## Package 'dde'

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**Title** Solve Delay Differential Equations

Version 1.0.7

Description Solves ordinary and delay differential equations, where the objective function is written in either R or C. Suitable only for non-stiff equations, the solver uses a 'Dormand-Prince' method that allows interpolation of the solution at any point. This approach is as described by Hairer, Norsett and Wanner (1993) <ISBN:3540604529>. Support is also included for iterating difference equations.

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URL https://github.com/mrc-ide/dde

BugReports https://github.com/mrc-ide/dde/issues

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2 difeq

## **Contents**

Index																		14
	difeq_replicate . dopri dopri_interpolate		 	 					 									4 7
	difeq								 									2

difeq

Solve difference equation

## **Description**

Solve a difference (or recurrence) equation by iterating it a number of times.

## Usage

```
difeq(y, steps, target, parms, ..., n_out = 0L, n_history = 0L,
  grow_history = FALSE, return_history = n_history > 0, dllname = "",
  parms_are_real = TRUE, ynames = names(y), outnames = NULL,
  return_by_column = TRUE, return_initial = TRUE, return_step = TRUE,
  return_output_with_y = TRUE, restartable = FALSE,
  return_minimal = FALSE)

difeq_continue(obj, steps, y = NULL, ..., copy = FALSE, parms = NULL,
  return_history = NULL, return_by_column = NULL, return_initial = NULL,
  return_step = NULL, return_output_with_y = NULL, restartable = NULL)

yprev(step, i = NULL)
```

## **Arguments**

у	The initial state of the system. Must be a numeric vector (and will be passed through as.numeric by this function).
steps	A vector of steps to return the system at. The <i>first</i> step is taken as step zero, and the solution will be recorded at every other step in the vector. So to step a system from time zero to times 1, 2, 3,, n use 0:n. Must be integer values and will be passed through as . integer (which may truncate or otherwise butcher non-integer values).
target	The target function to advance. This can either be an R function taking arguments n, i, t, y, parms or be a scalar character with the name of a compiled function with arguments size_t n, size_t step, double time, const double *y, double *dydt, size_t n_out, double *output void *data.
parms	Parameters to pass through to the difference function
•••	Dummy arguments - nothing is allowed here, but this means that all further arguments <i>must</i> be specified by name (not order) so I can easily reorder them

later on.

difeq 3

n\_out The number of output variables (in addition to the difference equation variables).

If given, then an R function must return an attribute output with the output

variables.

n\_history The number of iterations of history to save during the simulation. By default, no

history is saved.

grow\_history Logical indicating if history should be grown during the simulation. If FALSE

(the default) then when history is used it is overwritten as needed (so only the most recent n\_history elements are saved. This may require some tuning so that you have enough history to run your simulation (i.e. to the longest delay) or an error will be thrown when it underflows. The required history length will vary with your delay sizes and with the timestep for dopri. If TRUE, then history will grow as the buffer is exhausted. The growth is geometric, so every time it reaches the end of the buffer it will increase by a factor of about 1.6 (see the ring documentation). This may consume more memory than necessary, but may be useful where you don't want to care about picking the history length carefully.

return\_history Logical indicating if history is to be returned. By default, history is returned if

n\_history is nonzero.

dllname Name of the shared library (without extension) to find the function func in the

case where func refers to compiled function.

parms\_are\_real Logical, indicating if parms should be treated as vector of doubles by func

(when it is a compiled function). If TRUE (the default), then REAL(parms), which is double\* is passed through. If FALSE then if params is an externalptr type (EXTPTRSXP) we pass through the result of R\_ExternalPtrAddr, otherwise we pass params through unmodified as a SEXP. In the last case, in your target function you will need to include <Rinternals.h>, cast to SEXP and then pull

it apart using the R API (or Rcpp).

ynames Logical, indicating if the output should be named following the names of the

input vector y. Alternatively, if ynames is a character vector of the same length

as y, these will be used as the output names.

outnames An optional character vector, used when n\_out is greater than 0, to name the

model output matrix.

return\_by\_column

Logical, indicating if the output should be returned organised by column (rather than row). This incurs a slight cost for transposing the matrices. If you can work

with matrices that are transposed relative to deSolve, then set this to FALSE. return\_initial Logical, indicating if the output should include the initial conditions. Specifying

FALSE avoids binding this onto the output.

