

# Package ‘rodeo’

March 27, 2021

**Type** Package

**NeedsCompilation** yes

**Title** A Code Generator for ODE-Based Models

**Version** 0.7.7

**Date** 2021-03-27

**Author** David Kneis <david.kneis@tu-dresden.de>

**Maintainer** David Kneis <david.kneis@tu-dresden.de>

**Description** Provides an R6 class and several utility methods to facilitate the implementation of models based on ordinary differential equations. The heart of the package is a code generator that creates compiled 'Fortran' (or 'R') code which can be passed to a numerical solver. There is direct support for solvers contained in packages 'deSolve' and 'rootSolve'.

**URL** <https://github.com/dkneis/rodeo>

**License** GPL (>= 2)

**Imports** R6, deSolve

**VignetteBuilder** knitr

**Suggests** knitr, rmarkdown, xtable, rootSolve

**SystemRequirements** The tools to run 'R CMD SHLIB' on 'Fortran' code. The used 'Fortran' compiler must support pointer initialization which is a feature of the 2008 standard. The compiler from Oracle Developer Studio 12.6 on Solaris 10 currently does not meet this requirement.

**RoxygenNote** 7.1.1

**Repository** CRAN

**Date/Publication** 2021-03-27 18:20:02 UTC

**R topics documented:**

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rodeo-package

*Package to Facilitate ODE-Based Modeling***Description**

This package provides methods to

- import a conceptual ODE-based model stored in tabular form (i.e. as text files or spreadsheets).
- generate source code (either R or Fortran) to be passed to an ODE-solver.
- visualize and export basic information about a model, e.g. for documentation purposes.

**Details**

Consult the package vignette for details. The concept of writing an ODE system in tabular/matrix form is nicely introduced, e. g., in the book of Reichert, P., Borchardt, D., Henze, M., Rauch, W., Shanahan, P., Somlyódy, L., and Vanrolleghem, P. A. (2001): River water quality model No. 1, IWA publishing, ISBN 9781900222822.

The current source code repository is <https://github.com/dkneis/rodeo>.

### Class and class methods

See [rodeo-class](#) for the rodeo class and the corresponding class methods.

### Non-class methods

Type `help(package="rodeo")` or see the links below to access the documentation of non-class methods contained in the package.

- [forcingFunctions](#) Generation of forcing functions in Fortran.
- [exportDF](#) Export of data frames as TEX or HTML code.
- [stoiCreate](#) Creates a stoichiometry matrix from a set of chemical reactions.
- [stoiCheck](#) Validates a stoichiometry matrix by checking for conservation of mass.

### Author(s)

<david.kneis@tu-dresden.de>

---

compile

*Generate Executable Code*

---

### Description

Creates and 'compiles' a function for use with numerical methods from package [deSolve](#) or [rootSolve](#).

### Arguments

|         |  |
|---------|--|
| sources | Name(s) of source files(s) where functions appearing in process rates or stoichiometric factors are implemented. Can be NULL if no external functions are required, the name of a single file, or a vector of file names. See notes below.   |
| fortran | If TRUE, Fortran code is generated and compiled into a shared library. If FALSE, R code is generated.  |
| target  | Name of a 'target environment'. Currently, 'deSolve' is the only supported value.  |
| lib     | File path to be used for the generated library (without the platform specific extension). Note that any uppercase characters will be converted to lowercase. By default, the file is created in R's temporary folder under a random name.  |
| reuse   | If TRUE, an already existing library file will be loaded. Use this to prevent unnecessary re-compilation but note that R is likely to crash in case of any mismatches between the object and the existing library. Default is FALSE, i.e. the library is unconditionally build from scratch. |

### Value

`invisible(NULL)`

**Note**

The expected language of the external code passed in `sources` depends on the value of `fortran`.

If `fortran` is `FALSE`, R code is generated and made executable by `eval` and `parse`. Auxiliary code passed via `sources` is made available via `source`. The created R function is stored in the object.

If `fortran` is `TRUE`, the external code passed in `sources` must implement a module with the fixed name `'functions'`. This module must contain all user-defined functions referenced in process rates or stoichiometric factors.

If `fortran` is `TRUE`, a shared library is created. The library is immediately loaded with `dyn.load` and it is automatically unloaded with `dyn.unload` when the object's `finalize` method is called.

The name of the library (base name without extension) as well as the name of the function to compute the derivatives are stored in the object. These names can be queried with the `libName` and `libFunc` methods, respectively. Unless a file path is specified via the `lib` argument, the library is created in the folder returned by `tempdir` under a unique random name.

**Author(s)**

<david.kneis@tu-dresden.de>

**See Also**

This method internally calls `generate`.

**Examples**

```
data(vars, pars, funs, pros, stoi)
model <- rodeo$new(vars, pars, funs, pros, stoi, dim=c(1))
# This would trigger compilation assuming that 'functionsCode.f95' contains
# a Fortran implementation of all functions; see vignette for full example
## Not run:
model$compile(sources="functionsCode.f95")

## End(Not run)
```

---

dynamics

*Numerical Integration*

---

**Description**

Compute a dynamic solution with the numerical algorithms from package `deSolve`.

**Arguments**

|                       |  |
|-----------------------|--|
| <code>times</code>    | Times of interest (numeric vector).  |
| <code>fortran</code>  | Switch between compiled Fortran and R code (logical). Default is <code>TRUE</code> .     |
| <code>proNames</code> | Assign names to output columns holding the process rates? Default is <code>TRUE</code> . |
| <code>...</code>      | Auxiliary arguments passed to <code>ode</code> . See notes below.                        |

**Value**

The matrix returned by the integrator (see [ode](#)).

