FALSE, ...)
```

Arguments

x the rpart object to be converted

digits the number of digits to be used for numeric values in labels

use.n logical. Add cases to labels, see [text.rpart](#) for further information

... any other argument to be passed to generic sub implementations

Value

a data.tree object. The tree contains a field `rpart.id` which references back to the original node id in the row names of the rpart object.

See Also

Other as.Node: [as.Node.data.frame\(\)](#), [as.Node.dendrogram\(\)](#), [as.Node.list\(\)](#), [as.Node.phylo\(\)](#), [as.Node\(\)](#)

Examples

```
if (require(rpart)) {
  fit <- rpart(Kyphosis ~ Age + Number + Start, data = kyphosis)
  as.Node(fit)
}
```

as.phylo.Node

Convert a Node to a phylo object from the ape package.

Description

This method requires the ape package to be installed and loaded.

Usage

```
as.phylo.Node(x, heightAttribute = DefaultPlotHeight, ...)
```

Arguments

x	The root Node of the tree or sub-tree to be converted
heightAttribute	The attribute (field name or function) storing the height
...	any other argument

See Also

Other ape phylo conversions: [GetPhyloNr\(\)](#), [as.Node.phylo\(\)](#)

Examples

```
library(ape)
data(acme)
acmephylo <- as.phylo(acme)
#plot(acmephylo)
```

averageBranchingFactor

Calculate the average number of branches each non-leaf has

Description

Calculate the average number of branches each non-leaf has

Usage

```
averageBranchingFactor(node)
```

Arguments

node	The node to calculate the average branching factor for
------	--

CheckNameReservedWord	<i>Checks whether name is a reserved word, as defined in NODE_RESERVED_NAMES_CONST.</i>
-----------------------	---

Description

Checks whether name is a reserved word, as defined in NODE_RESERVED_NAMES_CONST.

Usage

```
CheckNameReservedWord(name, check = c("check", "no-warn", "no-check"))
```

Arguments

name	the name to check
check	Either • "check": if the name conformance should be checked and warnings should be printed in case of non-conformance (the default) • "no-warn": if the name conformance should be checked, but no warnings should be printed in case of non-conformance (if you expect non-conformance) • "no-check" or FALSE: if the name conformance should not be checked; use this if performance is critical. However, in case of non-conformance, expect cryptic follow-up errors

Climb

*Climb a tree from parent to children, by provided criteria.***Description**

This method lets you climb the tree, from crutch to crutch. On each Node, the Climb finds the first child having attribute value equal to the the provided argument.

Usage

```
#node$Climb(...)
Climb(node, ...)
```

Arguments

node	The root Node of the tree or subtree to climb
...	an attribute-value pairlist to be searched. For brevity, you can also provide a character vector to search for names.

Value

the Node having path ..., or NULL if such a path does not exist

See Also

[Node](#)
[Navigate](#)

Examples

```
data(acme)

#the following are all equivalent
Climb(acme, 'IT', 'Outsource')
Climb(acme, name = 'IT', name = 'Outsource')
Climb(acme, 'IT')$Climb('Outsource')
Navigate(acme, path = "IT/Outsource")

Climb(acme, name = 'IT')

Climb(acme, position = c(2, 1))
#or, equivalent:
Climb(acme, position = 2, position = 1)
Climb(acme, name = "IT", cost = 250000)

tree <- CreateRegularTree(5, 2)
tree$Climb(c("1", "1"), position = c(2, 2))$path
```

Clone	<i>Clone a tree (creates a deep copy)</i>
-------	---

Description

The method also clones object attributes (such as the formatters), if desired. If the method is called on a non-root, then the parent relationship is not cloned, and the resulting [Node](#) will be a root.

Usage

```
Clone(node, pruneFun = NULL, attributes = FALSE)
```

Arguments

node	the root node of the tree or sub-tree to clone
pruneFun	allows providing a prune criteria, i.e. a function taking a Node as an input, and returning TRUE or FALSE. If the pruneFun returns FALSE for a Node, then the Node and its entire sub-tree will not be considered.
attributes	if FALSE, then R class attributes (e.g. formatters and grViz styles) are not cloned. This makes the method faster.

Value

the clone of the tree or sub-tree

See Also

[SetFormat](#)

Examples

```
data(acme)
acmeClone <- Clone(acme)
acmeClone$name <- "New Acme"
# acmeClone does not point to the same reference object anymore:
acme$name

#cloning a subtree
data(acme)
itClone <- Clone(acme$IT)
itClone$isRoot
```

CreateRandomTree	Create a tree for demo and testing
------------------	------------------------------------

Description

Create a tree for demo and testing

Usage

CreateRandomTree(nodes = 100, root = Node\$new("1"), id = 1)

Arguments

- | | |
|-------|--|
| nodes | The number of nodes to create |
| root | the previous node (for recursion, typically use default value) |
| id | The id (for recursion) |

CreateRegularTree	Create a tree for demo and testing
-------------------	------------------------------------

Description

Create a tree for demo and testing

Usage

CreateRegularTree(height = 5, branchingFactor = 3, parent = Node\$new("1"))

Arguments

- | | |
|-----------------|---------------------------------|
| height | the number of levels |
| branchingFactor | the number of children per node |
| parent | the parent node (for recursion) |

Cumulate	<i>Cumulate values among siblings</i>
----------	---------------------------------------

Description

For example, you can sum up values of siblings before this Node.

Usage

```
Cumulate(node, attribute, aggFun, ...)
```

Arguments

node	The node on which we want to cumulate
attribute	determines what is collected. The attribute can be • a.) the name of a field or a property/active of each Node in the tree, e.g. <code>acme\$Get("p")</code> or <code>acme\$Get("position")</code> • b.) the name of a method of each Node in the tree, e.g. <code>acme\$Get("levelZeroBased")</code>, where e.g. <code>acme\$levelZeroBased <- function() acme\$level - 1</code> • c.) a function, whose first argument must be a Node e.g. <code>acme\$Get(function(node) node\$cost * node\$p)</code>
aggFun	the aggregation function to be applied to the children's attributes
...	any arguments to be passed on to attribute (in case it's a function)

Examples

```
data(acme)
acme$Do(function(x) x$cost <- Aggregate(x, "cost", sum), traversal = "post-order")
acme$Do(function(x) x$cumCost <- Cumulate(x, "cost", sum))
print(acme, "cost", "cumCost")
```

DefaultPlotHeight	<i>Calculates the height of a Node given the height of the root.</i>
-------------------	--

Description

This function puts leafs at the bottom (not hanging), and makes edges equally long. Useful for easy plotting with third-party packages, e.g. if you have no specific height attribute, e.g. with [as.dendrogram.Node](#), [ToNewick](#), and [as.phylo.Node](#)

Usage

```
DefaultPlotHeight(node, rootHeight = 100)
```

Arguments

node	The node
rootHeight	The height of the root

Examples

```
data(acme)
dacme <- as.dendrogram(acme, heightAttribute = function(x) DefaultPlotHeight(x, 200))
plot(dacme, center = TRUE)
```

Distance	<i>Find the distance between two nodes of the same tree</i>
----------	---

Description

The distance is measured as the number of edges that need to be traversed to reach node2 when starting from node1.

Usage

```
Distance(node1, node2)
```

Arguments

node1	the first node in the tree
node2	the second node in the same tree

Examples

