

# Package ‘SBMLR’

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**Title** SBML-R Interface and Analysis Tools

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**Description** This package contains a systems biology markup language (SBML) interface to R.

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**Depends** XML, deSolve

**Suggests** rsbml

**License** GPL-2

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## Contents

|                         |   |
|-------------------------|---|
| Ops.SBMLR . . . . .     | 2 |
| readSBML . . . . .      | 3 |
| readSBMLR . . . . .     | 3 |
| S4toS3 . . . . .        | 4 |
| saveSBML . . . . .      | 5 |
| saveSBMLR . . . . .     | 6 |
| sim . . . . .           | 7 |
| summary.SBMLR . . . . . | 9 |

|              |           |
|--------------|-----------|
| <b>Index</b> | <b>11</b> |
|--------------|-----------|

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Ops.SBMLR

*Check the equality of the species and reactions of two SBMLR models*

---

### Description

This function tests the equivalence of two models with respect to the species and reaction data frames generated by summary.

### Usage

```
## S3 method for class 'SBMLR'  
Ops(e1,e2)
```

### Arguments

e1                    The first of the two model objects of class SBMLR which are to be compared.  
e2                    The second model object.

### Value

A list containing the following two boolean dataframes

species              The equality of species information tabularized as a data frame.  
reactions            The equality of reaction information tabularized as a dataframe.

### Author(s)

Tom Radivoyevitch

### See Also

[summary.SBMLR](#)

### Examples

```
library(SBMLR)  
curto1=readSBMLR(file.path(system.file(package="SBMLR"), "models/curto.r"))  
curto2=readSBML(file.path(system.file(package="SBMLR"), "models/curto.xml"))  
curto1==curto2
```

---

|          |   |
|----------|---|
| readSBML | <i>Convert an SBML file into an R object of class SBMLR</i> |
|----------|---|

---

**Description**

This function converts an SBML level 2 file into a corresponding R model structure of class SBMLR.

**Usage**

```
readSBML(filename)
```

**Arguments**

filename            An SBML level 2 model input file.

**Details**

A limited subset of SBML level 2 models is currently supported, e.g. events and function definitions are not covered.

**Value**

A corresponding SBMLR model object in R.

**Author(s)**

Tom Radivoyevitch

**See Also**

[readSBMLR](#)

**Examples**

```
library(SBMLR)
curtoX=readSBML(file.path(system.file(package="SBMLR"), "models/curto.xml"))
curtoR=readSBMLR(file.path(system.file(package="SBMLR"), "models/curto.r"))
curtoX==curtoR
```

---

|           |  |
|-----------|--|
| readSBMLR | <i>Convert an SBMLR file into an R model object of class SBMLR</i> |
|-----------|--|

---

**Description**

This function converts an SBMLR model file into a corresponding SBMLR model object. This is more than a source-ing: the file is simpler than the object since things are generated, such as, rate law and rule R expressions and functions, and mathML.

**Usage**

```
readSBMLR(filename)
```

**Arguments**

filename            An SBMLR model definition file.

**Details**

A limited subset of SBML level 2 models is currently supported, e.g. events and function definitions are not covered.

**Value**

A corresponding SBMLR model object.

**Note**

This function replaces the use of `source` in older versions of SBMLR. It converts rate law and rule strings to R functions and expressions and to MathML.

**Author(s)**

Tom Radivoyevitch

**See Also**

[readSBML](#)

**Examples**

```
library(SBMLR)
curtoX=readSBML(file.path(system.file(package="SBMLR"), "models/curto.xml"))
curtoR=readSBMLR(file.path(system.file(package="SBMLR"), "models/curto.r"))
curtoX==curtoR
```

---

S4toS3

*Converts an S4 class SBML object created by `rsbml` into an S3 object of class `SBMLR`*

---

**Description**

This function provides a path from `rsbml` to `SBMLR`. The latter, being `S3`, is less cluttered with empty fields/slots than the former. The advantage of the `S4` object is that it comes from more robust SBML reading: `rsbml` uses `libsbml` to parse SBML, `SBMLR` uses the R package `XML`. NOTE: As `rsbml` is no longer supported on the MAC, this function no longer works on the Mac.

**Usage**

```
S4toS3(dom)
```

**Arguments**

dom                An S4 DOM object of class `SBML` produced by `rsbml`.

**Details**

Carried over are compartments, species, global parameters, rules and reactions.

