

SimBiology

For Use with MATLAB®

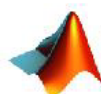
■ Computation

■ Visualization

■ Programming

Reference

Version 2



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SimBiology Reference

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Functions — By Category

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Object Constructors (p. 1-5)	MATLAB functions that create SimBiology objects
Units (p. 1-6)	Functions for unit conversion and creating user defined units

Tools

<code>sbioconsmoiety</code>	Find conserved moieties in SimBiology model
<code>sbiodesktop</code>	Open SimBiology modeling and simulation GUI
<code>sbioensembleplot</code>	Show results of ensemble run using 2-D or 3-D plots
<code>sbioensimblerun</code>	Multiple stochastic ensemble runs of SimBiology model
<code>sbioensemblestats</code>	Get statistics from ensemble data created using <code>sbioensimblerun</code>
<code>sbiogetmodel</code>	Get model object that generated simulation data
<code>sbiogetnamedstate</code>	Get state and time data from simulation results
<code>sbiogetsensmatrix</code>	3-D sensitivity matrix from simulation results
<code>sbiohelp</code>	Help for SimBiology functions
<code>sbiolasterror</code>	SimBiology last error message
<code>sbiolastwarning</code>	SimBiology last warning message
<code>sbioparamestim</code>	Perform parameter estimation
<code>sbioreset</code>	Delete all model and simulation objects
<code>sbioselect</code>	Search for objects with specified constraints
<code>sbiosimulate</code>	Simulate model object

Projects

<code>sbioaddtolibrary</code>	Add abstract kinetic law to user-defined library
<code>sbicopylibrary</code>	Copy library to disk
<code>sbioloadproject</code>	Load project from file
<code>sbioremovefromlibrary</code>	Remove abstract kinetic law or unit from user-defined library
<code>sbiosaveproject</code>	Save all models in root object
<code>sbiowhos</code>	Show contents of project file, library file or SimBiology root object

SBML Models

<code>sbmlexport</code>	Export SimBiology model to SBML file
<code>sbmlimport</code>	Import SBML formatted file

Object Constructors

<code>sbioabstractkineticlaw</code>	Construct abstract kinetic law object
<code>sbiomodel</code>	Construct model object
<code>sbioparameter</code>	Construct parameter object
<code>sbioreaction</code>	Construct reaction object
<code>sbioroot</code>	Return SimBiology root object
<code>sbiorule</code>	Construct rule object
<code>sbiospecies</code>	Construct species object

Units

<code>sbiococonvertunits</code>	Convert unit and unit value to new unit
<code>sbioregisterunit</code>	Create user-defined unit
<code>sbioregisterunitprefix</code>	Create user-defined unit prefix
<code>sbioshowunitprefixes</code>	Information about registered unit prefixes
<code>sbioshowunits</code>	Information about registered units
<code>sbiounitcalculator</code>	Convert value between units
<code>sbiounregisterunit</code>	Remove user-defined unit from root and library
<code>sbiounregisterunitprefix</code>	Remove user-defined unit prefix from root and library

Functions — Alphabetical List

sbioabstractkineticlaw

Purpose	Construct abstract kinetic law object
Syntax	<pre>abstkineticlawObj = sbioabstractkineticlaw('Name') abstkineticlawObj = sbioabstractkineticlaw('Name', 'Expression') abstkineticlawObj = sbioabstractkineticlaw(...'PropertyName', PropertyNameValue...)</pre>
Arguments	
<i>Name</i>	Enter a name for the abstract kinetic law. Name must be unique in the user-defined kinetic law library. Name is referenced by <i>kineticlawObj</i> .
<i>Expression</i>	The mathematical expression that defines the kinetic law.
Description	<p>A SimBiology abstract kinetic law defines a reaction rate expression, species variables and parameter variables for a kinetic law.</p> <p><i>abstkineticlawObj = sbioabstractkineticlaw('Name')</i> creates an abstract kinetic law object, with name <i>Name</i> and returns it to <i>abstkineticlawObj</i>.</p> <p>The <i>abstract kinetic law</i> provides a mechanism for applying a specific rate law to multiple reactions. It acts as a mapping template for the reaction rate. The abstract kinetic law defines a reaction rate expression, which is shown in the property <i>Expression</i>, and the species and parameter variables used in the expression. The species variables are defined in the <i>SpeciesVariables</i> property, and the parameter variables are defined in the <i>ParameterVariables</i> property of the abstract kinetic law object.</p> <p>In order to use <i>abstkineticlawObj</i> when constructing a kinetic law object with the method <i>addkineticlaw</i>, <i>abstkineticlawObj</i> must be added to the user-defined library with the <i>sbioaddtolibrary</i> function. To get the abstract kinetic law objects in the user-defined library, use the command <i>get(sbioroot, 'UserDefinedKineticLaws')</i></p>