**Note**

This method can only be used after [compile](#) has been called.

The `...` argument should *not* be used to assign values to any of `y`, `parms`, `times`, `func`. If `fortran` is `TRUE` it should also not assign values to `dllname`, `nout`, or `outnames`. All these arguments of [ode](#) get their appropriate values automatically.

**Author(s)**

<david.kneis@tu-dresden.de>

**See Also**

Use [step](#) for integration over a single time step with a built-in, Fortran-based solver.

---

exportDF

*Export a Data Frame as HTML/TEX Code*

---

**Description**

Generates code to include tabular data in a tex document or web site.

**Usage**

```
exportDF(  
  x,  
  tex = FALSE,  
  colnames = NULL,  
  width = NULL,  
  align = NULL,  
  funHead = NULL,  
  funCell = NULL,  
  lines = TRUE,  
  indent = 2  
)
```

**Arguments**

|                       |  |
|-----------------------|--|
| <code>x</code>        | The data frame being exported.   |
| <code>tex</code>      | Logical. Allows to switch between generation of TEX code and HTML.   |
| <code>colnames</code> | Displayed column names. If <code>NULL</code> , the original names of <code>x</code> are used. Otherwise it must be a named vector with element names corresponding to column names in <code>x</code> . It is OK to supply alternative names for selected columns only. |

|         |   |
|---------|---|
| width   | Either NULL (all columns get equal width) or a named vector with element names corresponding to column names in <code>x</code> . If <code>tex == TRUE</code> , values (between 0 and 1) are needed for columns with align code 'p' only. They are interpreted as a multiplier for <code>'\textwidth'</code> . If <code>tex == FALSE</code> , values (between 0 and 100) should be supplied for all columns of <code>x</code> .        |
| align   | Either NULL (to use automatic alignment) or a named vector with element names corresponding to column names in <code>x</code> . If <code>tex == FALSE</code> valid alignment codes are 'left', 'right', 'center'. If <code>tex == TRUE</code> valid alignment codes are 'l', 'r', 'c', and 'p'. For columns with code 'p' a corresponding value of width should be set. It is OK to supply alignment codes for selected columns only. |
| funHead | Either NULL or a list of functions whose names correspond to column names of <code>x</code> . The functions should have a single formal argument; the respective column names of <code>x</code> are used as the actual arguments. It is OK to supply functions for selected columns only (an empty function is applied to the remaining columns). See below for some typical examples.  |
| funCell | Like funHead but these functions are applied to the cells in columns rather than to the column names.   |
| lines   | Logical. Switches table borders on/off.   |
| indent  | Integer. Number of blanks used to indent the generated code.  |

**Value**

A character string (usually needs to be exported to a file).

**Note**

The functions `funHead` and `funCell` are useful to apply formatting or character replacement. For example, one could use

```
function(x) {paste0("\\bold{", toupper(x), "}")}
```

to generate bold, uppercase column names in a TEX table.

**Author(s)**

David Kneis <david.kneis@tu-dresden.de>

**See Also**

The `xtable` package provides similar functionality with more sophisticated options. Consider the 'pandoc' software do convert documents from one markup language to another one. Finally, consider the latex package 'datatools' for direct inclusion of delimited text files (e.g. produced by `write.table`) in tex documents.

**Examples**

```
# Create example table
df <- data.frame(stringsAsFactors=FALSE, name= c("growth", "dead"),
  unit= c("1/d", "1/d"), expression= c("r * N * (1 - N/K)", "d * N"))
```

```

# Export as TEX: header in bold, 1st colum in italics, last column as math
tex <- exportDF(df, tex=TRUE,
  colnames=c(expression="process rate expression"),
  width=c(expression=0.5),
  align=c(expression="p"),
  funHead=setNames(replicate(ncol(df),
    function(x){paste0("\\textbf{" , x, "}")}),names(df)),
  funCell=c(name=function(x){paste0("\\textit{" , x, "}")},
    expression=function(x){paste0("$" , x, "$")})
)
cat(tex, "\n")

# Export as HTML: non-standard colors are used for all columns
tf <- tempfile(fileext=".html")
write(x= exportDF(df, tex=FALSE,
  funHead=setNames(replicate(ncol(df),
    function(x){paste0("<font color='red'>" , x, "</font>")}),names(df)),
  funCell=setNames(replicate(ncol(df),
    function(x){paste0("<font color='blue'>" , x, "</font>")}),names(df))
), file=tf)
## Not run:
  browseURL(tf)
  file.remove(tf)

## End(Not run)

```

---

finalize

*Clean-up a rodeo Object*


---

## Description

Clean-up method for objects of the [rodeo-class](#).

## Value

The method is called implicitly for its side effects when a [rodeo](#) object is destroyed.

## Note

At present, the method just unloads the object-specific shared libraries created with the [compile](#) or [initStepper](#) methods.

## Author(s)

<david.kneis@tu-dresden.de>

---

 forcingFunctions

*Generation of Forcing Functions in Fortran*


---

### Description

Generates Fortran code to return the current values of forcing functions based on interpolation in tabulated time series data.

### Usage

```
forcingFunctions(x)
```

### Arguments

`x` Data frame with columns 'name', 'file', 'column', 'mode', 'default'. See below for expected entries.