```
data(acme)
Distance(FindNode(acme, "Outsource"), FindNode(acme, "Research"))
```

Do	<i>Executes a function on a set of nodes</i>
----	--

Description

Executes a function on a set of nodes

Usage

```
# 00-style:
# node$Do(fun,
#         ...,
#         traversal = c("pre-order", "post-order", "in-order", "level", "ancestor"),
#         pruneFun = NULL,
#         filterFun = NULL)

# traditional:
Do(nodes, fun, ...)
```

Arguments

nodes	The nodes on which to perform the Get (typically obtained via Traverse)
fun	the function to execute. The function is expected to be either a Method, or to take a Node as its first argument
...	any additional parameters to be passed on to fun

See Also

[Node](#)
[Get](#)
[Set](#)
[Traverse](#)

Examples

```
data(acme)
traversal <- Traverse(acme)
Do(traversal, function(node) node$expectedCost <- node$p * node$cost)
print(acme, "expectedCost")
```

FindNode

*Find a node by name in the (sub-)tree***Description**

Scans the entire sub-tree spanned by node and returns the first [Node](#) having the name specified. This is mainly useful for trees whose name is unique. If [AreNamesUnique](#) is FALSE, i.e. if there is more than one Node called name in the tree, then it is undefined which one will be returned. Also note that this method is not particularly fast. See examples for a faster way to index large trees, if you need to do multiple searches. See [Traverse](#) if you need to find multiple Nodes.

Usage

```
FindNode(node, name)
```

Arguments

node	The root Node of the tree or sub-tree to search
name	The name of the Node to be returned

Value

The first Node whose name matches, or NULL if no such Node is found.

See Also

AreNamesUnique, Traverse

Examples

```
data(acme)
FindNode(acme, "Outsource")

#re-usable hashed index for multiple searches:
if(!AreNamesUnique(acme)) stop("Hashed index works for unique names only!")
trav <- Traverse(acme, "level")
names(trav) <- Get(trav, "name")
nameIndex <- as.environment(trav)
#you could also use hash from package hash instead!
#nameIndex <- hash(trav)
nameIndex$Outsource
nameIndex$IT
```

FormatFixedDecimal *Format a Number as a Decimal*

Description

Simple function that can be used as a format function when converting trees to a data.frame

Usage

```
FormatFixedDecimal(x, digits = 3)
```

Arguments

x	a numeric scalar or vector
digits	the number of digits to print after the decimal point

Value

A string corresponding to x, suitable for printing

Examples

```
data(acme)
print(acme, prob = acme$Get("p", format = function(x) FormatFixedDecimal(x, 4)))
```

FormatPercent	<i>Format a Number as a Percentage</i>
---------------	--

Description

This utility method can be used as a format function when converting trees to a `data.frame`

Usage

```
FormatPercent(x, digits = 2, format = "f", ...)
```

Arguments

<code>x</code>	A number
<code>digits</code>	The number of digits to print
<code>format</code>	The format to use
<code>...</code>	Any other argument passed to <code>formatC</code>

Value

A string corresponding to `x`, suitable for printing

See Also

`formatC`

Examples

```
data(acme)
print(acme, prob = acme$Get("p", format = FormatPercent))
```

Get

*Traverse a Tree and Collect Values***Description**

The `Get` method is one of the most important ones of the `data.tree` package. It lets you traverse a tree and collect values along the way. Alternatively, you can call a method or a function on each [Node](#).

Usage

```
# OO-style:
#node$Get(attribute,
#           ...,
#           traversal = c("pre-order", "post-order", "in-order", "level", "ancestor"),
#           pruneFun = NULL,
#           filterFun = NULL,
#           format = FALSE,
#           inheritFromAncestors = FALSE)

# traditional:
Get(nodes,
     attribute,
     ...,
     format = FALSE,
     inheritFromAncestors = FALSE,
     simplify = c(TRUE, FALSE, "array", "regular"))
```

Arguments

<code>nodes</code>	The nodes on which to perform the <code>Get</code> (typically obtained via Traverse)
<code>attribute</code>	determines what is collected. The attribute can be - a.) the name of a field or a property/active of each <code>Node</code> in the tree, e.g. <code>acme\$Get("p")</code> or <code>acme\$Get("position")</code> - b.) the name of a method of each <code>Node</code> in the tree, e.g. <code>acme\$Get("levelZeroBased")</code>, where e.g. <code>acme\$levelZeroBased <- function() acme\$level - 1</code> - c.) a function, whose first argument must be a <code>Node</code> e.g. <code>acme\$Get(function(node) node\$cost * node\$p)</code>
<code>...</code>	in case the attribute is a function or a method, the ellipsis is passed to it as additional arguments.
<code>format</code>	if <code>FALSE</code> (the default), no formatting is being used. If <code>TRUE</code> , then the first formatter (if any) found along the ancestor path is being used for formatting (see SetFormat). If <code>format</code> is a function, then the collected value is passed to that function, and the result is returned.

`inheritFromAncestors` if TRUE, then the path above a Node is searched to get the attribute in case it is NULL.

`simplify` same as [sapply](#), i.e. TRUE, FALSE or "array". Additionally, you can specify "regular" if each returned value is of length > 1, and equally named. See below for an example.

Value

a vector containing the attributes collected during traversal, in traversal order. NULL is converted to NA, such that `length(Node$Get) == Node$totalCount`

See Also

[Node](#)

[Set](#)

[Do](#)

[Traverse](#)

Examples

```
data(acme)
acme$Get("level")
acme$Get("totalCount")
```

```
acme$Get(function(node) node$cost * node$p,
          filterFun = isLeaf)
```

```
#This is equivalent:
nodes <- Traverse(acme, filterFun = isLeaf)
Get(nodes, function(node) node$cost * node$p)
```

```
#simplify = "regular" will preserve names
acme$Get(function(x) c(position = x$position, level = x$level), simplify = "regular")
```

GetAttribute

Get an attribute from a Node.

Description

Get an attribute from a Node.

Usage

```
GetAttribute(
  node,
  attribute,
  ...,
  format = FALSE,
  inheritFromAncestors = FALSE,
  nullAsNa = TRUE
)
```

Arguments

node	The Node from which the attribute should be fetched.
attribute	determines what is collected. The attribute can be - a.) the name of a field or a property/active of each Node in the tree, e.g. <code>acme\$Get("p")</code> or <code>acme\$Get("position")</code> - b.) the name of a method of each Node in the tree, e.g. <code>acme\$Get("levelZeroBased")</code>, where e.g. <code>acme\$levelZeroBased <- function() acme\$level - 1</code> - c.) a function, whose first argument must be a Node e.g. <code>acme\$Get(function(node) node\$cost * node\$p)</code>
...	in case the attribute is a function or a method, the ellipsis is passed to it as additional arguments.
format	if FALSE (the default), no formatting is being used. If TRUE, then the first formatter (if any) found along the ancestor path is being used for formatting (see SetFormat). If format is a function, then the collected value is passed to that function, and the result is returned.
inheritFromAncestors	if TRUE, then the path above a Node is searched to get the attribute in case it is NULL.
nullAsNa	If TRUE (the default), then NULL is returned as NA. Otherwise it is returned as NULL.

Examples

