

Package ‘nLTT’

January 13, 2020

Type Package

Title Calculate the NLTT Statistic

Version 1.4.3

Description Provides functions to calculate the normalised Lineage-Through-Time (nLTT) statistic, given two phylogenetic trees. The nLTT statistic measures the difference between two Lineage-Through-Time curves, where each curve is normalised both in time and in number of lineages.

License GPL-2

Imports ape, coda, deSolve, testit

Suggests DDD, ggplot2, Hmisc, knitr, microbenchmark, plyr, reshape2, rmarkdown, TESS, testthat, TreeSim

NeedsCompilation no

RoxygenNote 6.1.1

Encoding UTF-8

VignetteBuilder knitr

URL <https://github.com/thijsjanzen/nLTT>

BugReports <https://github.com/thijsjanzen/nLTT/issues>

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Package providing functions to visualize the normalized Lineage-Through-Time statistic, and calculate the difference between two nLTT curves

Description

This package provides a function to visualize the normalized Lineage-Through-Time (nLTT) statistic, where the number of lineages relative to the maximum number of lineages in a phylogenetic tree is plotted against the relative time between the most common recent ancestor and the present. Furthermore the package provides a function to calculate the difference between two nLTT curves, including two different distance measurements.

Updates: Version 1.4.3: Added support for log transformation before normalization.

Version 1.4: Added the following four functions: `get_branching_times`, `get_n_lineages`, `get_norm_brts` and `get_norm_n`. Furthermore, vignette building has improved, and the underlying code base has been polished up as well.

Version 1.3.1: Added walkthrough vignette, and updated several typos in the manual

Version 1.3: Version 1.3 adds a lot of extended functionality: firstly, we have added functions to calculate, and plot, the average nLTT across a number of phylogenies. Furthermore, we have added vignettes, and we have added a GitHub repository. On the GitHub repository the vignettes are separately accessible through the wiki. Lastly we have added an extra option to the nLTT functions, where the user can specify if the used trees are rooted, or not. Under the hood, some changes have been made as well, the majority of the code is now conforming to the lintR code conventions, and we have written formalized tests that check correctness of all code (code coverage 100

Version 1.2.1: updated comments and coding style to adhere to the general coding rules. Backwards compatibility has been favoured for the nLTT stat functions. ABC related functions are no longer

backwards compatible (variable names have been changed to adhere to coding style).

Version 1.2: added an "exact" nLTT function. This function is faster for small trees, and provides an exact measurement of the nLTT function. Comparison between "old" and "exact" estimates show that these are highly correlated, although the "exact" values are slightly higher than the "old" values. The "exact" function should generally be preferred, unless dealing with extremely large trees (500+ tips) in which case the old function is much faster.

Version 1.2: updated the example for the ABC_SMC_nLTT function, prior generating and prior density functions are now more realistic

Version 1.1.1: fixed a minor bug in the ABC_SMC_nLTT function

Version 1.1.1: removed some intermediate output in ABC_SMC_nLTT function

Version 1.1: Made a universal nLTT function called "nLTTstat", with argument "distanceMethod", this serves as a more elegant wrapper for the functions "normLTTdiffABS" and "normLTTdiffSQ"

Version 1.1: Updated references in the manual

Details

Package:	nLTT
Type:	Package
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Author(s)

Thijs Janzen

Maintainer: Thijs Janzen <thijsjanzen@gmail.com>

References

Janzen,T. Hoehna,S., Etienne,R.S. (2015) Approximate Bayesian Computation of diversification rates from molecular phylogenies: introducing a new efficient summary statistic, the nLTT. Methods in Ecology and Evolution. doi: 10.1111/2041-210X.12350

abc_smc_nltt

A function to perform Approximate Bayesian Computation within a Sequential Markov Chain (ABC-SMC), for diversification analysis of phylogenetic trees.

Description

This function performs ABC-SMC as described in Toni 2009 for given diversification model, provided a phylogenetic tree. ABC-SMC is not limited to only using the normalized LTT as statistic.