senting step is included.

return\_output\_with\_y

Logical, indicating if the output should be bound together with the returned matrix y (as it is with deSolve). If FALSE, then output will be returned as the

attribute output.

restartable Logical, indicating if the problem should be restartable. If TRUE, then the return value of a simulation can be passed to difeq\_restart to continue the simula-

tion after arbitrary changes to the state or the parameters. Note that this is really

4 difeq\_replicate

only useful for delay difference equations where you want to keep the history but make changes to the parameters or to the state vector while keeping the history of the problem so far.

return\_minimal Shorthand option - if set to TRUE then it sets all of return\_by\_column, return\_initial,

return\_time, return\_output\_with\_y to FALSE

obj An object to continue from; this must be the results of running a simulation

with the option restartable = TRUE. Note that continuing a problem moves the pointer along in time (unless copy = TRUE, and that the incoming time (times[[1]])

must equal the previous time exactly.

copy Logical, indicating if the pointer should be copied before continuing. If TRUE,

this is non-destructive with respect to the data in the original pointer so the problem can be restarted multiple times. By default this is FALSE because there

is a (potentially very small) cost to this operation.

step The step to access (not that this is not an offset, but the actual step; within your

target function you'd write things like yprev(step - 1) to get the previous step.

i index within the state vector y to return. The index here is R-style base-1 index-

ing, so pass 1 in to access the first element. This can be left NULL to return all

the elements or a vector longer than one.

#### **Examples**

```
# Here is a really simple equation that just increases by 'p' each # time (p is the parameter vector and could be any R structure). rhs <- function(i, y, p) y + p y0 <- 1 \\ t <- 0:10 \\ p <- 5 \\ dde::difeq(y0, t, rhs, p)
```

difeq\_replicate

Solve difference equations repeatedly

## **Description**

Solve a replicate set of difference (or recurrence) equation by iterating it a number of times. This is a wrapper around difeq that does not (yet) do anything clever to avoid many allocations.

## Usage

```
difeq_replicate(n, y, steps, target, parms, ..., n_out = 0L,
    n_history = 0L, grow_history = FALSE, return_history = n_history > 0,
    dllname = "", parms_are_real = TRUE, ynames = NULL, outnames = NULL,
    return_by_column = TRUE, return_initial = TRUE, return_step = TRUE,
    return_output_with_y = TRUE, restartable = FALSE,
    return_minimal = FALSE, as_array = TRUE)
```

difeq\_replicate 5

#### **Arguments**

n Number of replicates. It is an error to request zero replicates.

y The initial state of the system. Must be either a numeric vector or a list of

numeric vectors. If the latter, it must have length n.

steps A vector of steps to return the system at. The *first* step is taken as step zero,

and the solution will be recorded at every other step in the vector. So to step a system from time zero to times 1, 2, 3, ..., n use 0:n. Must be integer values and will be passed through as.integer (which may truncate or otherwise butcher

non-integer values).

target The target function to advance. This can either be an R function taking ar-

guments n, i, t, y, parms or be a scalar character with the name of a compiled function with arguments size\_t n, size\_t step, double time, const double \*y, double \*dydt, size\_t n\_out, double \*output void \*data.

parms Parameters to pass through to the difference function

... Dummy arguments - nothing is allowed here, but this means that all further

arguments must be specified by name (not order) so I can easily reorder them

later on.

n\_out The number of output variables (in addition to the difference equation variables).

If given, then an R function must return an attribute output with the output

variables.

n\_history The number of iterations of history to save during the simulation. By default, no

history is saved.

grow\_history Logical indicating if history should be grown during the simulation. If FALSE

(the default) then when history is used it is overwritten as needed (so only the most recent n\_history elements are saved. This may require some tuning so that you have enough history to run your simulation (i.e. to the longest delay) or an error will be thrown when it underflows. The required history length will vary with your delay sizes and with the timestep for dopri. If TRUE, then history will grow as the buffer is exhausted. The growth is geometric, so every time it reaches the end of the buffer it will increase by a factor of about 1.6 (see the ring documentation). This may consume more memory than necessary, but may be useful where you don't want to care about picking the history length carefully.

return\_history Logical indicating if history is to be returned. By default, history is returned if

n\_history is nonzero.

dllname Name of the shared library (without extension) to find the function func in the

case where func refers to compiled function.

parms\_are\_real Logical, indicating if parms should be treated as vector of doubles by func

(when it is a compiled function). If TRUE (the default), then REAL(parms), which is double\* is passed through. If FALSE then if params is an externalptr type (EXTPTRSXP) we pass through the result of R\_ExternalPtrAddr, otherwise we pass params through unmodified as a SEXP. In the last case, in your target function you will need to include <Rinternals.h>, cast to SEXP and then pull

it apart using the R API (or Rcpp).