```
data(acme)
GetAttribute(acme$IT$Outsource, "cost")
```

GetPhyloNr

Determine the number a Node has after conversion to a phylo object

Description

Use this function when plotting a Node as a phylo, e.g. to set custom labels to plot.

Usage

```
GetPhyloNr(x, type = c("node", "edge"))
```

Arguments

x	The Node
type	Either "node" (the default) or "edge" (to get the number of the edge from x to its parent)

Value

an integer representing the node

See Also

Other ape phylo conversions: [as.Node.phylo\(\)](#), [as.phylo.Node\(\)](#)

Examples

```
library(ape)
library(data.tree)
data(acme)
ap <- as.phylo(acme)
#plot(ap)
#nodeLabels("IT Dep.", GetPhyloNr(Climb(acme, "IT")))
#edgeLabels("Good!", GetPhyloNr(Climb(acme, "IT", "Switch to R"), "edge"))
```

isLeaf	<i>Check if a Node is a leaf</i>
--------	----------------------------------

Description

Check if a Node is a leaf

Usage

```
isLeaf(node)
```

Arguments

node	The Node to test.
------	-------------------

Value

TRUE if the Node is a leaf, FALSE otherwise

isNotLeaf	<i>Check if a Node is not a leaf</i>
-----------	--------------------------------------

Description

Check if a Node is not a leaf

Usage

isNotLeaf(node)

Arguments

node The Node to test.

Value

FALSE if the Node is a leaf, TRUE otherwise

isNotRoot	<i>Check if a Node is not a root</i>
-----------	--------------------------------------

Description

Check if a Node is not a root

Usage

isNotRoot(node)

Arguments

node The Node to test.

Value

FALSE if the Node is the root, TRUE otherwise

isRoot	<i>Check if a Node is the root</i>
--------	------------------------------------

Description

Check if a Node is the root

Usage

```
isRoot(node)
```

Arguments

node	The Node to test.
------	-------------------

Value

TRUE if the Node is the root, FALSE otherwise

mushroom	<i>Sample Data: Data Used by the ID3 Vignette</i>
----------	---

Description

mushroom contains attributes of mushrooms. We can use this data to predict a mushroom's toxicity based on its attributes. The attributes available in the data set are:

Usage

```
data(mushroom)
```

Format

```
data.frame
```

Details

- color the color of a mushroom
- size whether a mushroom is small or large
- points whether a mushroom has points
- edibility whether a mushroom is edible or toxic

 Navigate

Navigate to another node by relative path.

Description

Navigate to another node by relative path.

Usage

```
Navigate(node, path)
```

Arguments

node	The starting Node to navigate
path	A string or a character vector describing the path to navigate

Details

The path is always relative to the node. Navigation to the parent is defined by `..`, whereas navigation to a child is defined via the child's name. If path is provided as a string, then the navigation steps are separated by `'/'`.

See Also

[Climb](#)

Examples

```
data(acme)
Navigate(acme$Research, "../IT/Outsource")
Navigate(acme$Research, c("../", "IT", "Outsource"))
```

 Node

Create a data.tree Structure With Nodes

Description

Node is at the very heart of the `data.tree` package. All trees are constructed by tying together Node objects.

Usage

```
# n1 <- Node$new("Node 1")
```

Format

An `R6Class` generator object

Details

Assemble `Node` objects into a `data.tree` structure and use the traversal methods to set, get, and perform operations on it. Typically, you construct larger tree structures by converting from `data.frame`, `list`, or other formats.

Most methods (e.g. `node$Sort()`) also have a functional form (e.g. `Sort(node)`)

Active bindings

`name` Gets or sets the name of a `Node`. For example `Node$name <- "Acme"`.

`printFormatters` gets or sets the formatters used to print a `Node`. Set this as a list to a root node.

The different formatters are `h` (horizontal), `v` (vertical), `l` (L), `j` (junction), and `s` (separator). For example, you can set the formatters to `list(h = "\u2500", v = "\u2502", l = "\u2514", j = "\u251C", s = " ")` to get a similar behavior as in `fs::dir_tree()`. The defaults are: `list(h = "--", v = "\u00A6", l = "\u00B0", j = "\u00A6", s = " ")`

`parent` Gets or sets the parent `Node` of a `Node`. Only set this if you know what you are doing, as you might mess up the tree structure!

`children` Gets or sets the children list of a `Node`. Only set this if you know what you are doing, as you might mess up the tree structure!

`isLeaf` Returns `TRUE` if the `Node` is a leaf, `FALSE` otherwise

`isRoot` Returns `TRUE` if the `Node` is the root, `FALSE` otherwise

`count` Returns the number of children of a `Node`

`totalCount` Returns the total number of `Nodes` in the tree

`path` Returns a vector of mode character containing the names of the `Nodes` in the path from the root to this `Node`

`pathString` Returns a string representing the path to this `Node`, separated by backslash

`position` The position of a `Node` within its siblings

`attributes` The attributes defined on this specific node

`attributesAll` The distinct union of attributes defined on all the nodes in the tree spanned by this `Node`

`levelName` Returns the name of the `Node`, preceded by level times `'*'`. Useful for printing and not typically called by package users.

`leaves` Returns a list containing all the leaf `Nodes`

`leafCount` Returns the number of leaves are below a `Node`

`level` Returns an integer representing the level of a `Node`. For example, the root has level 1.

`height` Returns `max(level)` of any of the `Nodes` of the tree

`isBinary` Returns `TRUE` if all `Nodes` in the tree (except the leaves) have `count = 2`

`root` Returns the root of a `Node` in a tree.

`siblings` Returns a list containing all the siblings of this `Node`

`averageBranchingFactor` Returns the average number of crotches below this `Node`

Methods

Public methods:

- `Node$new()`
- `Node$AddChild()`
- `Node$AddChildNode()`
- `Node$AddSibling()`
- `Node$AddSiblingNode()`
- `Node$RemoveChild()`
- `Node$RemoveAttribute()`
- `Node$Sort()`
- `Node$Revert()`
- `Node$Prune()`
- `Node$Climb()`
- `Node$Navigate()`
- `Node$Get()`
- `Node$Do()`
- `Node$Set()`
- `Node$clone()`

Method `new()`: Create a new Node object. This is often used to create the root of a tree when creating a tree programmatically.

Usage:

```
Node$new(name, check = c("check", "no-warn", "no-check"), ...)
```

Arguments:

`name` the name of the node to be created

`check` Either

- "check": if the name conformance should be checked and warnings should be printed in case of non-conformance (the default)
- "no-warn": if the name conformance should be checked, but no warnings should be printed in case of non-conformance (if you expect non-conformance)
- "no-check" or FALSE: if the name conformance should not be checked; use this if performance is critical. However, in case of non-conformance, expect cryptic follow-up errors

... A name-value mapping of node attributes

Returns: A new 'Node' object

Examples:

```
node <- Node$new("mynode", x = 2, y = "value of y")
node$y
```

Method `AddChild()`: Creates a Node and adds it as the last sibling as a child to the Node on which this is called.

Usage:

```
Node$AddChild(name, check = c("check", "no-warn", "no-check"), ...)
```

Arguments:

name the name of the node to be created

check Either

- "check": if the name conformance should be checked and warnings should be printed in case of non-conformance (the default)
- "no-warn": if the name conformance should be checked, but no warnings should be printed in case of non-conformance (if you expect non-conformance)
- "no-check" or FALSE: if the name conformance should not be checked; use this if performance is critical. However, in case of non-conformance, expect cryptic follow-up errors

... A name-value mapping of node attributes

Returns: The new Node (invisibly)

Examples:

```
root <- Node$new("myroot", myname = "I'm the root")
root$AddChild("child1", myname = "I'm the favorite child")
child2 <- root$AddChild("child2", myname = "I'm just another child")
child3 <- child2$AddChild("child3", myname = "Grandson of a root!")
print(root, "myname")
```

Method AddChildNode(): Adds a Node as a child to this node.

Usage:

```
Node$AddChildNode(child)
```

Arguments:

child The child "Node" to add.

Returns: the child node added (this lets you chain calls)

Examples:

```
root <- Node$new("myroot")
child <- Node$new("mychild")
root$AddChildNode(child)
```

Method AddSibling(): Creates a new Node called name and adds it after this Node as a sibling.

Usage:

```
Node$AddSibling(name, check = c("check", "no-warn", "no-check"), ...)
```

Arguments:

name the name of the node to be created

check Either

- "check": if the name conformance should be checked and warnings should be printed in case of non-conformance (the default)
- "no-warn": if the name conformance should be checked, but no warnings should be printed in case of non-conformance (if you expect non-conformance)

- "no-check" or FALSE: if the name conformance should not be checked; use this if performance is critical. However, in case of non-conformance, expect cryptic follow-up errors

... A name-value mapping of node attributes

Returns: the sibling node (this lets you chain calls)

Examples:

```
#' root <- Node$new("myroot")
child <- root$AddChild("child1")
sibling <- child$AddSibling("sibling1")
```

Method AddSiblingNode(): Adds a Node after this Node, as a sibling.

Usage:

```
Node$AddSiblingNode(sibling)
```

Arguments:

sibling The "Node" to add as a sibling.

Returns: the added sibling node (this lets you chain calls, as in the examples)

Examples:

```
root <- Node$new("myroot")
child <- Node$new("mychild")
sibling <- Node$new("sibling")
root$AddChildNode(child)$AddSiblingNode(sibling)
```

Method RemoveChild(): Remove the child Node called name from a Node and returns it.

Usage:

```
Node$RemoveChild(name)
```

Arguments:

name the name of the node to be created

Returns: the subtree spanned by the removed child.

Examples:

```
node <- Node$new("myroot")$AddChild("mychild")$root
node$RemoveChild("mychild")
```

Method RemoveAttribute(): Removes attribute called name from this Node.

Usage:

```
Node$RemoveAttribute(name, stopIfNotAvailable = TRUE)
```

Arguments:

name the name of the node to be created

stopIfNotAvailable Gives an error if stopIfNotAvailable and the attribute does not exist.

Examples:

```
node <- Node$new("mynode")
node$RemoveAttribute("age", stopIfNotAvailable = FALSE)
node$age <- 27
node$RemoveAttribute("age")
node
```

Method Sort(): Sort children of a Node or an entire data.tree structure

Usage:

```
Node$Sort(attribute, ..., decreasing = FALSE, recursive = TRUE)
```

Arguments:

attribute determines what is collected. The attribute can be

- a.) the name of a **field** or a **property/active** of each Node in the tree, e.g. acme\$Get("p") or acme\$Get("position")
- b.) the name of a **method** of each Node in the tree, e.g. acme\$Get("levelZeroBased"), where e.g. acme\$levelZeroBased <- function() acme\$level - 1
- c.) a **function**, whose first argument must be a Node e.g. acme\$Get(function(node) node\$cost * node\$p)

... any parameters to be passed on the the attribute (in case it's a method or a function)

decreasing sort order

recursive if TRUE, the method will be called recursively on the Node's children. This allows sorting an entire tree.

Details: You can sort with respect to any argument of the tree. But note that sorting has side-effects, meaning that you modify the underlying, original data.tree object structure.

See also [Sort](#) for the equivalent function.

Returns: Returns the node on which Sort is called, invisibly. This can be useful to chain Node methods.

Examples:

```
data(acme)
acme$Do(function(x) x$totalCost <- Aggregate(x, "cost", sum), traversal = "post-order")
Sort(acme, "totalCost", decreasing = FALSE)
print(acme, "totalCost")
```

Method Revert(): Reverts the sort order of a Node's children.

See also [Revert](#) for the equivalent function.

Usage:

```
Node$Revert(recursive = TRUE)
```

Arguments:

recursive if TRUE, the method will be called recursively on the Node's children. This allows sorting an entire tree.

Returns: returns the Node invisibly (for chaining)

Method Prune(): Prunes a tree.

Pruning refers to removing entire subtrees. This function has side-effects, it modifies your data.tree structure!

See also [Prune](#) for the equivalent function.

Usage:

```
Node$Prune(pruneFun)
```

Arguments:

pruneFun allows providing a a prune criteria, i.e. a function taking a Node as an input, and returning TRUE or FALSE. If the pruneFun returns FALSE for a Node, then the Node and its entire sub-tree will not be considered.

Returns: the number of nodes removed

Examples:

```
data(acme)
acme$Do(function(x) x$cost <- Aggregate(x, "cost", sum))
Prune(acme, function(x) x$cost > 700000)
print(acme, "cost")
```

Method Climb(): Climb a tree from parent to children, by provided criteria.

Usage:

```
Node$Climb(...)
```

Arguments:

... an attribute-value pairlist to be searched. For brevity, you can also provide a character vector to search for names.

node The root [Node](#) of the tree or subtree to climb

Details: This method lets you climb the tree, from crutch to crutch. On each Node, the Climb finds the first child having attribute value equal to the the provided argument.

See also [Climb](#) and [Navigate](#)

```
Climb(node, ...)
```

Returns: the Node having path ..., or NULL if such a path does not exist

Examples:

```
data(acme)

#the following are all equivalent
Climb(acme, 'IT', 'Outsource')
Climb(acme, name = 'IT', name = 'Outsource')
Climb(acme, 'IT')$Climb('Outsource')
Navigate(acme, path = "IT/Outsource")

Climb(acme, name = 'IT')

Climb(acme, position = c(2, 1))
#or, equivalent:
Climb(acme, position = 2, position = 1)
```

```
Climb(acme, name = "IT", cost = 250000)

tree <- CreateRegularTree(5, 2)
tree$Climb(c("1", "1"), position = c(2, 2))$path
```

Method `Navigate()`: Navigate to another node by relative path.

Usage:

```
Node$Navigate(path)
```

Arguments:

path A string or a character vector describing the path to navigate

node The starting [Node](#) to navigate

Details: The path is always relative to the Node. Navigation to the parent is defined by `..`, whereas navigation to a child is defined via the child's name. If path is provided as a string, then the navigation steps are separated by `'/'`.

See also [Navigate](#) and [Climb](#)

Examples:

```
data(acme)
Navigate(acme$Research, "../IT/Outsource")
Navigate(acme$Research, c("../", "IT", "Outsource"))
```

Method `Get()`: Traverse a Tree and Collect Values

Usage:

```
Node$Get(
  attribute,
  ...,
  traversal = c("pre-order", "post-order", "in-order", "level", "ancestor"),
  pruneFun = NULL,
  filterFun = NULL,
  format = FALSE,
  inheritFromAncestors = FALSE,
  simplify = c(TRUE, FALSE, "array", "regular")
)
```

Arguments:

attribute determines what is collected. The attribute can be

- a.) the name of a **field** or a **property/active** of each Node in the tree, e.g. `acme$Get("p")` or `acme$Get("position")`
- b.) the name of a **method** of each Node in the tree, e.g. `acme$Get("levelZeroBased")`, where e.g. `acme$levelZeroBased <- function() acme$level - 1`
- c.) a **function**, whose first argument must be a Node e.g. `acme$Get(function(node) node$cost * node$p)`

... in case the attribute is a function or a method, the ellipsis is passed to it as additional arguments.

traversal defines the traversal order to be used. This can be

pre-order Go to first child, then to its first child, etc.

post-order Go to the first branch's leaf, then to its siblings, and work your way back to the root

in-order Go to the first branch's leaf, then to its parent, and only then to the leaf's sibling

level Collect root, then level 2, then level 3, etc.

ancestor Take a node, then the node's parent, then that node's parent in turn, etc. This ignores the pruneFun

function You can also provide a function, whose sole parameter is a [Node](#) object. The function is expected to return the node's next node, a list of the node's next nodes, or NULL.

Read the `data.tree` vignette for a detailed explanation of these traversal orders.

`pruneFun` allows providing a prune criteria, i.e. a function taking a `Node` as an input, and returning TRUE or FALSE. If the `pruneFun` returns FALSE for a `Node`, then the `Node` and its entire sub-tree will not be considered.

`filterFun` allows providing a filter, i.e. a function taking a `Node` as an input, and returning TRUE or FALSE. Note that if filter returns FALSE, then the node will be excluded from the result (but not the entire subtree).

`format` if FALSE (the default), no formatting is being used. If TRUE, then the first formatter (if any) found along the ancestor path is being used for formatting (see [SetFormat](#)). If `format` is a function, then the collected value is passed to that function, and the result is returned.

`inheritFromAncestors` if TRUE, then the path above a `Node` is searched to get the attribute in case it is NULL.

`simplify` same as [sapply](#), i.e. TRUE, FALSE or "array". Additionally, you can specify "regular" if each returned value is of length > 1, and equally named. See below for an example.

Details: The `Get` method is one of the most important ones of the `data.tree` package. It lets you traverse a tree and collect values along the way. Alternatively, you can call a method or a function on each [Node](#).

See also [Get](#), [Node](#), [Set](#), [Do](#), [Traverse](#)

Returns: a vector containing the attributes collected during traversal, in traversal order. NULL is converted to NA, such that `length(Node$Get) == Node$totalCount`

Examples:

```
data(acme)
acme$Get("level")
acme$Get("totalCount")
```

```
acme$Get(function(node) node$cost * node$p,
          filterFun = isLeaf)
```

```
#This is equivalent:
nodes <- Traverse(acme, filterFun = isLeaf)
Get(nodes, function(node) node$cost * node$p)
```

```
#simplify = "regular" will preserve names
acme$Get(function(x) c(position = x$position, level = x$level), simplify = "regular")
```

Method Do(): Executes a function on a set of nodes

Usage:

```
Node$Do(
  fun,
  ...,
  traversal = c("pre-order", "post-order", "in-order", "level", "ancestor"),
  pruneFun = NULL,
  filterFun = NULL
)
```

Arguments:

fun the function to execute. The function is expected to be either a Method, or to take a Node as its first argument

... A name-value mapping of node attributes

traversal defines the traversal order to be used. This can be

pre-order Go to first child, then to its first child, etc.

post-order Go to the first branch's leaf, then to its siblings, and work your way back to the root

in-order Go to the first branch's leaf, then to its parent, and only then to the leaf's sibling

level Collect root, then level 2, then level 3, etc.

ancestor Take a node, then the node's parent, then that node's parent in turn, etc. This ignores the pruneFun

function You can also provide a function, whose sole parameter is a [Node](#) object. The function is expected to return the node's next node, a list of the node's next nodes, or NULL.

Read the `data.tree` vignette for a detailed explanation of these traversal orders.

pruneFun allows providing a prune criteria, i.e. a function taking a Node as an input, and returning TRUE or FALSE. If the pruneFun returns FALSE for a Node, then the Node and its entire sub-tree will not be considered.

filterFun allows providing a filter, i.e. a function taking a Node as an input, and returning TRUE or FALSE. Note that if filter returns FALSE, then the node will be excluded from the result (but not the entire subtree).

Details: See also [Node](#), [Get](#), [Set](#), [Traverse](#)

Examples:

```
data(acme)
acme$Do(function(node) node$expectedCost <- node$p * node$cost)
print(acme, "expectedCost")
```

Method Set(): Traverse a Tree and Assign Values

Usage:

```
Node$Set(
  ...,
  traversal = c("pre-order", "post-order", "in-order", "level", "ancestor"),
  pruneFun = NULL,
  filterFun = NULL
)
```

Arguments:

... each argument can be a vector of values to be assigned. Recycled.

`traversal` defines the traversal order to be used. This can be

pre-order Go to first child, then to its first child, etc.

post-order Go to the first branch's leaf, then to its siblings, and work your way back to the root

in-order Go to the first branch's leaf, then to its parent, and only then to the leaf's sibling

level Collect root, then level 2, then level 3, etc.

ancestor Take a node, then the node's parent, then that node's parent in turn, etc. This ignores the `pruneFun`

function You can also provide a function, whose sole parameter is a [Node](#) object. The function is expected to return the node's next node, a list of the node's next nodes, or `NULL`.

Read the `data.tree` vignette for a detailed explanation of these traversal orders.

`pruneFun` allows providing a prune criteria, i.e. a function taking a `Node` as an input, and returning `TRUE` or `FALSE`. If the `pruneFun` returns `FALSE` for a `Node`, then the `Node` and its entire sub-tree will not be considered.

`filterFun` allows providing a filter, i.e. a function taking a `Node` as an input, and returning `TRUE` or `FALSE`. Note that if filter returns `FALSE`, then the node will be excluded from the result (but not the entire subtree).

Details: The method takes one or more vectors as an argument. It traverses the tree, whereby the values are picked from the vector. Also available as OO-style method on [Node](#).

See also [Node](#), [Get](#), [Do](#), [Traverse](#)

Returns: invisibly returns the nodes (useful for chaining)

Examples:

```
data(acme)
acme$Set(departmentId = 1:acme$totalCount, openingHours = NULL, traversal = "post-order")
acme$Set(head = c("Jack Brown",
                  "Mona Moneyhead",
                  "Dr. Frank N. Stein",
                  "Eric Nerdahl"
                ),
        filterFun = function(x) !x$isLeaf
      )
print(acme, "departmentId", "head")
```

Method `clone()`: The objects of this class are cloneable with this method.

Usage:

```
Node$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

See Also

For more details see the [data.tree](#) documentations, or the `data.tree` vignette: `vignette("data.tree")`

[Node](#)

[Sort](#)