Usage

```
abc_smc_nltt(
  tree, statistics, simulation_function, init_epsilon_values,
  prior_generating_function, prior_density_function,
  number_of_particles = 1000, sigma = 0.05, stop_rate = 1e-05
)
```

Arguments

<code>tree</code>	an object of class "phylo"; the tree upon which we want to fit our diversification model
<code>statistics</code>	A vector containing functions that take a tree as an argument and return a single scalar value (the statistic).
<code>simulation_function</code>	A function that implements the diversification model and returns an object of class "phylo".
<code>init_epsilon_values</code>	A vector containing the initial threshold values for the summary statistics from the vector <code>statistics</code> .
<code>prior_generating_function</code>	Function to generate parameters from the prior distribution of these parameters (e.g. a function returning lambda and mu in case of the birth-death model)
<code>prior_density_function</code>	Function to calculate the prior probability of a set of parameters.
<code>number_of_particles</code>	Number of particles to be used per iteration of the ABC-SMC algorithm.
<code>sigma</code>	Standard deviation of the perturbation distribution (perturbation distribution is a gaussian with mean 0).
<code>stop_rate</code>	If the acceptance rate drops below <code>stopRate</code> , stop the ABC-SMC algorithm and assume convergence.

Value

A matrix with n columns, where n is the number of parameters you are trying to estimate.

Author(s)

Thijs Janzen

References

Toni, T., Welch, D., Strelkowa, N., Ipsen, A., & Stumpf, M.P.H. (2009). Approximate Bayesian computation scheme for parameter inference and model selection in dynamical systems. *Journal of the Royal Society Interface*, 6(31), 187-202.

Examples

```

## Not run:

prior_gen <- function() {
  return( rexp(n=2, rate=0.1) )
}

prior_dens <- function(val) {
  return( dexp( val[1], rate = 0.1) * dexp( val[2], rate = 0.1) )
}

require(TESS)

treeSim <- function(params) {
  t <- TESS.sim.age(n=1, lambda = params[1], mu = params[2], age = 10)[[1]]
  return(t)
}

obs <- treeSim(c(0.5,0.1))

statWrapper <- function(tree1) {
  return( nLTstat_exact(tree1, obs, "abs"))
}

stats <- c(statWrapper)

results <- abc.smc.nltt(
  obs, stats, treeSim, init_epsilon_values = 0.2,
  prior_generating_function = prior_gen,
  prior_density_function = prior_dens,
  number_of_particles = 1000, sigma = 0.05, stop_rate = 1e-5
)

## End(Not run) # end of dontrun

```

exampleTrees

example trees to test the functionality of the package

Description

100 phylogenetic trees of class `phylo`, generated using the `sim.globalBiDe.age` function from the `TESS` package, with $\lambda = 0.3$, $\mu = 0.1$, $age = 10$.

Usage

```
data(exampleTrees)
```

Format

A list containing objects of class phylo.

Examples

```
data(exampleTrees);
obs <- exampleTrees[[1]];
nltt_plot(obs);
```

get_average_nltt_matrix

Get the average nLTT from a collection of phylogenies

Description

Get the average nLTT from a collection of phylogenies

Usage

```
get_average_nltt_matrix(phylogenies, dt = 0.001)
```

Arguments

phylogenies	the phylogenies, supplied as either a list or a multiPhylo object, where the phylogenies are of type 'phylo'
dt	The timestep resolution, where 1/dt is the number of points evaluated

Value

A matrix of timepoints with the average number of (normalized) lineages through (normalized) time

Author(s)

Richel Bilderbeek

Examples

```
get_average_nltt_matrix(c(ape::rcoal(10), ape::rcoal(20)))
```

get_branching_times *Collect the branching times from the stem age*

Description

Collect the branching times from the stem age

Usage

```
get_branching_times(phylogeny)
```

Arguments

phylogeny a phylogeny of class 'phylo'

Value

branching times, in time units before the present

Author(s)

Richel Bilderbeek

Examples

```
phylogeny <- ape::read.tree(text = "((a:2,b:2):1,c:3);")  
phylogeny$root.edge <- 2 # nolint ape variable name  
testthat::expect_true(  
  all(nLTT::get_branching_times(phylogeny) == c(5, 3, 2)))
```

get_nltt_values *Get the nLTT values in time*

Description

Collect the nLTT values in time over all phylogenies in the long form.