6 difeq\_replicate

ynames Logical, indicating if the output should be named following the names of the

input vector y. Alternatively, if ynames is a character vector of the same length

as y, these will be used as the output names.

outnames An optional character vector, used when n\_out is greater than 0, to name the

model output matrix.

return\_by\_column

Logical, indicating if the output should be returned organised by column (rather than row). This incurs a slight cost for transposing the matrices. If you can work with matrices that are transposed relative to deSolve, then set this to FALSE.

return\_initial Logical, indicating if the output should include the initial conditions. Specifying

FALSE avoids binding this onto the output.

senting step is included.

return\_output\_with\_y

Logical, indicating if the output should be bound together with the returned matrix y (as it is with deSolve). If FALSE, then output will be returned as the

attribute output.

restartable Logical, indicating if the problem should be restartable. If TRUE, then the return

value of a simulation can be passed to difeq\_restart to continue the simulation after arbitrary changes to the state or the parameters. Note that this is really only useful for delay difference equations where you want to keep the history but make changes to the parameters or to the state vector while keeping the history

of the problem so far.

return\_minimal Shorthand option - if set to TRUE then it sets all of return\_by\_column, return\_initial,

return\_time, return\_output\_with\_y to FALSE

as\_array (Defunct) Logical, indicating if the output should be converted into an array. If

TRUE then res[, ,i] will contain the i'th replicate, if FALSE then res[[i]] does instead. If both as\_array and restartable are TRUE, then the attributes ptr and restart\_data will be present as a list of restarting information for

difeq\_continue, though using these is not yet supported.

#### **Details**

It is not currently possible to replicate over a set of parameters at once yet; the same parameter set will be used for all replications.

The details of how replication is done here are all considered implementation details and are up for change in the future - in particular if the models are run in turn or simultaneously (and the effect that has on the random number stream). Logic around naming output may change in future too; note that varying names in the y here will have some unexpected behaviours.

## **Examples**

```
# Here is a really simple equation that does a random walk with # steps that are normally distributed: rhs <- function(i, y, p) y + runif(1) y0 <- 1 t <- 0:10
```

```
p <- 5
dde::difeq_replicate(10, y0, t, rhs, p)</pre>
```

dopri

Integrate ODE/DDE with dopri

#### **Description**

Integrate an ODE or DDE with dopri.

## Usage

```
dopri(y, times, func, parms, ..., n_out = 0L, output = NULL,
  rtol = 1e-06, atol = 1e-06, step_size_min = 0, step_size_max = Inf,
  step_size_initial = 0, step_max_n = 100000L,
 step_size_min_allow = FALSE, tcrit = NULL, event_time = NULL,
 event_function = NULL, method = "dopri5", stiff_check = 0,
 verbose = FALSE, callback = NULL, n_history = 0,
  grow_history = FALSE, return_history = n_history > 0, dllname = "",
 parms_are_real = TRUE, ynames = names(y), outnames = NULL,
  return_by_column = TRUE, return_initial = TRUE, return_time = TRUE,
  return_output_with_y = TRUE, return_statistics = FALSE,
  restartable = FALSE, return_minimal = FALSE)
dopri5(y, times, func, parms, ...)
dopri853(y, times, func, parms, ...)
dopri_continue(obj, times, y = NULL, ..., copy = FALSE, parms = NULL,
  tcrit = NULL, return_history = NULL, return_by_column = NULL,
 return_initial = NULL, return_statistics = NULL, return_time = NULL,
 return_output_with_y = NULL, restartable = NULL)
ylag(t, i = NULL)
```

#### **Arguments**

У	Initial conditions for the integration
times	Times where output is needed. Unlike deSolve we won't actually stop at these times, but instead interpolate back to get the result.
func	Function to integrate. Can be an R function of arguments t, y, parms, returning a numeric vector, or it can be the name or address of a C function with arguments size_t n, double t, const double *y, double *dydt, void *data.
parms	Parameters to pass through to the derivatives.
	Dummy arguments - nothing is allowed here, but this means that all further arguments <i>must</i> be specified by name (not order) so I can easily reorder them later on.