### Value

A character string holding generated Fortran code. Must be written to disk, e.g. using `write`, prior to compilation.

### Note

The fields of the input data frame are interpreted as follows:

- `name` Name of the forcing function as declared in the table of functions.
- `file` Name of the text file containing the time series data either as an absolute or relative path. Time information is expected as numeric values in the first column (e.g. as number of seconds after some reference date). The period is used as the decimal character in floating point numbers, numeric values can also be given in scientific format (e.g. as `0.314e+1`). Allowed column delimiters are blank, tab, or comma. A sequence of white spaces collapses to a single delimiter but this is not the case for commas. It is strictly recommended to use a consistent delimiter character within a particular file. Blank lines are allowed everywhere in the file, comment lines must start with a '#'. The first non-blank, non-comment line is interpreted as column headers and the name of the first column (holding time info) is essentially ignored.
- `column` Name of the column in `file` from which data are to be read.
- `mode` Integer code to control how the interpolation is performed. Use 0 for constant interpolation with full weight given to the value at the end of a time interval. Use 1 for constant interpolation with full weight given to the value at the begin of a time interval. Any other values (`< 0` or `> 1`) result in linear interpolation with weights being set automatically.
- `default` Logical. If `FALSE`, the generated function has the interface 'f(time)'. If `TRUE`, the generated function has a two-argument interface 'f(time, z)'. If the actual argument 'z' is `NaN`, the function behaves just like the single-argument version, i.e. interpolation in tabulated data is performed. If 'z' is not `NaN`, the function returns the value of 'z'.



The generated code provides a single module named 'forcings' which defines as many forcing functions as there are rows in  $x$ . The module 'forcings' needs to be made available to the compiler (either at the command line or via inclusion in another file with Fortran's include mechanism). In addition, it must be referenced in the module 'functions' with an appropriate 'use' statement (see example below).

The generated function return scalar values of type double precision. If an error condition is encountered, the return value of a functions equals the largest possible double precision value (generated by Fortran's 'huge' function). In addition, errors trigger calls of the subroutines 'rexit' (at top level) or 'rwarn' (at lower levels). These two functions are available automatically if the Fortran code is compiled using 'R CMD SHLIB'. Otherwise, the two functions need to be defined (see examples below).

In the two-argument version, the second argument is tested against NaN using 'ISNAN'. This function is not part of the Fortran standard but it is supported by most compilers, including gfortan. The Fortran 2003 standard conformal function would be 'IS\_IEEE\_NAN' which is not yet supported by compiler versions normally installed with R (March 2016).

### Author(s)

David Kneis <david.kneis@tu-dresden.de>

### Examples

```
## Not run:
! Example of a Fortran file to define functions
include 'forcings.f95' ! include generated forcings file in compilation
module functions
use forcings          ! make forcings available as functions
implicit none
contains
! ... any non-forcing functions go here ...
end module

## End(Not run)

## Not run:
! Definition of 'rexit' and 'rwarn' for testing of the generated code
! outside of R
subroutine rexit (x)
  character(len=*), intent(in):: x
  write(*,*) "ERROR: ",trim(adjustl(x))
  stop 1
end subroutine

subroutine rwarn (x)
  character(len=*), intent(in):: x
  write(*,*) "WARNING: ",trim(adjustl(x))
end subroutine

## End(Not run)
```

---

 funs

*Declaration of Functions*


---

### Description

Declaration of functions referenced at the ODE's right hand sides of the bacteria growth example model.

### Format

A data frame with the following fields:

- name : Name of the function
- units : Unit of the return value
- description : Short description (text)

---

 generate

*Code Generator*


---

### Description

This is a low-level method to translate the ODE-model specification into a function that computes process rates and the state variables derivatives (either in R or Fortran). You probably want to use the high-level method [compile](#) instead, which uses `generate` internally.

### Arguments

|      |   |
|------|---|
| lang | Character string to select the language of the generated source code. Currently either 'f95' (for Fortran) or 'r' (for R).                  |
| name | Name for the generated function (character string). It should start with a letter, optionally followed by letters, numbers, or underscores. |

### Value

The generated source code as a string. Must be written to disk, e.g. using [write](#), prior to compilation.

### Note

Details of this low-level method may change in the future.

### Author(s)

<david.kneis@tu-dresden.de>

**See Also**

See other methods of the [rodeo-class](#), especially [compile](#) which internally uses this method.

**Examples**

```
data(vars, pars, funs, pros, stoi)
model <- rodeo$new(vars, pars, funs, pros, stoi, dim=c(1))
fortranCode <- model$generate(lang="f95")
## Not run:
write(fortranCode, file="")

## End(Not run)
```

---

 getPars

*Query Values of Parameters*


---

**Description**

Query values of parameters of a [rodeo](#)-based model.

**Arguments**

|          |  |
|----------|--|
| asArray  | Logical. If FALSE, the values of parameters are returned as vector irrespective of the model's spatial resolution. If TRUE, the values are returned as an <a href="#">array</a> with properly named dimensions. The array's last dimension represents the parameters and its first (fastest cycling) dimension, if any, refers to the model's highest spatial dimension.   |
| useNames | Logical. Used to enable/disable element names for the return vector when asArray is FALSE. The names follow the pattern 'x.i.j' where 'x' is the parameter name and 'i', 'j' are indices of the sub-units in the first and second spatial dimension. The actual suffix is controlled by the number of dimensions and in the 0-dimensional case, no suffix is applied at all, i.e. the pure parameter names are used to label the elements of the vector. If isArray is TRUE, this argument is simply ignored, hence the dimensions of a returned array are always named. |

**Value**

A numeric vector or array.