`abstkineticlawObj = sbioabstractkineticlaw('Name', 'Expression')`, constructs a SimBiology abstract kinetic law object, `abstkineticlawObj` with name, `Name` and with expression, `'Expression'` and returns it to `abstkineticlawObj`.

`abstkineticlawObj = sbioabstractkineticlaw(...'PropertyName', PropertyValue...)` defines optional properties. The property name/property value pairs can be in any format supported by the function set (for example, name-value string pairs, structures, and name-value cell array pairs).

Additional `abstkineticlawObj` properties can be viewed with the `get` command. `abstkineticlawObj` properties can be modified with the `set` command.

Method Summary

<code>copyobj (any object)</code>	Copy SimBiology object and its children
<code>delete (any object)</code>	Delete SimBiology object
<code>display (any object)</code>	Display summary of SimBiology object

Property Summary

<code>Annotation</code>	Property with information about a SimBiology object
<code>Expression</code>	Property containing the expression used to determine the reaction rate equation
<code>Name</code>	Property with name of object
<code>Notes</code>	Property with HTML text describing SimBiology object
<code>ParameterVariables</code>	Property showing parameters in abstract kinetic law

sbioabstractkineticlaw

Parent	Property indicating the parent object
SpeciesVariables	Property showing species in abstract kinetic law
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Example

- 1 Create an abstract kinetic law.

```
abstkineticlawObj = sbioabstractkineticlaw('mylaw1', '(k1*s)/(k2+k1*s)');
```

- 2 Assign the parameter and species variables in the expression

```
set (abstkineticlawObj, 'SpeciesVariables', {'s'});
set (abstkineticlawObj, 'ParameterVariables', {'k1', 'k2'});
```

- 3 Add the new abstract kinetic law to the user-defined library.

```
sbioaddtolibrary(abstkineticlawObj);
```

SimBiology adds the abstract kinetic law to the user-defined library.
You can verify this using sbiowhos.

```
sbiowhos -kineticlaw -userdefined
```

```
Abstract Kinetic Law Object Array
```

Index:	Library:	Name:	Expression:
1	UserDefined	mylaw1	(k1*s)/(k2+k1*s)

- 4 Use the new abstract kinetic law when defining a reaction's kinetic law.

```
modelObj = sbiomodel('cell');
reactionObj = addreaction(modelObj, 'A + B <-> B + C');
kineticlawObj = addkineticlaw(reactionObj, 'mylaw1');
```

Remember to specify the `SpeciesVariableNames` and the `ParameterVariableNames` in the `kineticlawObj` to fully define the `ReactionRate` of the reaction.

See Also

`sbiomodel`, `addreaction`, `addkineticlaw`, `addparameter`

sbioaddtolibrary

Purpose Add abstract kinetic law to user-defined library

Syntax `sbioaddtolibrary (abstkineticlawObj)`

Description `sbioaddtolibrary (abstkineticlawObj)` adds the abstract kinetic law object (`abstkineticlawObj`) to the user-defined library. `abstkineticlawObj` is added to the SimBiology root object's `UserDefinedKineticLaws` list. `abstkineticlawObj` is available automatically in future MATLAB sessions. You can use the abstract kinetic law objects in the built-in and user-defined library to construct a kinetic law object with the method `addkineticlaw`.

To get the abstract kinetic law objects in the built-in and user-defined libraries, use the commands `get(sbioroot, 'BuiltInKineticLaws')`, `get(sbioroot, 'UserDefinedKineticLaws')`.

To remove an abstract kinetic law from the user-defined library, use the method `sbioremovefromlibrary`. You will not be able to remove an abstract kinetic law object being used by a kinetic law object.

Example This example shows how to create an abstract kinetic law and add it to the user-defined library.