Usage

```
get_nltt_values(phylogenies, dt)
```

Arguments

phylogenies the phylogenies, supplied as either a list or a multiPhylo object, where the phylogenies are of type 'phylo'
dt The timestep resolution, where 1/dt is the number of points evaluated

Value

A dataframe of timepoints with the nLTT value of each phylogeny in time

Author(s)

Richel Bilderbeek

See Also

Use [nlts_diff](#) to compare nLTT statistic between one focal tree and a set of one or more other trees

Examples

```
library(ape)
library(ggplot2)
library(nLTT)

# Create some random phylogenies
phylogeny1 <- rcoal(10)
phylogeny2 <- rcoal(20)
phylogeny3 <- rcoal(30)
phylogeny4 <- rcoal(40)
phylogeny5 <- rcoal(50)
phylogeny6 <- rcoal(60)
phylogeny7 <- rcoal(70)
phylogenies <- c(phylogeny1, phylogeny2, phylogeny3,
               phylogeny4, phylogeny5, phylogeny6, phylogeny7
)
)

# Obtain the nLTT values
dt <- 0.2
nltt_values <- get_nltt_values(phylogenies, dt = dt)

# Check properties of nltt_values
testit::assert(names(nltt_values) == c("id", "t", "nltt"))
nltt_values_per_phylogeny <- (1 + (1 / dt))
n_phylogenies <- length(phylogenies)
testit::assert(nrow(nltt_values)
               == nltt_values_per_phylogeny * n_phylogenies
)
)

# Plot the phylogenies, where the individual nLTT values are visible
qplot(t, nltt, data = nltt_values, geom = "point",
      ylim = c(0,1),
      main = "Average nLTT plot of phylogenies", color = id, size = I(0.1)
) + stat_summary(
  fun.data = "mean_cl_boot", color = "red", geom = "smooth"
)

# Plot the phylogenies, where the individual nLTT values are omitted
qplot(
  t, nltt, data = nltt_values, geom = "blank", ylim = c(0,1),
```

```

  main = "Average nLTT plot of phylogenies"
) + stat_summary(
  fun.data = "mean_cl_boot", color = "red", geom = "smooth"
)

```

get_norm_brts

*Collect the normalized branching times from the stem age***Description**

Collect the normalized branching times from the stem age

Usage

```
get_norm_brts(phylogeny)
```

Arguments

phylogeny a phylogeny of class 'phylo'

Value

branching times, in time units before the present

Author(s)

Richel Bilderbeek

Examples

```

phylogeny <- ape::read.tree(text = "((a:2,b:2):1,c:3);")
phylogeny$root.edge <- 2 # nolint ape variable name
testthat::expect_true(
  all(nLTT::get_branching_times(phylogeny) == c(5, 3, 2)))

```

get_norm_n

*Collect the normalized number of lineages from the stem age***Description**

Collect the normalized number of lineages from the stem age

Usage

```
get_norm_n(phylogeny)
```

Arguments

`phylogeny` a phylogeny of class 'phylo'

Value

branching times, in time units before the present

Author(s)

Richel Bilderbeek

Examples

```
phylogeny <- ape::read.tree(text = "((a:2,b:2):1,c:3);")
phylogeny$root.edge <- 2 # nolint ape variable name
testthat::expect_true(
  all(nLT::get_branching_times(phylogeny) == c(5, 3, 2)))
```

`get_n_lineages` *Collect the number of lineages from the stem age*

Description

Collect the number of lineages from the stem age

Usage

`get_n_lineages(phylogeny)`

Arguments

`phylogeny` a phylogeny of class 'phylo'

Value

number of lineages, will go from 1 to the number of tips, if there is a stem, will go from 2 to the number of tips if there is no stem

Author(s)

Richel Bilderbeek

Examples

```
phylogeny <- ape::read.tree(text = "((a:2,b:2):1,c:3);")
testthat::expect_true(
  all(nLT::get_n_lineages(phylogeny) == c(2, 3)))
phylogeny$root.edge <- 2 # nolint ape variable name
testthat::expect_true(
  all(nLT::get_n_lineages(phylogeny) == c(1, 2, 3)))
```

get_phylogeny_nltt_matrix

Extract the nLTT matrix from a phylogeny

Description

Extract the nLTT matrix from a phylogeny

Usage

```
get_phylogeny_nltt_matrix(phylogeny)
```

Arguments

phylogeny A phylogeny of type phylo

Value

a matrix

Author(s)

Richel Bilderbeek

mcmc_nltt

Code to perform Metropolis-Hastings MCMC for a diversification model, given a phylogenetic tree. This function was used in the MEE paper to calculate the likelihood reference estimates.