**Value**

A corresponding SBMLR model object, i.e. an S3 list of lists type of object.

**Author(s)**

Tom Radivoyevitch

**Examples**

```
## Not run:
library(rsbml)
(dom <- rsbml_read(file.path(system.file(package="SBMLR"), "models/sod.xml")))
library(SBMLR)
(mod=S4toS3(dom))
summary(mod)

## End(Not run)
```

---

saveSBML

*Saves an SBMLR object to an SBML file.*

---

**Description**

This function converts a class SBMLR model object into an SBML level 2 version 1 file.

**Usage**

```
saveSBML(model, filename)
```

**Arguments**

|          |                            |
|----------|----------------------------|
| model    | The S3 SBMLR model object. |
| filename | The name of the SBML file  |

**Details**

The output file is SBML level 2.

**Value**

No value returned.

**Warning**

SBML events and function definitions are NOT implemented.

**Note**

The SBML file is written incrementally, rather than first built as a DOM in R and then saved using `xmlSave`.

**Author(s)**

Tom Radivoyevitch

**References**

Radivoyevitch, T. A two-way interface between limited Systems Biology Markup Language and R. *BMC Bioinformatics* 5, 190 (2004).

**See Also**

[saveSBMLR](#)

**Examples**

```
library(SBMLR)
curtoR=readSBMLR(file.path(system.file(package="SBMLR"), "models/curto.r"))
saveSBML(curtoR,"curtoR.xml")
curtoX=readSBML("curtoR.xml")
curtoX==curtoR
summary(curtoR)
unlink("curtoR.xml")
```

---

saveSBMLR

*Save an R model object of class SBMLR to a file.*

---

**Description**

This function converts an SBMLR model object in R into an SBMLR model definition file. Rate laws are provided only in string form. Redundancy is eliminated to make the file easier to edit.

**Usage**

```
saveSBMLR(model, filename)
```

**Arguments**

`model`            The SBMLR model object to be mapped into the SBMLR model definition file.  
`filename`        The file name of the destination SBMLR model definition file.

**Value**

No value returned.

**Warning**

SBML events and function definitions are NOT implemented.

**Note**

Similar to saveSBML, the file is written incrementally.

**Author(s)**

Tom Radivoyevitch

**References**

Radivoyevitch, T. A two-way interface between limited Systems Biology Markup Language and R. BMC Bioinformatics 5, 190 (2004).

**See Also**

[saveSBML](#)

**Examples**

```
library(SBMLR)
curto=readSBMLR(file.path(system.file(package="SBMLR"), "models/curto.r"))
saveSBMLR(curto,"curtoR.r")
curtoR=readSBMLR("curtoR.r")
curto==curtoR
summary(curtoR)
unlink("curtoR.r")
```

---

sim

*Simulate a model of S3 class SBMLR*

---

**Description**

This function simulates a model given report times and optional modulators. It uses lsoda of the deSolve package.

**Usage**

```
sim(model, times, modulator=NULL,X0=NULL, ...)
```

**Arguments**

|           |   |
|-----------|---|
| model     | The S3 model object to be simulated. Initial conditions are passed through this object.   |
| times     | The sequence of time points to be sampled and provided as rows of the output matrix.  |
| modulator | Null if there are no modulators (default), a vector of numbers if there are steady state Vmax modulators, and a list of interpolating functions if there are time course Vmax modulators. |
| X0        | Override model initial conditions in simulations, particularly piece-wise perturbation simulations.   |
| ...       | To pass extra args such as event data frames to deSolve.  |

**Details**

This is a wrapper for ode.

**Value**

The data frame output that comes out of ode.

**Note**

Rules are implemented through time varying boundary conditions updated at each time point as a side effect within the (now internal) function `fderiv`.

**Author(s)**

Tom Radivoyevitch

**References**

For the folate cycle example given below: Morrison PF, Allegra CJ: Folate cycle kinetics in human breast cancer cells. *JBiolChem* 1989, 264(18):10552-10566.