Examples

```
library(data.tree)
acme <- Node$new("Acme Inc.")
accounting <- acme$AddChild("Accounting")$
  AddSibling("Research")$
  AddChild("New Labs")$
  parent$
  AddSibling("IT")$
  AddChild("Outsource")
print(acme)
```

```
## -----
## Method `Node$new`
## -----
```

```
node <- Node$new("mynode", x = 2, y = "value of y")
node$y
```

```
## -----
## Method `Node$AddChild`
## -----
```

```
root <- Node$new("myroot", myname = "I'm the root")
root$AddChild("child1", myname = "I'm the favorite child")
child2 <- root$AddChild("child2", myname = "I'm just another child")
child3 <- child2$AddChild("child3", myname = "Grandson of a root!")
print(root, "myname")
```

```
## -----
## Method `Node$AddChildNode`
## -----
```

```
root <- Node$new("myroot")
child <- Node$new("mychild")
root$AddChildNode(child)
```

```

## -----
## Method `Node$AddSibling`
## -----

#' root <- Node$new("myroot")
child <- root$AddChild("child1")
sibling <- child$AddSibling("sibling1")

## -----
## Method `Node$AddSiblingNode`
## -----

root <- Node$new("myroot")
child <- Node$new("mychild")
sibling <- Node$new("sibling")
root$AddChildNode(child)$AddSiblingNode(sibling)

## -----
## Method `Node$RemoveChild`
## -----

node <- Node$new("myroot")$AddChild("mychild")$root
node$RemoveChild("mychild")

## -----
## Method `Node$RemoveAttribute`
## -----

node <- Node$new("mynode")
node$RemoveAttribute("age", stopIfNotAvailable = FALSE)
node$age <- 27
node$RemoveAttribute("age")
node

## -----
## Method `Node$Sort`
## -----

data(acme)
acme$Do(function(x) x$totalCost <- Aggregate(x, "cost", sum), traversal = "post-order")
Sort(acme, "totalCost", decreasing = FALSE)
print(acme, "totalCost")

## -----
## Method `Node$Prune`
## -----

```

```

data(acme)
acme$Do(function(x) x$cost <- Aggregate(x, "cost", sum))
Prune(acme, function(x) x$cost > 700000)
print(acme, "cost")

## -----
## Method `Node$Climb`
## -----

data(acme)

#the following are all equivalent
Climb(acme, 'IT', 'Outsource')
Climb(acme, name = 'IT', name = 'Outsource')
Climb(acme, 'IT')$Climb('Outsource')
Navigate(acme, path = "IT/Outsource")

Climb(acme, name = 'IT')

Climb(acme, position = c(2, 1))
#or, equivalent:
Climb(acme, position = 2, position = 1)
Climb(acme, name = "IT", cost = 250000)

tree <- CreateRegularTree(5, 2)
tree$Climb(c("1", "1"), position = c(2, 2))$path

## -----
## Method `Node$Navigate`
## -----

data(acme)
Navigate(acme$Research, "../IT/Outsource")
Navigate(acme$Research, c("../", "IT", "Outsource"))

## -----
## Method `Node$Get`
## -----

data(acme)
acme$Get("level")
acme$Get("totalCount")

acme$Get(function(node) node$cost * node$p,
          filterFun = isLeaf)

#This is equivalent:
nodes <- Traverse(acme, filterFun = isLeaf)

```

```

Get(nodes, function(node) node$cost * node$p)

#simplify = "regular" will preserve names
acme$Get(function(x) c(position = x$position, level = x$level), simplify = "regular")

## -----
## Method `Node$Do`
## -----

data(acme)
acme$Do(function(node) node$expectedCost <- node$p * node$cost)
print(acme, "expectedCost")

## -----
## Method `Node$Set`
## -----

data(acme)
acme$Set(departmentId = 1:acme$totalCount, openingHours = NULL, traversal = "post-order")
acme$Set(head = c("Jack Brown",
                  "Mona Moneyhead",
                  "Dr. Frank N. Stein",
                  "Eric Nerdahl"
                ),
        filterFun = function(x) !x$isLeaf
      )
print(acme, "departmentId", "head")

```

NODE_RESERVED_NAMES_CONST

Names that are reserved by the Node class.

Description

These are reserved by the Node class, you cannot use these as attribute names. Note also that all attributes starting with a . are reserved.

Usage

```
NODE_RESERVED_NAMES_CONST
```

Format

An object of class character of length 41.

plot.Node	<i>Plot a graph, or get a graphviz dot representation of the tree</i>
-----------	---

Description

Use these methods to style your graph, and to plot it. The functionality is built around the DiagrammeR package, so for anything that goes beyond simple plotting, it is recommended to read its documentation at <https://rich-iannone.github.io/DiagrammeR/docs.html>. Note that DiagrammeR is only suggested by data.tree, so 'plot' only works if you have installed it on your system.

Usage

```
## S3 method for class 'Node'
plot(
  x,
  ...,
  direction = c("climb", "descend"),
  pruneFun = NULL,
  output = "graph"
)
```

```
ToDiagrammeRGraph(root, direction = c("climb", "descend"), pruneFun = NULL)
```

```
SetNodeStyle(node, inherit = TRUE, keepExisting = FALSE, ...)
```

```
SetEdgeStyle(node, inherit = TRUE, keepExisting = FALSE, ...)
```

```
SetGraphStyle(root, keepExisting = FALSE, ...)
```

```
GetDefaultTooltip(node)
```

Arguments

x	The root node of the data.tree structure to plot
...	For the SetStyle methods, this can be any stlyeName / value pair. See https://graphviz.org/Documentation for details. For the plot.Node generic method, this is not used.
direction	when converting to a network, should the edges point from root to children ("climb") or from child to parent ("descend")?
pruneFun	allows providing a prune criteria, i.e. a function taking a Node as an input, and returning TRUE or FALSE. If the pruneFun returns FALSE for a Node, then the Node and its entire sub-tree will not be considered.
output	A string specifying the output type; graph (the default) renders the graph using the <code>grViz()</code> function and visNetwork renders the graph using the <code>visnetwork()</code> function.
root	The root <code>Node</code> of the data.tree structure to visualize.

node	The Node of the data.tree structure on which you would like to set style attributes.
inherit	If TRUE, then children will inherit this node's style. Otherwise they inherit from this node's parent. Note that the inherit always applies to the node, i.e. all style attributes of a node and not to a single style attribute.
keepExisting	If TRUE, then style attributes are added to possibly existing style attributes on the node.

Details

Use `SetNodeStyle` and `SetEdgeStyle` to define the style of your plot. Use `plot` to display a graphical representation of your tree.

The most common styles that can be set on the nodes are:

- color
- fillcolor
- fixedsize true or false
- fontcolor
- fontname
- fontsize
- height
- penwidth
- shape box, ellipse, polygon, circle, box, etc.
- style
- tooltip
- width

The most common styles that can be set on the edges are:

- arrowhead e.g. normal, dot, vee
- arrowsize
- arrowtail
- color
- dir forward, back, both, none
- fontcolor
- fontname
- fontsize
- headport
- label
- minlen
- penwidth
- tailport

- tooltip

A good source to understand the attributes is <https://graphviz.org/Documentation.php>. Another good source is the DiagrammeR package documentation, or more specifically: <https://rich-iannone.github.io/DiagrammeR/do>

In addition to the standard GraphViz functionality, the `data.tree` plotting infrastructure takes advantage of the fact that `data.tree` structure are always hierarchic. Thus, style attributes are inherited from parents to children on an individual basis. For example, you can set the `fontcolor` to red on a parent, and then all children will also have red font, except if you specifically disallow inheritance. Labels and tooltips are never inherited.