Description

This function performs Metropolis-Hastings MCMC, where the user provides a likelihood function and a phylogenetic tree.

Usage

```
mcmc_nltt(phy, likelihood_function, parameters, logtransforms,  
iterations, burnin = round(iterations / 3), thinning = 1, sigma=1)
```

Arguments

<code>phy</code>	an object of class "phylo"; the tree upon which we want to fit our diversification model
<code>likelihood_function</code>	Function that calculates the likelihood of our diversification model, given the tree. Function should me of the format function(parameters,phy).
<code>parameters</code>	Initial parameters to start the chain.
<code>logtransforms</code>	Whether to perform jumps on logtransformed parameters (TRUE) or not (FALSE)
<code>iterations</code>	Length of the chain
<code>burnin</code>	Length of the burnin, default is 30
<code>thinning</code>	Size of thinning, default = 1
<code>sigma</code>	Standard deviation of the jumping distribution, which is N(0, sigma).

Value

An MCMC object, as used by the package "coda".

Author(s)

Sebastian Hoehna & Thijs Janzen

Examples

```
## MCMC examples are typically very slow #####
## Not run:

require(TESS);

obs <- TESS.sim.age(n = 1, lambda = 0.5, mu = 0.1, age = 10)[[1]];

LL_BD <- function(params, phy) {
  lnl <- tess.likelihood(phy, lambda = params[1], mu = params[2],
    samplingProbability = 1, log = TRUE);
  prior1 <- dunif( params[1], 0, 100, log = TRUE)
  prior2 <- dunif( params[2], 0, 100, log = TRUE);
  return(lnl + prior1 + prior2);
}

require(coda);

mcmc_out <- mcmc_nltt(obs, LL_BD, c(0.5, 0.1), c(TRUE, TRUE),
  iterations = 1000, burnin = 100, thinning = 10, sigma = 1)
plot(mcmc_out);

## End(Not run)
```

nLTTstat	<i>Calculate the difference between two normalized Lineage-Through-Time curves, given two phylogenetic trees.</i>
----------	---

Description

This function takes two ultrametric phylogenetic trees, calculates the normalized Lineage-Through-Time statistic for both trees and then calculates the difference between the two statistics.

Usage

```
nLTTstat(tree1, tree2, distance_method = "abs", ignore_stem = TRUE, log_transform = FALSE)
```

Arguments

tree1	an object of class "phylo"
tree2	an object of class "phylo"
distance_method	Chosen measurement of distance between the two nLTT curves, options are (case sensitive): - "abs": use the absolute distance - "squ": use the squared distance;
ignore_stem	a boolean whether to ignore the stem length
log_transform	a boolean whether to log-transform the number of lineages before normalization

Value

The difference between the two nLTT statistics

Author(s)

Thijs Janzen

Examples

```
data(exampleTrees)
nltt_plot(exampleTrees[[1]])
nltt_lines(exampleTrees[[2]], lty=2)
nLTTstat(
  exampleTrees[[1]], exampleTrees[[2]],
  distance_method = "abs", ignore_stem = TRUE)
```

<code>nLTTstat_exact</code>	<i>Calculate the exact difference between two normalized Lineage-Through-Time curves, given two phylogenetic trees.</i>
-----------------------------	---

Description

This function takes two ultrametric phylogenetic trees, calculates the normalized Lineage-Through-Time statistic for both trees and then calculates the exact difference between the two statistics. Whereas the function `nLTTstat` uses an approximation to calculate the difference (which is faster for large trees), the function `nLTTstat_exact` calculates the exact difference, and should generally be preferred. Although the estimates are highly similar, `nLTTstat_exact` tends to return slightly higher values.