**Author(s)**

<david.kneis@tu-dresden.de>

**See Also**

The corresponding 'set' method is [setPars](#) and examples can be found there. Use [getVar](#)s to query the values of variables rather than parameters.

---

|         |  |
|---------|--|
| getVars | <i>Query Values of State Variables</i> |
|---------|--|

---

**Description**

Query values of state variables of a [rodeo](#)-based model.

**Arguments**

|          |  |
|----------|--|
| asArray  | Logical. If FALSE, the values of variables are returned as vector irrespective of the model's spatial resolution. If TRUE, the values are returned as an <a href="#">array</a> with properly named dimensions. The array's last dimension represents the variables and its first (fastest cycling) dimension, if any, refers to the model's highest spatial dimension.   |
| useNames | Logical. Used to enable/disable element names for the return vector when asArray is FALSE. The names follow the pattern 'x.i.j' where 'x' is the variable name and 'i', 'j' are indices of the sub-units in the first and second spatial dimension. The actual suffix is controlled by the number of dimensions and in the 0-dimensional case, no suffix is applied at all, i.e. the pure variable names are used to label the elements of the vector. If isArray is TRUE, this argument is simply ignored, hence the dimensions of a returned array are always named. |

**Value**

A numeric vector or array.

**Author(s)**

<david.kneis@tu-dresden.de>

**See Also**

The corresponding 'set' method is [setVars](#) and examples can be found there. Use [getPars](#) to query the values of parameters rather than variables.

---

|            |                                  |
|------------|----------------------------------|
| initialize | <i>Initialize a rodeo Object</i> |
|------------|----------------------------------|

---

**Description**

Initializes an object of the [rodeo-class](#) with data frames holding the specification of an ODE system.

**Arguments**

|                       |   |
|-----------------------|---|
| <code>vars</code>     | Declaration of state variables appearing in the ODE system. Data frame with mandatory columns 'name', 'unit', 'description'.  |
| <code>pars</code>     | Declaration of parameters (i.e. constants) appearing in the ODE system. Data frame with the same mandatory columns as <code>vars</code> .   |
| <code>funs</code>     | Declaration of functions being referenced in the ODE system. Data frame with the same mandatory columns as <code>vars</code> or <code>NULL</code> if no function calls are present at the ODEs' right-hand sides.   |
| <code>pros</code>     | Declaration of process rates. Data frame with mandatory columns 'name', 'unit', 'description', 'expression'.  |
| <code>stoi</code>     | Declaration of stoichiometric factors. A data frame with mandatory columns 'variable', 'process', 'expression', if <code>asMatrix</code> is <code>FALSE</code> . The 'expression' column holds the stoichiometric factors. If <code>asMatrix</code> is <code>TRUE</code> , this must be a matrix of type character with row names (processes) and column names (variables). Empty or <code>NA</code> matrix elements are interpreted as zero stoichiometry factors. |
| <code>asMatrix</code> | Logical. Specifies whether stoichiometry information is given in matrix or data base format.  |
| <code>dim</code>      | An integer vector, specifying the number of boxes in each spatial dimension. Use <code>c(1)</code> to create a zero-dimensional (i.e. single-box) model. This is the default. Use, e.g. <code>c(5)</code> to create a 1-dimensional model with 5 boxes. To create, e.g., a 2-dimensional model with 4 x 5 boxes, use <code>c(4, 5)</code> .   |

**Value**

The method is called implicitly for its side effects when a `rodeo` object is instantiated with `new`. It has no accessible return value.

**Note**

The mandatory fields of the input data frames should be of type character. Additional fields may be present in these data frames and the contents becomes part of the `rodeo` object. The 'expression' fields of `pros` and `stoi` (or the contents of the stoichiometry matrix) should be valid mathematical expressions in R and Fortran. These can involve the names of declared state variables, parameters, and functions as well as numeric constants or basic math operators. Branching or loop constructs are not allowed (but these can appear inside referenced functions). There are currently few reserved words that cannot be used as variable, parameter, function, or process names. The reserved words are 'time', 'left', and 'right'.

Initialization does not assign numeric values to state variables or parameters. Use the dedicated methods `setVars` and `setPars` for that purpose.

**Author(s)**

<david.kneis@tu-dresden.de>

**See Also**

See the package vignette for examples.

**Examples**

```
data(vars, pars, funs, pros, stoi)
model <- rodeo$new(vars, pars, funs, pros, stoi, dim=c(1))
print(model)
```

---

|             |                                       |
|-------------|---------------------------------------|
| initStepper | <i>Initialize Internal ODE Solver</i> |
|-------------|---------------------------------------|

---

**Description**

Initializes rodeo's built-in ODE solver. This method must be called prior to using [step](#).

**Arguments**

|         |  |
|---------|--|
| sources | Name(s) of fortran source file(s) where a module with the fixed name 'functions' is implemented. This module must contain all user-defined functions referenced in process rates or stoichiometric factors. Can be NULL, the name of a single file, or a vector of file names if the Fortran code is split over several files. |
| method  | Name of a the solver. Currently, 'rk5' is the only supported value (Runge-Kutta method of Cash and Karp).  |

**Value**

invisible(NULL)

**Note**

After this method was called, [step](#) can be used to perform the integration.

**Author(s)**

<david.kneis@tu-dresden.de>

**See Also**

To perform integration with the solvers from package [deSolve](#) use [compile](#) instead of this method.