Another feature concerns functions: Instead of setting a fixed value (e.g. `SetNodeStyle(acme, label = "Acme. Inc")`), you can set a function (e.g. `SetNodeStyle(acme, label = function(x) x$name)`). The function must take a `Node` as its single argument. Together with inheritance, this becomes a very powerful tool.

The `GetDefaultTooltip` method is a utility method that can be used to print all attributes of a `Node`.

There are some more examples in the 'applications' vignette, see `vignette('applications', package = "data.tree")`

Examples

```
data(acme)
SetGraphStyle(acme, rankdir = "TB")
SetEdgeStyle(acme, arrowhead = "vee", color = "blue", penwidth = 2)
#per default, Node style attributes will be inherited:
SetNodeStyle(acme, style = "filled,rounded", shape = "box", fillcolor = "GreenYellow",
             fontname = "helvetica", tooltip = GetDefaultTooltip)
SetNodeStyle(acme$IT, fillcolor = "LightBlue", penwidth = "5px")
#inheritance can be avoided:
SetNodeStyle(acme$Accounting, inherit = FALSE, fillcolor = "Thistle",
             fontcolor = "Firebrick", tooltip = "This is the accounting department")
SetEdgeStyle(acme$Research$`New Labs`,
             color = "red",
             label = "Focus!",
             penwidth = 3,
             fontcolor = "red")
#use Do to set style on specific nodes:
Do(acme$leaves, function(node) SetNodeStyle(node, shape = "egg"))
plot(acme)

#print p as label, where available:
SetNodeStyle(acme, label = function(node) node$p)
plot(acme)
```

Description

Print a Node in a human-readable fashion.

Usage

```
## S3 method for class 'Node'
print(
  x,
  ...,
  pruneMethod = c("simple", "dist", NULL),
  limit = 100,
  pruneFun = NULL,
  row.names = T
)
```

Arguments

<code>x</code>	The Node
<code>...</code>	Node attributes to be printed. Can be either a character (i.e. the name of a Node field), a Node method, or a function taking a Node as a single argument. See <code>Get</code> for details on the meaning of attribute.
<code>pruneMethod</code>	The method can be used to prune for printing in a simple way. If <code>NULL</code> , the entire tree is displayed. If "simple", then only the first <code>limit</code> nodes are displayed. If "dist", then Nodes are removed everywhere in the tree, according to their level. If <code>pruneFun</code> is provided, then <code>pruneMethod</code> is ignored.
<code>limit</code>	The maximum number of nodes to print. Can be <code>NULL</code> if the entire tree should be printed.
<code>pruneFun</code>	allows providing a prune criteria, i.e. a function taking a Node as an input, and returning <code>TRUE</code> or <code>FALSE</code> . If the <code>pruneFun</code> returns <code>FALSE</code> for a Node, then the Node and its entire sub-tree will not be considered.
<code>row.names</code>	If <code>TRUE</code> (default), then the row names are printed out. Else, they are not.

Examples

```
data(acme)
print(acme, "cost", "p")
print(acme, "cost", probability = "p")
print(acme, expectedCost = function(x) x$cost * x$p)
do.call(print, c(acme, acme$attributesAll))

tree <- CreateRegularTree(4, 5)
# print entire tree:
print(tree, pruneMethod = NULL)
# print first 20 nodes:
print(tree, pruneMethod = "simple", limit = 20)
# print 20 nodes, removing leafs first:
print(tree, pruneMethod = "dist", limit = 20)
# provide your own pruning function:
```

```
print(tree, pruneFun = function(node) node$position != 2)
```

Prune

Prunes a tree.

Description

Pruning refers to removing entire subtrees. This function has side-effects, it modifies your data.tree structure!

Usage

```
Prune(node, pruneFun)
```

Arguments

node	The root of the sub-tree to be pruned
pruneFun	allows providing a prune criteria, i.e. a function taking a Node as an input, and returning TRUE or FALSE. If the pruneFun returns FALSE for a Node, then the Node and its entire sub-tree will not be considered.

Value

the number of nodes removed

See Also

[Node](#)

Examples

```
data(acme)
acme$Do(function(x) x$cost <- Aggregate(x, "cost", sum))
Prune(acme, function(x) x$cost > 700000)
print(acme, "cost")
```

Revert	<i>Reverts the sort order of a Node's children.</i>
--------	---

Description

Reverts the sort order of a Node's children.

Usage

Revert(node, recursive = TRUE)

Arguments

- node the Node whose childrens' sort order is to be reverted
- recursive If TRUE, then revert is called recursively on all children.

Value

returns the Node invisibly (for chaining)

See Also

- [Node](#)
- [Sort](#)

Set	<i>Traverse a Tree and Assign Values</i>
-----	--

Description

The method takes one or more vectors as an argument. It traverses the tree, whereby the values are picked from the vector. Also available as OO-style method on [Node](#).

Usage

```
#OO-style:
# node$Set(...,
#   traversal = c("pre-order", "post-order", "in-order", "level", "ancestor"),
#   pruneFun = NULL,
#   filterFun = NULL)
#traditional:
Set(nodes, ...)
```

Arguments

nodes	The nodes on which to perform the Get (typically obtained via Traverse)
...	each argument can be a vector of values to be assigned. Recycled.

Value

invisibly returns the nodes (useful for chaining)

See Also

- [Node](#)
- [Get](#)
- [Do](#)
- [Traverse](#)

Examples

```
data(acme)
acme$Set(departmentId = 1:acme$totalCount, openingHours = NULL, traversal = "post-order")
acme$Set(head = c("Jack Brown",
                  "Mona Moneyhead",
                  "Dr. Frank N. Stein",
                  "Eric Nerdahl"
                ),
        filterFun = function(x) !x$isLeaf
      )
print(acme, "departmentId", "head")
```

SetFormat	<i>Set a formatter function on a specific node</i>
-----------	--

Description

Formatter functions set on a Node act as a default formatter when printing and using the [Get](#) method. The formatter is inherited, meaning that whenever Get fetches an attribute from a Node, it checks on the Node or on any of its ancestors whether a formatter is set.

Usage

```
SetFormat(node, name, formatFun)
```

Arguments

node	The node on which to set the formatter
name	The attribute name for which to set the formatter
formatFun	The formatter, i.e. a function taking a value as an input, and formatting returning the formatted value

See Also

Get
print.Node

Examples

```
data(acme)
acme$Set(id = 1:(acme$totalCount))
SetFormat(acme, "id", function(x) FormatPercent(x, digits = 0))
SetFormat(Climb(acme, "IT"), "id", FormatFixedDecimal)
print(acme, "id")
# Calling Get with an explicit formatter will overwrite the default set on the Node:
print(acme, id = acme$Get("id", format = function(x) paste0("id:", x)))

# Or, to avoid formatters, even though you set them on a Node:
print(acme, id = acme$Get("id", format = identity))
```

Sort	<i>Sort children of a Node or an entire data.tree structure</i>
------	---

Description

You can sort with respect to any argument of the tree. But note that sorting has side-effects, meaning that you modify the underlying, original data.tree object structure.