Usage

```
nLTTstat_exact(tree1, tree2, distance_method = "abs",
                ignore_stem = TRUE, log_transform = FALSE)
```

Arguments

tree1	an object of class "phylo"
tree2	an object of class "phylo"
distance_method	Chosen measurement of distance between the two nLTT curves, options are (case sensitive): - "abs": use the absolute distance. - "squ": use the squared distance
ignore_stem	a boolean whether to ignore the stem length
log_transform	a boolean whether to log-transform the number of lineages before normalization

Value

The exact difference between the two nLTT statistics

Author(s)

Thijs Janzen

Examples

```
data(exampleTrees)
nltt_plot(exampleTrees[[1]])
nltt_lines(exampleTrees[[2]], lty = 2)
nLTTstat_exact(
  exampleTrees[[1]],
  exampleTrees[[2]],
  distance_method = "abs",
```

```
    ignore_stem = TRUE
)
```

nlts_diff

Calculates the nLTT statistic between each phylogeny in a collection compared to a same focal/reference tree

Description

Calculates the nLTT statistic between each phylogeny in a collection compared to a same focal/reference tree

Usage

```
nlts_diff(tree, trees, distance_method = "abs", ignore_stem = TRUE,
          log_transform = FALSE)
```

Arguments

tree	One phylogenetic tree
trees	A collection of one or more phylogenetic trees
distance_method	(string) absolute, or squared distance?
ignore_stem	(logical) Should the phylogeny its stem be ignored?
log_transform	(logical) Should the number of lineages be log-transformed before normalization?

Value

the nLTT statistic values, as a numeric vector of the same length as `trees`

Author(s)

Richel J.C. Bilderbeek

See Also

use [nltt_diff](#) to compare two phylogenies

Examples

```
tree <- ape::rcoal(4)
trees <- c(ape::rcoal(4), ape::rcoal(4))
nlts <- nlts_diff(tree, trees)
testit::assert(all(nlts >= 0.0 && nlts <= 1.0))
```

nltts_plot*Get the average nLTT from a collection of phylogenies***Description**

Get the average nLTT from a collection of phylogenies

Usage

```
nltts_plot(phylogenies, dt = 0.001, plot_nltts = FALSE,
           xlab = "Normalized Time", ylab = "Normalized Lineages",
           replot = FALSE, ...)
```

Arguments

<code>phylogenies</code>	the phylogenies, where the phylogenies are of type 'phylo'
<code>dt</code>	The timestep resolution, where 1/dt is the number of points evaluated
<code>plot_nltts</code>	Also plot each nLLT line
<code>xlab</code>	Label on the x axis
<code>ylab</code>	Label on the y axis
<code>replot</code>	If false, start a clean plot. If true, plot the new data over the current
<code>...</code>	Plotting options

Value

Nothing

Author(s)

Richel Bilderbeek

Examples

```
nltts_plot(c(ape::rcoal(10), ape::rcoal(10)))
nltts_plot(c(ape::rcoal(10), ape::rcoal(20)), dt = 0.1)
```

nltt_diff

Calculates the exact difference between the lineage through time curves of tree1 & tree2 (normalized in time and for the number of lineages)

Description

Calculates the exact difference between the lineage through time curves of tree1 & tree2 (normalized in time and for the number of lineages)

Usage

```
nltt_diff(tree1, tree2, distance_method = "abs", ignore_stem = TRUE,  
log_transform = FALSE)
```

Arguments

tree1	(phylo) First phylogenetic tree
tree2	(phylo) Second phylogenetic tree
distance_method	(string) absolute, or squared distance?
ignore_stem	logical Should the phylogeny its stem be ignored?
log_transform	(logical) Should the number of lineages be log-transformed before normalization?

Value

(scalar) normalized Lineage-Through-Time difference between tree1 & tree2

Author(s)

Thijs Janzen

See Also

use [nltts_diff](#) to compare a collection of phylogenies to one focal/reference tree

<code>nltt_diff_exact</code>	<i>Calculates the exact, difference between the lineage through time curves of tree1 & tree2 (normalized in time and for the number of lineages)</i>
------------------------------	--

Description

Calculates the exact, difference between the lineage through time curves of tree1 & tree2 (normalized in time and for the number of lineages)

Usage

```
nltt_diff_exact(tree1, tree2, distance_method = "abs",
                 ignore_stem = TRUE, log_transform = FALSE)
```

Arguments

<code>tree1</code>	(phylo) First phylogenetic tree
<code>tree2</code>	(phylo) Second phylogenetic tree
<code>distance_method</code>	(string) absolute, or squared distance?
<code>ignore_stem</code>	(logical) Should the phylogeny its stem be ignored?
<code>log_transform</code>	(logical) Should the number of lineages be log-transformed before normalization?