---

|         |  |
|---------|--|
| libFunc | <i>Return name of library function</i> |
|---------|--|

---

**Description**

Return the name of the library function for use with [deSolve](#) or [rootSolve](#) methods.

**Value**

The name of the function to compute derivatives which is contained in the library built with [compile](#). This name has to be supplied as the func argument of the solver methods in [deSolve](#) or [rootSolve](#).

**Author(s)**

<david.kneis@tu-dresden.de>

---

|         |                            |
|---------|----------------------------|
| libName | <i>Return library name</i> |
|---------|----------------------------|

---

**Description**

Return the pure name of the shared library for use with [deSolve](#) or [rootSolve](#) methods.

**Value**

The base name of the shared library file created with [compile](#) after stripping of the the platform specific extension. This name has to be supplied as the dllname argument of the solver methods in [deSolve](#) or [rootSolve](#).

**Author(s)**

<david.kneis@tu-dresden.de>

---

|      |                                  |
|------|----------------------------------|
| pars | <i>Declaration of Parameters</i> |
|------|----------------------------------|

---

**Description**

Declaration of parameters of the bacteria growth example model.

**Format**

A data frame with the following fields:

- name : Name of the parameter
- units : Unit
- description : Short description (text)

---

plotStoichiometry      *Plot Qualitative Stoichiometry Matrix*

---

### Description

Visualizes the stoichiometry matrix using standard plot methods. The sign of stoichiometric factors is displayed as upward and downward pointing triangles, optionally colored.

### Arguments

|             |   |
|-------------|---|
| box         | A vector of positive integers pointing to a spatial sub-unit of the model.                    |
| time        | Time. The value is ignored in the case of autonomous models.                                  |
| cex         | Character expansion factor.   |
| shade       | Logical. If TRUE, the dependence of process rates on state variables is indicated by shading. |
| colPositive | Color for positive stoichiometric factors.  |
| colNegative | Color for negative stoichiometric factors.  |
| colGrid     | Color of a grid (can be identical to background color).                                       |
| colShade    | Shading color, if used.   |

### Note

The values of state variables and parameters must have been set using the [setVars](#) and [setPars](#) methods. If the stoichiometric factors are mathematical expressions involving function references, these functions must be defined in R (even if the numerical computations are based on generated Fortran code).

### Author(s)

<david.kneis@tu-dresden.de>

### See Also

See other methods of the [rodeo-class](#) or [stoichiometry](#) for computing the stoichiometric factors only. Alternative options for displaying stoichiometry information are described in the package vignette.

### Examples

```
data(vars, pars, funs, pros, stoi)
model <- rodeo$new(vars, pars, funs, pros, stoi, dim=c(1))
model$setVars(c(bac=0.1, sub=0.5))
model$setPars(c(mu=0.8, half=0.1, yield= 0.1, vol=1000, flow=50, sub_in=1))
monod <- function(c,h) {c/(c+h)}
model$plotStoichiometry(box=c(1))
```



---

pros *Declaration of Processes*

---

### Description

Definition of processes of the bacteria growth example model.

### Format

A data frame with the following fields:

- name : Name of the process
- units : Unit of the rate expression
- description : Short description (text)
- expression : Mathematical expression (as a string)

---

rodeo-class *rodeo Class*

---

### Description

This documents the rodeo class to represent an ODE-based model. See the [rodeo-package](#) main page or type `help(package="rodeo")` for an introduction to the package of the same name.

### Fields

`prosTbl` A data frame with fields 'name', 'unit', 'description', and 'expression' defining the process rates.

`stoiTbl` A data frame with fields 'variable', 'process', and 'expression' representing the stoichiometry matrix in data base format.

`varsTbl` A data frame with fields 'name', 'unit', 'description' declaring the state variables of the model. The declared names become valid identifiers to be used in the expression fields of `prosTbl` or `stoiTbl`.

`parsTbl` A data frame of the same structure as `vars` declaring the parameters of the model. The declared names become valid identifiers to be used in the expression fields of `prosTbl` or `stoiTbl`.

`funstTbl` A data frame of the same structure as `vars` declaring any functions referenced in the expression fields of `prosTbl` or `stoiTbl`.

`dim` Integer vector specifying the spatial dimensions.

`vars` Numeric vector, holding values of state variables.

`pars` Numeric vector, holding values of parameters.

## Class methods

For most of the methods below, a separate help page is available describing its arguments and usage.

- [initialize](#) Initialization method for new objects.
- [namesVars](#), [namesPars](#), [namesFuns](#), [namesPros](#) Functions returning the names of declared state variables, parameters, functions, and processes, respectively (character vectors). No arguments.
- [lenVars](#), [lenPars](#), [lenFuns](#), [lenPros](#) Functions returning the number of declared state variables, parameters, functions and processes, respectively (integer). No arguments.
- [getVarTable](#), [getParsTable](#), [getFunsTable](#), [getProsTable](#), [getStoiTable](#) Functions returning the data frames used to initialize the object. No arguments
- [getDim](#) Returns the spatial dimensions as an integer vector. No arguments.
- [compile](#) Compiles a Fortran library for use with numerical methods from packages [deSolve](#) or [rootSolve](#).
- [generate](#) Translate the ODE-model specification into a function that computes process rates and the state variables' derivatives (either in R or Fortran). Consider to use the high-level method [compile](#).
- [setVars](#) Assign values to state variables.
- [setPars](#) Assign values to parameters.
- [getVar](#) Returns the values of state variables.
- [getPars](#) Returns the values of parameters.
- [stoichiometry](#) Returns the stoichiometry matrix, either evaluated (numeric) or as text.
- [plotStoichiometry](#) Plots qualitative stoichiometry information.