**Examples**

```
##---- The following perturbs PRPP from 5 to 50 uM in Curto et al.'s model.
library(SBMLR)
curto=readSBML(file.path(system.file(package="SBMLR"), "models/curto.xml"))
(dPRPP10 <- data.frame(var = "PRPP", time = 0, value = 10,method = "mult"))
(out=sim(curto,times=seq(-20,70,1),events = list(data = dPRPP10) ) )
plot(out,which=c("PRPP","den","IMP","HX","Gua","aprt","XMP","Xa","UA"))

# which should be the same plots as
curto=readSBMLR(file.path(system.file(package="SBMLR"), "models/curto.r"))
(dPRPP10 <- data.frame(var = "PRPP", time = 0, value = 10,method = "mult"))
(out=sim(curto,times=seq(-20,70,1),events = list(data = dPRPP10) ) )
plot(out,which=c("PRPP","den","IMP","HX","Gua","aprt","XMP","Xa","UA"))

##---- The following generates Morrison's folate system response to 1uM MTX
morr=readSBMLR(file.path(system.file(package="SBMLR"), "models/morrison.r"))
out1=sim(morr,seq(-20,0,1))
morr$species$EMTX$ic=1
out2=sim(morr,0:30)
outs=data.frame(rbind(out1,out2))
attach(outs)
par(mfrow=c(3,4))
plot(time,FH2b,type="l",xlab="Hours")
plot(time,FH2f,type="l",xlab="Hours")
plot(time,DHFRf,type="l",xlab="Hours")
plot(time,DHFRtot,type="l",xlab="Hours")
plot(time,CHOFH4,type="l",xlab="Hours")
plot(time,FH4,type="l",xlab="Hours")
plot(time,CH2FH4,type="l",xlab="Hours")
plot(time,CH3FH4,type="l",xlab="Hours")
plot(time,AICARsyn,type="l",xlab="Hours")
plot(time,MTR,type="l",xlab="Hours")
plot(time,TYMS,type="l",xlab="Hours")
```



```
#plot(time,EMTX,type="l",xlab="Hours")
plot(time,DHFReductase,type="l",xlab="Hours")
par(mfrow=c(1,1))
detach(outs)
morr$species$EMTX$ic=0

## Note: This does not work, since EMTX is not a state variable, it is a bc
##(dEMTX1 <- data.frame(var = "EMTX", time = 0, value = 1,method = "add"))
##(out=simulate(morr,times=seq(-20,30,1),events = list(data = dEMTX1) ) )
```

summary.SBMLR

*Get summary information from an SBMLR model***Description**

This function extracts information from a model of class SBMLR and returns it as a list. The list includes species and reaction information tabularized as data frames.

**Usage**

```
## S3 method for class 'SBMLR'
summary(object,...)
```

**Arguments**

|        |  |
|--------|--|
| object | A model object of class SBMLR from which information is to be extracted. |
| ...    | For compatibility with summary of the base package.                      |

**Details**

no details

**Value**

A list containing the following elements

|            |   |
|------------|---|
| BC         | A logical vector indicating which species are not state variables, i.e. which species are boundary conditions or auxillary variables.   |
| y0         | The initial state (boundary conditions excluded!).  |
| nStates    | The length of the state vector, i.e. the number of system states.   |
| S0         | The full set of species initial values.   |
| nReactions | The number of reactions.  |
| nSpecies   | The number of species, including states, boundary conditions and possibly auxillary variables such as the total concentration of dihydrofolate reductase in the morrison.r model.   |
| incid      | The incidence/stoichiometry matrix. This usually contains ones and minus ones corresponding to fluxes either synthesizing or degrading (respectively) a state variable chemical species. This matrix multiplied by the flux vector on its right yields the corresponding concentration state variable time derivatives. |

|           |  |
|-----------|--|
| species   | Species information (i.e. names, ICs, BCs, and compartments) as a data frame.              |
| reactions | Reaction information tabularized as a dataframe, including string laws and initial fluxes. |

**Note**

The list output can be attached to immediately define many model variables of interest.

**Author(s)**

Tom Radivoyevitch

**Examples**

```
library(SBMLR)
curto=readSBMLR(file.path(system.file(package="SBMLR"), "models/curto.r"))
summary(curto)
```

# Index

## \* **arith**

- readSBML, 3
- readSBMLR, 3
- S4toS3, 4
- saveSBML, 5
- saveSBMLR, 6

## \* **math**

- Ops . SBMLR, 2
- readSBML, 3
- readSBMLR, 3
- S4toS3, 4
- saveSBML, 5
- saveSBMLR, 6
- sim, 7
- summary . SBMLR, 9

Ops . SBMLR, 2

readSBML, 3, 4  
readSBMLR, 3, 3

S4toS3, 4  
saveSBML, 5, 7  
saveSBMLR, 6, 6  
sim, 7  
summary (summary . SBMLR), 9  
summary . SBMLR, 2, 9