n\_out Number of "output" variables (not differential equation variables) to compute

via the routine output.

output The output routine; either an R function taking arguments t, y, parms or the

 $name/address\ of\ a\ C\ function\ taking\ arguments\ {\tt size\_t}\ {\tt n,\ double}\ {\tt t,\ const}$ 

double \*y,size\_t n\_out, double \*out, void \*data.

rtol The per-step relative tolerance. The total accuracy will be less than this.

atol The per-step absolute tolerance.

step\_size\_min The minimum step size. The actual minimum used will be the largest of the

absolute value of this step\_size\_min or .Machine\$double.eps. If the integration attempts to make a step smaller than this, it will throw an error by default, stopping the integration (note that this differs from the treatment of hmin in deSolve::lsoda). See allow\_step\_size\_min to change this behaviour.

step\_size\_max The largest step size. By default there is no maximum step size (Inf) so the

solver can take as large a step as it wants to. If you have short-lived fluctuations in your rhs that the solver may skip over by accident, then specify a smaller

maximum step size here (or use tcrit below).

step\_size\_initial

The initial step size. By default the integrator will guess the step size automati-

cally, but one can be given here instead.

step\_max\_n The maximum number of steps allowed. If the solver takes more steps than this

it will throw an error. Note the number of evaluations of func will be about 6

times the number of steps (or 11 times if using method = "dopri853").

step\_size\_min\_allow

Logical, indicating if when a step size is driven down to step\_size\_min we

should allow it to proceed. This is the behaviour in of hmin in deSolve::lsoda.

tcrit An optional vector of critical times that the solver must stop at (rather than

interpolating over). This can include an end time that we can't go past, or points within the integration that must be stopped at exactly (for example cases where the derivatives change abruptly). Note that this differs from the interpretation of this parameter in deSolve; there tcrit is a single time that integration may not go past — with dde we never go past the final time, and this is just for times that

fall within the range of times in times.

event\_time Vector of times to fire events listed in event\_function at

event\_function Function to fire at events. For R models (func is an R function and dllname

is empty), this must be either a single R function (same function for all events) or a list of R functions. For C models, this must be a single C function (same

requirements as func or output or a list/vector of these as appropriate).

method The integration method to use, as a string. The supported methods are "dopri5"

(5th order method with 4th order dense output) and "dopri853" (8th order method with 7th order output and embedded 5th and 3rd order schemes). Alter-

natively, use the functions dopri5 or dopri853 which simply sets this argument.

stiff\_check How often to check that the problem has become stiff. If zero, then the problem is never checked, and if positive then the problem is checked every stiff\_check

accepted steps. The actual check is based off the algorithm in Hairer's implementation of the solvers and may be overly strict, especially for delay equations

with the 853 method (in my limited experience with it).

verbose Be verbose, and print information about each step. This may be useful for

learning about models that misbehave. Valid values are TRUE (enable debugging) or FALSE (disable debugging) or use one of dopri:::VERBOSE\_QUIET, dopri:::VERBOSE\_STEP or VERBOSE:::VERBOSE\_EVAL. If an R function is provided as the argument callback then this function will also be called at each

step or evaluation (see below for details).

callback Callback function that can be used to make verbose output more useful. This can

be used to return more information about the evaluation as it proceeds, generally as information printed to the screen. The function must accept arguments t, y

and dydt. See Details for further information.

n\_history Number of history points to retain. This needs to be greater than zero for delay differential equations to work. Alternatively, this may be greater than zero to

return model outputs that can be inspected later.

grow\_history Logical indicating if history should be grown during the simulation. If FALSE

> (the default) then when history is used it is overwritten as needed (so only the most recent n\_history elements are saved. This may require some tuning so that you have enough history to run your simulation (i.e. to the longest delay) or an error will be thrown when it underflows. The required history length will vary with your delay sizes and with the timestep for dopri. If TRUE, then history will grow as the buffer is exhausted. The growth is geometric, so every time it reaches the end of the buffer it will increase by a factor of about 1.6 (see the ring

> documentation). This may consume more memory than necessary, but may be useful where you don't want to care about picking the history length carefully.