1 Create an abstract kinetic law.

```
abstkineticlawObj = sbioabstractkineticlaw('mylaw1', '(k1*s)/(k2+k1*s)');
```

2 Assign the parameter and species variables in the expression.

```
set (abstkineticlawObj, 'SpeciesVariables', {'s'});
set (abstkineticlawObj, 'ParameterVariables', {'k1', 'k2'});
```

3 Add the new abstract kinetic law to the user-defined library.

```
sbioaddtolibrary(abstkineticlawObj);
```

SimBiology adds the abstract kinetic law to the user-defined library. You can verify this using `sbiowhos`.

```
sbiowhos -kineticlaw -userdefined
```

Abstract Kinetic Law Object Array

Index:	Library:	Name:	Expression:
1	UserDefined	mylaw1	(k1*s)/(k2+k1+s)

- 4** Use the new abstract kinetic law when defining a reaction's kinetic law.

```
modelObj = sbiomodel('cell');
reactionObj = addreaction(modelObj, 'A + B <-> B + C');
kineticlawObj = addkineticlaw(reactionObj, 'mylaw1');
```

Remember to specify the `SpeciesVariableNames` and the `ParameterVariableNames` in the `KineticLawObj` to fully define the `ReactionRate` of the reaction.

See Also

`addkineticlaw`, `sbioabstractkineticlaw`, `sbioregisterunit`,
`sbioregisterunitprefix`, `sbioroot`

sbioconsmoiet

Purpose Find conserved moieties in SimBiology model

Syntax

```
[G, Sp]=  
sbioconsmoiet(modelObj)  
[G, Sp] = sbioconsmoiet(modelObj, alg)  
H = sbioconsmoiet(modelObj, alg, 'p')  
H = sbioconsmoiet(modelObj, alg, 'p', FormatArg)  
[SI,SD,LO,NR,ND] = sbioconsmoiet(modelObj, 'link')
```

Arguments

<i>G</i>	An m -by- n matrix, where m is the number of conserved quantities found and n is the number of species in the model. Each row of <i>G</i> specifies a linear combination of species whose rate of change over time is zero.
<i>Sp</i>	Cell array of species names that labels the columns of <i>G</i> .
<i>modelObj</i>	Model object to be evaluated for conserved moieties.
<i>alg</i>	Specify algorithm to use during evaluation of conserved moieties. Valid values are 'qr' , 'rreduce' , or 'semipos' .
<i>H</i>	Cell array of strings containing the conserved moieties.
<i>p</i>	Prints the output to a cell array of strings.
<i>FormatArg</i>	Specifies formatting for the output <i>H</i> . <i>FormatArg</i> should either be a C-style format string, or a positive integer specifying the maximum number of digits of precision used.

<i>interpolation</i>	String variable denoting the interpolation scheme to be used if data should be interpolated to get a consistent time vector. Valid values are 'linear' (linear interpolation), 'zoh' (zero-order hold), or 'off' (no interpolation). Default is 'off'. If interpolation is on, the data is interpolated to match the time vector with the smallest simulation stop time.
<i>SI</i>	Cell array containing the names of independent species in the model.
<i>SD</i>	Cell array containing the names of dependent species in the model.
<i>L0</i>	Link matrix relating <i>SI</i> and <i>SD</i> . The link matrix <i>L0</i> satisfies <i>ND</i> = <i>L0</i> * <i>NR</i> . For the 'link' functionality, species with their BoundaryCondition or ConstantAmount properties set to true are treated as having stoichiometry of zero in all reactions.
<i>NR</i>	Reduced stoichiometry matrices containing one row for each independent species. The concatenated matrix [<i>NR</i> ; <i>ND</i>] is a row-permuted version of the full stoichiometry matrix of <i>modelObj</i> .
<i>ND</i>	Reduced stoichiometry matrices containing one row for each dependent species. The concatenated matrix [<i>NR</i> ; <i>ND</i>] is a row-permuted version of the full stoichiometry matrix of <i>modelObj</i> .

Description

[*G*, *Sp*] = sbioconsmoiety(*modelObj*) calculates a complete set of linear conservation relations for the species in the SimBiology model object *modelObj*.

sbioconsmoiety computes conservation relations by analyzing the structure of the model object's stoichiometry matrix. Thus, **sbioconsmoiety** does not include species that are governed by algebraic or rate rules.