Value

(scalar) normalized Lineage-Through-Time difference between tree1 & tree2

Author(s)

Thijs Janzen

<code>nltt_diff_exact_brts</code>	<i>Calculates the exact difference between the nLTT curves of the branching times</i>
-----------------------------------	---

Description

Calculates the exact difference between the nLTT curves of the branching times

Usage

```
nltt_diff_exact_brts(b_times, lineages, b_times2, lineages2,
                      distance_method = "abs", time_unit = "since")
```

Arguments

b_times	branching times of the first phylogeny,
lineages	the number of lineages, usually one to the number of lineages
b_times2	branching times of the second phylogeny
lineages2	the number of lineages, usually one to the number of lineages
distance_method	how the difference between the two nLTTs is summed <ul style="list-style-type: none">• "abs": "the absolute distance between the two nLTTs is summed"• "squ": "the squared distance between the two nLTTs is summed"
time_unit	the time unit of the branching times <ul style="list-style-type: none">• "ago": "the branching times are positive, as these are in time units ago"• "since": "the branching times are negative, as these are in time units since present"

Author(s)

Thijs Janzen and Richel Bilderbeek

nltt_diff_exact_norm_brts

Calculates the exact difference between the nLTT curves of the branching times

Description

Calculates the exact difference between the nLTT curves of the branching times

Usage

```
nltt_diff_exact_norm_brts(b_times_n, lineages_n, b_times2_n, lineages2_n,
                           distance_method)
```

Arguments

b_times_n	branching times of the first phylogeny
lineages_n	the number of lineages, usually one to the number of lineages
b_times2_n	branching times of the second phylogeny
lineages2_n	the number of lineages, usually one to the number of lineages
distance_method	(string) absolute, or squared distance?

Author(s)

Thijs Janzen and Richel Bilderbeek

nltt_lines*Normalized version of the ape function ltt.lines.***Description**

This is a modified version of the `ape` function `ltt.lines`: add the normalized Lineage-Through-Time statistic of a phylogenetic tree to an already existing plot

Usage

```
nltt_lines(phy, ...)
```

Arguments

<code>phy</code>	an object of class "phylo"
...	further graphical arguments that can be passed to <code>lines()</code>

Author(s)

Thijs Janzen

Examples

```
data(exampleTrees)
nltt_plot(exampleTrees[[1]])
nltt_lines(exampleTrees[[2]], lty=2)
```

nltt_plot*Normalized version of the ape function ltt.plot***Description**

This function uses a modified version of the `ltt.plot` function from "ape" to plot the normalized number of lineages through normalized time, where the number of lineages is normalized by dividing by the number of tips of the tree, and the time is normalized by the total time between the most common recent ancestor and the present, such that $t(\text{MRCA}) = 0$ & $t(\text{present}) = 1$.

Usage

```
nltt_plot(
  phy, xlab = "Normalized Time", ylab = "Normalized Lineages", ...)
```

Arguments

phy	an object of class "phylo"
xlab	a character string (or a variable of mode character) giving the label for the <i>x</i> -axis (default is "Normalized Time").
ylab	a character string (or a variable of mode character) giving the label for the <i>y</i> -axis (default is "Normalized Lineages").
...	further graphical arguments that can be passed to plot()

Author(s)

Thijs Janzen

Examples

```
data(exampleTrees)
nltt_plot(exampleTrees[[1]])
```

stretch_nltt_matrix *Stretch matrix 'm' with a timestep resolution of 'dt'.*

Description

Stretch matrix 'm' with a timestep resolution of 'dt'.

Usage

```
stretch_nltt_matrix(m, dt, step_type)
```

Arguments

m	A matrix of 2 columns and at least 2 rows
dt	The resolution, a value $\in [0.0001, 1]$. If 'dt' is set to a very small value, this function will stop
step_type	can be 'lower' or 'upper'

Value

The stretched matrix

Author(s)

Richel Bilderbeek

Examples

```
m <- matrix( c(c(0.0, 1.0), c(0.5, 1.0)), ncol = 2, nrow = 2)
expected <- matrix(
  c(
    c(0.0, 0.5, 1.0), # Timepoints
    c(0.5, 0.5, 1.0) # Values
  ),
  ncol = 2, nrow = 3
)
result <- stretch_nltt_matrix(m = m, dt = 0.5, step_type = "lower")
testit::assert(identical(result, expected))
```

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