return\_history Logical indicating if history should be returned alongside the output or discarded. By default, history is retained if n\_history is greater than 0, but that

> might change (and may not be desirable unless you plan on actually using it). Name of the shared library (without extension) to find the function func (and

> output if given) in the case where func refers to compiled function.

parms\_are\_real Logical, indicating if parms should be treated as vector of doubles by func (when it is a compiled function). If TRUE (the default), then REAL(parms), which is double\* is passed through. If FALSE then if params is an externalptr type (EXTPTRSXP) we pass through the result of R\_ExternalPtrAddr, otherwise we pass params through unmodified as a SEXP. In the last case, in your target function you will need to include <Rinternals.h>, cast to SEXP and then pull it apart using the R API (or Rcpp).

> Logical, indicating if the output should be named following the names of the input vector y. Alternatively, if ynames is a character vector of the same length as y, these will be used as the output names.

> An optional character vector, used when n\_out is greater than 0, to name the model output matrix.

Logical, indicating if the output should be returned organised by column (rather

than row). This incurs a slight cost for transposing the matrices. If you can work with matrices that are transposed relative to deSolve, then set this to FALSE.

return\_initial Logical, indicating if the output should include the initial conditions. Specifying FALSE avoids binding this onto the output.

dllname

ynames

outnames

return\_by\_column

return\_time Logical, indicating if a row (or column if return\_by\_column is TRUE) representing time is included. If FALSE, this is not added.

return\_output\_with\_y

Logical, indicating if the output should be bound together with the returned matrix y (as it is with deSolve). If FALSE, then output will be returned as the attribute output.

return\_statistics

Logical, indicating if statistics about the run should be included. If TRUE, then an integer vector containing the number of target evaluations, steps, accepted steps and rejected steps is returned (the vector is named).

restartable

Logical, indicating if the problem should be restartable. If TRUE, then the return value of an integration can be passed to dopri\_restart to continue the integration after arbitrary changes to the state or the parameters. Note that when using delay models, the integrator is fairly naive about how abrupt changes in the state space are dealt with, and may perform very badly with method = "dopri853" which assumes a fairly smooth problem. Note that this is really only useful for delay differential equations where you want to keep the history but make changes to the parameters or to the state vector while keeping the history of the problem so far.

return\_minimal Shorthand option - if set to TRUE then it sets all of return\_by\_column, return\_initial, return\_time, return\_output\_with\_y to FALSE

obj An object to continue from; this must be the results of running an integration with the option restartable = TRUE. Note that continuing a problem moves the pointer along in time (unless copy = TRUE, and that the incoming time (times[[1]]) must equal the previous time *exactly*.

Logical, indicating if the pointer should be copied before continuing. If TRUE, this is non-destructive with respect to the data in the original pointer so the problem can be restarted multiple times. By default this is FALSE because there

is a (potentially very small) cost to this operation.

t The time to access (not that this is not an offset, but the actual time; within your target function you'd write things like tlag(t - 1) to get 1 time unit ago.

index within the state vector y to return. The index here is R-style base-1 indexing, so pass 1 in to access the first element. This can be left NULL to return all the elements or a vector longer than one.

#### **Details**

i

copy

Like deSolve::lsoda, this function has *many* arguments. This is far from ideal, and I would welcome any approach for simplifying it a bit.

The options return\_by\_column, return\_initial, return\_time, return\_output\_with\_y exist because these options all carry out modifications of the data at the end of solving the ODE and this can incur a small but measurable cost. When solving an ODE repeatedly (e.g., in the context of an MCMC or optimisation) it may be useful to do as little as possible. For simple problems this can save around 5-10% of the total computational time (especially the transpose). The shorthand option return\_minimal will set all to FALSE when used.

#### Value

At present the return value is transposed relative to deSolve. This might change in future.

#### Verbose output and callbacks

Debugging a failed integration can be difficult, but dopri provides a couple of tools to get more information about where a failure might have occurred. Most simply, one can pass verbose = TRUE which will print information about the time and the step size at each point just before the step is stated. Passing in verbose = dde:::VERBOSE\_EVAL will print information just before every evaluation of the target function (there are several evaluations per step).