`[G, Sp] = sbioconsmoiety(modelObj, alg)` provides an algorithm specification. For `alg` specify 'qr', 'rreduce', or 'semipos'.

When you specify 'qr', `sbioconsmoiety` uses an algorithm based on QR factorization. From a numerical standpoint, this is the most efficient and reliable approach.

When you specify 'rreduce', `sbioconsmoiety` uses an algorithm based on row reduction, which yields better numbers for smaller models. This is the default.

When you specify 'semipos', `sbioconsmoiety` returns conservation relations in which all the coefficients are greater than or equal to 0, permitting a more transparent interpretation in terms of physical quantities.

For larger models, the QR-based method is recommended. For smaller models, row reduction or the semipositive algorithm may be preferable. For row reduction and QR factorization, the number of conservation relations returned equals the row rank degeneracy of the model object's stoichiometry matrix. The semipositive algorithm may return a different number of relations. Mathematically speaking, this algorithm returns a generating set of vectors for the space of semipositive conservation relations.

`H = sbioconsmoiety(modelObj, alg, 'p')` returns a cell array of strings `H` containing the conserved quantities in `modelObj`.

`H = sbioconsmoiety(modelObj, alg, 'p', FormatArg)` specifies formatting for the output `H`. `FormatArg` should either be a C-style format string, or a positive integer specifying the maximum number of digits of precision used.

`[SI, SD, LO, NR, ND] = sbioconsmoiety(modelObj, 'link')` uses a QR-based algorithm to compute information relevant to the dimensional reduction, via conservation relations, of the reaction network in `modelObj`.

Examples

Example 1

Shows conserved moieties in a cycle.

- 1** Create a model that is a cycle. For convenience use arbitrary reaction rates, as this will not affect the result.

```
m = sbiomodel('cycle');
m.addreaction('a -> b','ReactionRate','1');
m.addreaction('b -> c','ReactionRate','b');
m.addreaction('c -> a','ReactionRate','2*c');
```

- 2** Look for conserved moieties.

```
[g sp] = sbioconsmoiety(m)
```

```
g =
```

```
1      1      1
```

```
sp =
```

```
'a'
'b'
'c'
```

Example 2

Explore semipositive conservation relations in the oscillator model.

```
m = sbmlimport('oscillator');
sbioconsmoiety(m,'semipos','p')

ans =
'pol + pol_OpA + pol_OpB + pol_OpC'
'OpB + pol_OpB + pA_OpB1 + pA_OpB_pA + pA_OpB2'
'OpA + pol_OpA + pC_OpA1 + pC_OpA2 + pC_OpA_pC'
'OpC + pol_OpC + pB_OpC1 + pB_OpC2 + pB_OpC_pB'
```

sbioconsmoiety

See Also

Moiety Conservation in the User's Guide, SimBiology method
`getstoichmatrix`

Purpose

Convert unit and unit value to new unit

Syntax

```
sbioconvertunits(Obj,  
'Unit')
```

Description

`sbioconvertunits(Obj, 'Unit')` converts the current *Units property on SimBiology object, *Obj* to the unit, *Unit*. This will configure the *Units property to *Unit* and update the corresponding value property. For example `sbioconvertunits` on a speciesObj updates the InitialAmount property value and the InitialAmountUnits property value.

Obj can be an array of SimBiology objects. *Obj* must be a SimBiology object that contains a unit property. The SimBiology objects that contain a unit property are SimBiology species and parameter objects. For example, if *Obj* is a species object with InitialAmount configured to 1 and InitialAmountUnits configured to mole, after the call to `sbioconvertunits` with *Unit* specified as molecule, speciesObj InitialAmount is 6.0221e23 and InitialAmountUnits is molecule.

Example

Convert the units of the initial amount of glucose from molecule to mole.

- 1 Create the species 'glucose' and assign an initial amount of 23 molecule.