## See Also

See the [rodeo-package](#) main page or type `help(package="rodeo")` to find the documentation of any non-class methods contained in the rodeo package.

## Examples

```
# Bacteria growth in a continuous flow culture
library("deSolve")

# Creation of model object
data(vars, pars, funs, pros, stoi)
model <- rodeo$new(vars, pars, funs, pros, stoi, dim=c(1))

# Parameters, initial values
model$setPars(c(mu=0.8, half=0.1, yield= 0.1, vol=1000, flow=50, sub_in=1))
model$setVars(c(bac=0.01, sub=0))

# Implementation of functions declared in 'funs'
monod <- function(c,h) {c/(c+h)}

# Creation of derivatives function
```

```
code <- model$generate(name="derivs", lang="r")
derivs <- eval(parse(text=code))

# Integration
times <- 0:96
out <- deSolve::ode(y=model$getVars(), times=times, func=derivs,
  parms=model$getPars())
colnames(out) <- c("time", model$namesVars(), model$namesPros())
plot(out)
```

---

setPars

*Assign Values to Parameters*

---

### Description

Assign values to parameters of a [rodeo](#)-based model.

### Arguments

**x** A numeric vector or array, depending on the model's spatial dimensions. Consult the help page of the sister method [setVars](#) for details on the required input.

### Value

NULL (invisible). The assigned numeric data are stored in the object and can be accessed by the [getPars](#) method.

### Note

Look at the notes and examples for the [setVars](#) method.

### Author(s)

<david.kneis@tu-dresden.de>

### See Also

The corresponding 'get' method is [getPars](#). Use [setVars](#) to assign values to variables rather than parameters. Consult the help page of the latter function for examples.

setVars

*Assign Values to State Variables***Description**

Assign values to state variables of a [rodeo](#)-based model.

**Arguments**

`x` A numeric vector or array, depending on the model's spatial dimensions. See details below.

**Value**

NULL (invisible). The assigned numeric data are stored in the object and can be accessed by the [getVars](#) method.

**Note**

For a 0-dimensional model (i.e. a model without spatial resolution), `x` must be a numeric vector whose length equals the number of state variables. The element names of `x` must match those returned by the object's `namesVars` method. See the examples for how to bring the vector elements into required order.

For models with a spatial resolution, `x` must be a numeric array of proper dimensions. The last dimension (cycling slowest) corresponds to the variables and the first dimension (cycling fastest) corresponds to the models' highest spatial dimension. Thus, `dim(x)` must be equal to `c(rev(obj$getDim()), obj$namesVars())`, where `obj` is the object whose variables are to be assigned. The names of the array's last dimension must match the return value of `obj$namesVars()`.

In the common 1-dimensional case, this just means that `x` must be a matrix with column names matching the return value of `obj$namesVars()` and as many rows as given by `obj$getDim()`.

**Author(s)**

<david.kneis@tu-dresden.de>

**See Also**

The corresponding 'get' method is [getVars](#). Use [setPars](#) to assign values to parameters rather than variables.

**Examples**

```
data(vars, pars, funs, pros, stoi)
x0 <- c(bac=0.1, sub=0.5)

# 0-dimensional model
model <- rodeo$new(vars, pars, funs, pros, stoi, dim=c(1))
model$setVars(x0)
```

```

print(model$getVars())

# How to sort vector elements
x0 <- c(sub=0.5, bac=0.1)          # doesn't match order of variables
model$setVars(x0[model$namesVars()])

# 1-dimensional model with 3 boxes
nBox <- 3
model <- rodeo$new(vars, pars, funs, pros, stoi, dim=c(nBox))
x1 <- matrix(rep(x0, each=nBox), nrow=nBox, ncol=model$lenVars(),
  dimnames=list(NULL, model$namesVars()))
model$setVars(x1)
print(model$getVars())
print(model$getVars(asArray=TRUE))

# 2-dimensional model with 3 x 4 boxes
model <- rodeo$new(vars, pars, funs, pros, stoi, dim=c(3,4))
x2 <- array(rep(x0, each=3*4), dim=c(4,3,model$lenVars()),
  dimnames=list(dim2=NULL, dim1=NULL, variables=model$namesVars()))
model$setVars(x2)
print(model$getVars())
print(model$getVars(asArray=TRUE))

```

---

step

*Numerical Integration Over a Single Time Step*


---

### Description

Performs integration over a single time step using a built-in ODE solver. At present, a single solver is implement with limited options. The interface of this method may change when support for other solvers is added.

### Arguments

|          |  |
|----------|--|
| t0       | Numeric. Initial time.   |
| h        | Numeric. Length of time step of interest.  |
| hmin     | Minimum tolerated internal step size. The default of NULL sets this to 10 times the value of <code>.Machine\$double.eps</code> .   |
| maxsteps | Maximum tolerated number of sub-steps.   |
| tol      | Numeric. Relative accuracy requested. This is currently a global value, i.e. one cannot set the accuracy per state variable.   |
| method   | String. Currently, 'rk5' is the only method implemented. This is a Runge-Kutta Cash-Karp solver adapted from Press et al. (2002), Numerical recipes in Fortran 90. It is designed to handle non-stiff problems only. |
| check    | Logical. Can be used to avoid repeated checks of arguments. This may increase performance in repeated calls.   |

**Value**

A named numeric vector holding the values of state variables and process rates in all boxes.