However, this does not provide information about the state just before failure. To get that, one must provide a callback function - this is an R function that will be called just before a step or evaluation (based on the value of the verbose argument) in place of the default print. Define a callback function with arguments t, h and y where t is the time (beginning of a step or location of an evaluation), h is the step size (or NA for an evaluation) and y is the state at the point of the step or evaluation. Your callback function can do anything - you can print to the screen (using cat or message), you can store results using a closure and <<- or you could conditionally use a browser() call to debug interactively. However, it is not possible for the callback to affect the solution short of throwing an error and interrupting it. See the Examples for an example of use.

#### See Also

dopri\_interpolate which can be used to efficiently sample from output of dopri, and the package vignette which shows in more detail how to solve delay differential equations and to use compiled objective functions.

## **Examples**

```
# The lorenz attractor:
lorenz <- function(t, y, p) {</pre>
  sigma <- p[[1L]]
  R \leftarrow p[[2L]]
  b <- p[[3L]]
  c(sigma * (y[[2L]] - y[[1L]]),
    R * y[[1L]] - y[[2L]] - y[[1L]] * y[[3L]],
    -b * y[[3L]] + y[[1L]] * y[[2L]])
}
p < -c(10, 28, 8 / 3)
y0 <- c(10, 1, 1)
tt <- seq(0, 100, length.out = 40000)
y <- dde::dopri(y0, tt, lorenz, p, return_time = FALSE)</pre>
plot(y[, c(1, 3)], type = "l", lwd = 0.5, col = "#00000066")
# If we want to print progress as the integration progresses we can
# use the verbose argument:
y \leftarrow dde::dopri(y0, c(0, 0.1), lorenz, p, verbose = TRUE)
# Or print the y values too using a callback:
```

12 dopri\_interpolate

dopri\_interpolate

Interpolate Dormand-Prince output

## **Description**

Interpolate the Dormand-Prince output after an integration. This only interpolates the core integration variables and not any additional output variables.

## Usage

```
dopri_interpolate(h, t)
```

## Arguments

h The interpolation history. This can be the output running dopri with return\_history = TRUE, or the history attribute of this object (retrievable with attr(res, "history")).

t The times at which interpolated output is required. These times must fall within the included history (i.e., the times that the original simulation was run) or an error will be thrown.

## **Details**

This decouples the integration of the equations and the generation of output; it is not necessary for use of the package, but may come in useful where you need to do (for example) root finding on the time course of a problem, or generate minimal output in some cases and interrogate the solution more deeply in others. See the examples and the package vignette for a full worked example.

#### Author(s)

Rich FitzJohn

## **Examples**

```
# Here is the Lorenz attractor implemented as an R function
lorenz <- function(t, y, p) {
    sigma <- p[[1L]]
    R <- p[[2L]]
    b <- p[[3L]]
    c(sigma * (y[[2L]] - y[[1L]]),
        R * y[[1L]] - y[[2L]] - y[[1L]] * y[[3L]],
        -b * y[[3L]] + y[[1L]] * y[[2L]])
}</pre>
```

dopri\_interpolate 13

```
# Standard parameters and a reasonable starting point:
p <- c(10, 28, 8 / 3)
y0 <- c(10, 1, 1)
# Run the integration for times [0, 50] and return minimal output,
# but *do* record and return history.
y <- dopri(y0, c(0, 50), lorenz, p,
           n_history = 5000, return_history = TRUE,
           return_time = FALSE, return_initial = FALSE,
           return_by_column = FALSE)
# Very little output is returned (just 3 numbers being the final
# state of the system), but the "history" attribute is fairly
# large matrix of history information. It is not printed though
# as its contents should not be relied on. What does matter is
# the range of supported times printed (i.e., [0, 50]) and the
# number of entries (~2000).
# Generate an interpolated set of variables using this; first for
# 1000 steps over the full range:
tt <- seq(0, 50, length.out = 1000)
yy <- dopri_interpolate(y, tt)</pre>
plot(yy[, c(1, 3)], type = "l")
# Then for 50000
tt <- seq(0, 50, length.out = 50000)
yy <- dopri_interpolate(y, tt)</pre>
plot(yy[, c(1, 3)], type = "l")
```

# **Index**

```
difeq, 2, 4
difeq_continue (difeq), 2
difeq_replicate, 4
dopri, 7
dopri5 (dopri), 7
dopri853 (dopri), 7
dopri_continue (dopri), 7
dopri_interpolate, 11, 12
ylag (dopri), 7
yprev (difeq), 2
```