Usage

```
Sort(node, attribute, ..., decreasing = FALSE, recursive = TRUE)
```

Arguments

node	The node whose children are to be sorted
attribute	determines what is collected. The attribute can be • a.) the name of a field or a property/active of each Node in the tree, e.g. <code>acme\$Get("p")</code> or <code>acme\$Get("position")</code> • b.) the name of a method of each Node in the tree, e.g. <code>acme\$Get("levelZeroBased")</code>, where e.g. <code>acme\$levelZeroBased <- function() acme\$level - 1</code> • c.) a function, whose first argument must be a Node e.g. <code>acme\$Get(function(node) node\$cost * node\$p)</code>
...	any parameters to be passed on the the attribute (in case it's a method or a function)
decreasing	sort order
recursive	if TRUE, Sort will be called recursively on the Node's children. This allows sorting an entire tree.

Value

Returns the node on which Sort is called, invisibly. This can be useful to chain Node methods.

See Also

[Node](#)
[Revert](#)

Examples

```
data(acme)
acme$Do(function(x) x$totalCost <- Aggregate(x, "cost", sum), traversal = "post-order")
Sort(acme, "totalCost", decreasing = FALSE)
print(acme, "totalCost")
```

ToNewick

Write a data.tree structure to Newick notation

Description

To read from Newick, you can use the ape package, and convert the resulting phylo object to a data.tree structure.

Usage

```
ToNewick(node, heightAttribute = DefaultPlotHeight, ...)
```

Arguments

node	The root Node of a tree or sub-tree to be converted
heightAttribute	The attribute (field name, method, or function) storing or calculating the height for each Node
...	parameters that will be passed on the the heightAttributeName, in case it is a function

See Also

Other Conversions from Node: [as.dendrogram.Node\(\)](#)

Examples

```
data(acme)
ToNewick(acme)
ToNewick(acme, heightAttribute = NULL)
ToNewick(acme, heightAttribute = function(x) DefaultPlotHeight(x, 200))
ToNewick(acme, rootHeight = 200)
```

Traverse

*Traverse a tree or a sub-tree***Description**

Traverse takes the root of a tree or a sub-tree, and "walks" the tree in a specific order. It returns a list of [Node](#) objects, filtered and pruned by `filterFun` and `pruneFun`.

Usage

```
Traverse(
  node,
  traversal = c("pre-order", "post-order", "in-order", "level", "ancestor"),
  pruneFun = NULL,
  filterFun = NULL
)
```

Arguments

<code>node</code>	the root of a tree or a sub-tree that should be traversed
<code>traversal</code>	any of 'pre-order' (the default), 'post-order', 'in-order', 'level', 'ancestor', or a custom function (see details)
<code>pruneFun</code>	allows providing a prune criteria, i.e. a function taking a Node as an input, and returning TRUE or FALSE. If the <code>pruneFun</code> returns FALSE for a Node , then the Node and its entire sub-tree will not be considered.
<code>filterFun</code>	allows providing a filter, i.e. a function taking a Node as an input, and returning TRUE or FALSE. Note that if filter returns FALSE, then the node will be excluded from the result (but not the entire subtree).

Details

The traversal order is as follows. (Note that these descriptions are not precise and complete. They are meant for quick reference only. See the `data.tree` vignette for a more detailed description).

pre-order Go to first child, then to its first child, etc.

post-order Go to the first branch's leaf, then to its siblings, and work your way back to the root

in-order Go to the first branch's leaf, then to its parent, and only then to the leaf's sibling

level Collect root, then level 2, then level 3, etc.

ancestor Take a node, then the node's parent, then that node's parent in turn, etc. This ignores the `pruneFun`

function You can also provide a function, whose sole parameter is a [Node](#) object. The function is expected to return the node's next node, a list of the node's next nodes, or NULL.

Value

a list of Nodes

See Also

[Node](#)

[Get](#)

[Set](#)

[Do](#)

Index

* Conversions from Node

as.dendrogram.Node, 8
ToNewick, 61

* Newick

ToNewick, 61

* ape phylo conversions

as.Node.phylo, 20
as.phylo.Node, 22
GetPhyloNr, 34

* as.Node

as.Node, 11
as.Node.data.frame, 13
as.Node.dendrogram, 15
as.Node.list, 16
as.Node.phylo, 20
as.Node.rpart, 21

* datasets

acme, 3
mushroom, 37
NODE_RESERVED_NAMES_CONST, 52

acme, 3

Aggregate, 3

AreNamesUnique, 5, 7, 29

as.data.frame.Node, 5, 14

as.dendrogram.Node, 8, 27, 61

as.igraph.Node, 9

as.list.Node, 10

as.Node, 11, 14, 16, 17, 21, 22

as.Node.BinaryTree, 12

as.Node.data.frame, 12, 13, 16, 17, 21, 22

as.Node.dendrogram, 12, 14, 15, 17, 21, 22

as.Node.list, 12, 14, 16, 16, 21, 22

as.Node.party, 18

as.Node.phylo, 12, 14, 16, 17, 20, 22, 35

as.Node.rpart, 12, 14, 16, 17, 21, 21

as.phylo.Node, 21, 22, 27, 35

averageBranchingFactor, 23

CheckNameReservedWord, 23

Climb, 24, 38, 44, 45

Clone, 25

CreateRandomTree, 26

CreateRegularTree, 26

Cumulate, 27

data.tree, 49

DefaultPlotHeight, 27

dendrogram, 8, 15

Distance, 28

Do, 28, 33, 46, 48, 59, 63

FindNode, 29

FormatFixedDecimal, 30

FormatPercent, 31

FromDataFrameNetwork

(as.Node.data.frame), 13

FromDataFrameTable

(as.Node.data.frame), 13

FromListExplicit (as.Node.list), 16

FromListSimple (as.Node.list), 16

Get, 4, 6, 29, 32, 46–48, 59, 63

GetAttribute, 33

GetDefaultTooltip (plot.Node), 53

GetPhyloNr, 21, 22, 34

grViz(), 53

isLeaf, 35

isNotLeaf, 36

isNotRoot, 36

isRoot, 37

mushroom, 37

Navigate, 24, 38, 44, 45

Node, 4, 24, 25, 29, 32–34, 38, 38, 44–49,
53–55, 57–59, 61–63

NODE_RESERVED_NAMES_CONST, 52

plot.Node, 53

print.Node, [55](#)
Prune, [44](#), [57](#)

R6Class, [39](#)
Revert, [43](#), [58](#), [61](#)
rpart, [21](#)

sapply, [33](#), [46](#)
Set, [29](#), [33](#), [46](#), [47](#), [58](#), [63](#)
SetEdgeStyle (plot.Node), [53](#)
SetFormat, [32](#), [34](#), [46](#), [59](#)
SetGraphStyle (plot.Node), [53](#)
SetNodeStyle (plot.Node), [53](#)
Sort, [43](#), [49](#), [58](#), [60](#)

ToDataFrameNetwork
 (as.data.frame.Node), [5](#)
ToDataFrameTable (as.data.frame.Node), [5](#)
ToDataFrameTree (as.data.frame.Node), [5](#)
ToDataFrameTypeCol
 (as.data.frame.Node), [5](#)
ToDiagrammerRGraph (plot.Node), [53](#)
ToListExplicit (as.list.Node), [10](#)
ToListSimple (as.list.Node), [10](#)
ToNewick, [8](#), [27](#), [61](#)
Traverse, [6](#), [29](#), [32](#), [33](#), [46–48](#), [59](#), [62](#)

visnetwork(), [53](#)