**Note**

This method can only be used after a call to `initStepper` has been made.

**Author(s)**

<david.kneis@tu-dresden.de>

**See Also**

Use `deSolve` for advanced solvers with more options and capabilities to handle stiff problems.

---

|      |                                       |
|------|---------------------------------------|
| stoi | <i>Specification of Stoichiometry</i> |
|------|---------------------------------------|

---

**Description**

Definition of the links between simulated processes and state variables in the bacteria growth example model.

**Format**

A data frame with the following fields:

- variable : Name of the state variable
- process : Name of the process
- expression : Mathematical expression (as a string)

---

|           |   |
|-----------|---|
| stoiCheck | <i>Validation of a Stoichiometry Matrix</i> |
|-----------|---|

---

**Description**

Validates the stoichiometry matrix by checking for conservation of mass (more typically conservation of moles).

**Usage**

```
stoiCheck(stoi, comp, env = globalenv(), zero = .Machine$double.eps * 2)
```

**Arguments**

|      |   |
|------|---|
| stoi | Stoichiometry matrix either in evaluated ( <b>numeric</b> ) or non-evaluated ( <b>character</b> ) form. A suitable matrix can be created with <code>stoiCreate</code> , for example.  |
| comp | Matrix defining the elemental composition of compounds. Column names of comp need to match column names of stoi (but additional columns are allowed and columns can be in different order). There must be one row per element whose balance is to be checked and the elements' names must appear as row names. The elements of the matrix specify how much of an element is contained in a certain amount of a compound. Typically, these are molar ratios. If one works with mass ratios (not being a good idea), the information in stoi must be based on mass concentrations as well. The elements of comp are treated as mathematical expressions. Any variables, functions, or operators needed to evaluate those expressions must be provided by the specified environment env. |
| env  | An environment or list supplying constants, functions, and operators needed to evaluate expressions in comp or stoi.  |
| zero | A number close to zero. If the absolute result value of a mass balance computation is less than this, the result is set to 0 (exactly).   |

**Value**

A numeric matrix with the following properties:

- There is one row for each process, thus the number and names of rows are the same as in stoi.
- There is one column per checked element, hence column names are equal to the row names of comp.
- A matrix element at position  $[i, k]$  represent the mass balance for the process in row  $i$  with respect to the element in column  $k$ . A value of zero indicates a close balance. Positive (negative) values indicate that mass is gained (lost) in the respective process.

**Author(s)**

David Kneis <david.kneis@tu-dresden.de>

**See Also**

Use `stoiCreate` to create a stoichiometry matrix from a set of reactions in common notation.

**Examples**

```
# Eq. 1 and 2 are from Soetaert et al. (1996), Geochimica et Cosmochimica
# Acta, 60 (6), 1019-1040. 'OM' is organic matter. Constants 'nc' and 'pc'
# represent the nitrogen/carbon and phosphorus/carbon ratio, respectively.
reactions <- c(
  oxicDegrad= "OM + O2 -> CO2 + nc * NH3 + pc * H3PO4 + H2O",
  denitrific= "OM + 0.8*HN03 -> CO2 + nc*NH3 + 0.4*N2 + pc*H3PO4 + 1.4*H2O",
  dissPhosp1= "H3PO4 <-> H + H2PO4",
  dissPhosp2= "H2PO4 <-> H + HPO4"
)
```

```

# Non-evaluated stoichiometry matrix
stoi <- stoiCreate(reactions, toRight="_f", toLeft="_b")
# Parameters ('nc' and 'pc' according to Redfield ratio)
pars <- list(nc=16/106, pc=1/106)
# Elemental composition
comp <- rbind(
  OM= c(C=1, N="nc", P="pc", H="2 + 3*nc + 3*pc"),
  O2= c(C=0, N=0, P=0, H=0),
  CO2= c(C=1, N=0, P=0, H=0),
  NH3= c(C=0, N=1, P=0, H=3),
  H3PO4= c(C=0, N=0, P=1, H=3),
  H2PO4= c(C=0, N=0, P=1, H=2),
  HPO4= c(C=0, N=0, P=1, H=1),
  H2O= c(C=0, N=0, P=0, H=2),
  HNO3= c(C=0, N=1, P=0, H=1),
  N2= c(C=0, N=2, P=0, H=0),
  H= c(C=0, N=0, P=0, H=1)
)
# We need the transposed form
comp <- t(comp)
# Mass balance check
bal <- stoiCheck(stoi, comp=comp, env=pars)
print(bal)
print(colSums(bal) == 0)

```

---

 stoichiometry

*Return the Stoichiometry Matrix*


---

## Description

Return and optionally evaluate the mathematical expression appearing in the stoichiometry matrix.

## Arguments

|      |  |
|------|--|
| box  | Either NULL or a vector of positive integers pointing to a spatial sub-unit of the model. If NULL, the mathematical expressions appearing in the stoichiometry matrix are not evaluated, hence, they are returned as character strings. If a spatial sub-unit is specified, a numeric matrix is returned. In the latter case, the values of state variables and parameters must have been set using the <a href="#">setVars</a> and <a href="#">setPars</a> methods. |
| time | Time. The value is ignored in the case of autonomous models.   |

## Value

A matrix of numeric or character type, depending on the value of box.

## Note

If the stoichiometric factors are mathematical expressions involving function references, these functions must be defined in R (even if the numerical computations are based on generated Fortran code).



**Author(s)**

<david.kneis@tu-dresden.de>

**See Also**

See other methods of the [rodeo-class](#) or [plotStoichiometry](#) for a graphical representation of the stoichiometric factors only.

**Examples**

```
data(vars, pars, funs, pros, stoi)
model <- rodeo$new(vars, pars, funs, pros, stoi, dim=c(1))
model$setPars(c(mu=0.8, half=0.1, yield= 0.1, vol=1000, flow=50, sub_in=1))
model$setVars(c(bac=0.1, sub=0.5))
monod <- function(c,h) {c/(c+h)}
print(model$stoichiometry(box=NULL))
print(model$stoichiometry(box=c(1)))
```

---

stoiCreate

*Stoichiometry Matrix from Reaction Equations*

---

**Description**

Creates a stoichiometry matrix from a set of reaction equations.

**Usage**

```
stoiCreate(
  reactions,
  eval = FALSE,
  env = globalenv(),
  toRight = "_forward",
  toLeft = "_backward"
)
```

**Arguments**

- |           |  |
|-----------|--|
| reactions | A named vector of character strings, each representing a (chemical) reaction. See syntax details below.  |
| eval      | Logical. If FALSE (default), the returned matrix is of type <a href="#">character</a> and any mathematical expressions are returned as text. If TRUE, an attempt is made to return a <a href="#">numeric</a> matrix by evaluating the expression making use env. |
| env       | Only relevant if eval is TRUE. Must be an environment or list supplying constants, functions, and operators needed to evaluate expressions in the generated matrix.  |

|         |  |
|---------|--|
| toRight | Only relevant for reversible reactions. The passed character string is appended to the name of the respective element of reactions to create a unique name for the forward reaction. |
| toLeft  | Like toRight, but this is the suffix for the backward reaction.  |

### Value

A matrix with the following properties:

- The number of columns equals the total number of components present in reactions. The components' names are used as column names.
- The number of rows equals the length of reactions plus the number of reversible reactions. Thus, a single row is created for each non-reversible reaction but two rows are created for reversible ones. The latter represent the forward and backward reaction (in that order). The row names are constructed from the names of reactions, making use of the suffixes toRight and toLeft in the case of reversible reactions.
- The matrix is filled with the stoichiometric factors extracted from reactions. Empty elements are set to zero.
- The type of the matrix ([character](#) or [numeric](#)) depends on the value of eval.

### Note

The syntax rules for reaction equations are as follows (see examples):

- There must be a left hand side and a right hand side. Sides must be separated by one of the arrows '>', '<', or '<->' with the latter indicating a reversible reaction.
- Names of component(s) must appear at each side of the reaction. These must be legal row/column names in R. If multiple components are consumed or produced, they must be separated by '+'.  
Example:  $\text{H}_2 + \text{O}_2 \rightarrow \text{H}_2\text{O}$
- Any stoichiometric factors need to appear before the respective component name using '\*' as the separating character. Stoichiometric factors being equal to unity can be omitted.  
Example:  $2\text{H}_2 + \text{O}_2 \rightarrow \text{H}_2\text{O}$
- A stoichiometric factor is treated as a mathematical expression. In common cases, it is just a numeric constant. However, the expression can also involve references to variables or functions. If such an expression contains math operators '\*' or '+' it needs to be enclosed in parenthesis.  
Example:  $\text{H}_2 + \text{O}_2 \rightarrow (\text{H}_2 + \text{O}_2)$

### Author(s)

David Kneis <david.kneis@tu-dresden.de>

### See Also

Use [stoiCheck](#) to validate the mass balance of the generated matrix.

## Examples

```
# EXAMPLE 1: From https://en.wikipedia.org/wiki/Petersen_matrix (July 2016)
#
reactions <- c(
  formS= "A + 2 * B -> S",
  equiES= "E + S <-> ES",
  decoES= "ES -> E + P"
)
stoi <- stoiCreate(reactions, eval=TRUE, toRight="_f", toLeft="_b")
print(stoi)

# EXAMPLE 2: Decomposition of organic matter (selected equations only)
#
# Eq. 1 and 2 are from Soetaert et al. (1996), Geochimica et Cosmochimica
# Acta, 60 (6), 1019-1040. 'OM' is organic matter. Constants 'nc' and 'pc'
# represent the nitrogen/carbon and phosphorus/carbon ratio, respectively.
reactions <- c(
  oxicDegrad= "OM + O2 -> CO2 + nc * NH3 + pc * H3PO4 + H2O",
  denitrific= "OM + 0.8*HN03 -> CO2 + nc*NH3 + 0.4*N2 + pc*H3PO4 + 1.4*H2O",
  dissPhosp1= "H3PO4 <-> H + H2PO4",
  dissPhosp2= "H2PO4 <-> H + HPO4"
)
# Non-evaluated matrix
stoi <- stoiCreate(reactions, toRight="_f", toLeft="_b")
print(stoi)
# Evaluated matrix ('nc' and 'pc' according to Redfield ratio)
pars <- list(nc=16/106, pc=1/106)
stoi <- stoiCreate(reactions, eval=TRUE, env=pars, toRight="_f", toLeft="_b")
print(stoi)
```

---

vars

*Declaration of Variables*

---

## Description

Declaration of variables of the bacteria growth example model.

## Format

A data frame with the following fields:

- name : Name of the variable
- units : Unit
- description : Short description (text)

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