At the command prompt type,

```
speciesObj = sbiospecies ('glucose', 23, 'InitialAmountUnits', 'molecule')
```

Species Object Array

Index:	Name:	InitialAmount:	InitialAmountUnits:
1	glucose	23	molecule

- 2 Convert the InitialAmountUnits of glucose from molecule to mole.

sbioconvertunits

```
sbioconvertunits (speciesObj, 'mole')
```

3 Verify the conversion of units and InitialAmount value.

Units are converted from molecule to mole.

```
get (speciesObj2, 'InitialAmountUnits')
```

```
ans =
```

```
mole
```

InitialAmount value is changed.

```
get (speciesObj2, 'InitialAmount')
```

```
ans =
```

```
3.8192e-023
```

See Also

[sbioshowunits](#)

Purpose	Copy library to disk
Syntax	<pre>sbiocopylibrary ('kineticlaw','LibraryFileName') sbiocopylibrary ('unit','LibraryFileName')</pre>
Description	<p><code>sbiocopylibrary</code> copies all user-defined abstract kinetic laws to a file.<code>sbiocopylibrary ('kineticlaw','LibraryFileName')</code> copies all user-defined abstract kinetic laws to the file <code>LibraryFileName.sbklib</code>.</p> <p><code>sbiocopylibrary ('unit','LibraryFileName')</code> copies all user-defined units and unit-prefixes to the file <code>LibraryFileName.sbulib</code>.</p> <p>To get the abstract kinetic law objects in the built-in and user-defined libraries, use the commands <code>get(sbioroot, 'BuiltInKineticLaws')</code>, <code>get(sbioroot, 'UserDefinedKineticLaws')</code>. To add an abstract kinetic law to the user-defined library, use the method <code>sbioaddtolibrary</code>.</p> <p>To add a unit to the user-defined library, use the <code>sbioregisterunit</code> function. To add a unit prefix to the user-defined library, use the <code>sbioregisterunitprefix</code> function.</p>
Example	Create an abstract kinetic law, add it to the user-defined library and then copy the user-defined kinetic law library to a .sbklib file.
	<ol style="list-style-type: none">1 Create an abstract kinetic law.<pre>abstkineticlawObj = sbioabstractkineticlaw('mylaw1', '(k1*s)/(k2+k1*s)');</pre>2 Add the new abstract kinetic law to the user-defined library.<pre>sbioaddtolibrary(abstkineticlawObj);</pre> <p>SimBiology adds the abstract kinetic law to the user-defined library. You can verify this using <code>sbiowhos</code>.</p> <pre>sbiowhos -kineticlaw -userdefined</pre>

sbiocopylibrary

Abstract Kinetic Law Object Array

Index:	Library:	Name:	Expression:
1	UserDefined	mylaw1	(k1*s)/(k2+k1*s)

- 3** Copy the user-defined kinetic law library.

```
sbiocopylibrary kineticLaw myLibFile
```

- 4** Verify with sbiowhos.

```
sbiowhos -kineticlaw myLibFile
```

See Also

`sbioaddtolibrary`, `sbioabstractkineticlaw`, `sbioregisterunit`,
`sbioregisterunitprefix`, `sbioremovefromlibrary`

Purpose

Open SimBiology modeling and simulation GUI

Syntax

```
sbidesktop  
sbidesktop(modelObj)
```

Arguments

modelObj

Model object or an array of model objects. Enter the variable name for a top-level SimBiology model object. If you enter an array of model objects, the SimBiology desktop opens with each model object in a separate model session.

Description

`sbidesktop` opens the SimBiology GUI. The SimBiology GUI enables you to:

- Build a SimBiology model using reaction pathways and enter kinetic data for the reactions.
- Import or export SimBiology models to and from the MATLAB workspace or from a Systems Biology Markup Language (SBML) file.
- Modify an existing SimBiology model.
- Simulate a SimBiology model.
- View results from the simulation.
- Create and/or modify user-defined units and unit prefixes.
- Create and/or modify user-defined abstract kinetic law objects.

`sbidesktop(modelObj)` opens the SimBiology GUI with a top-level SimBiology model object (*modelObj*). A top-level SimBiology model object has its property `Parent` set to the SimBiology root object.

In contrast, a SimBiology model object that has its property `Parent` set to another SimBiology model is a submodel and is not stored directly by the SimBiology root. You cannot load a submodel without loading the parent model.

sbidesktop

Example

Create a SimBiology model in the MATLAB workspace, and then open the GUI with the model.

```
modelObj = sbiomodel('cell');  
sbidesktop(modelObj)
```

See Also

[sbioroot](#)

Purpose

Show results of ensemble run using 2-D or 3-D plots

Syntax

```
sbioensembleplot(simdata, species_array)
sbioensembleplot(simdata, species_array, time)
FH = sbioensembleplot(simdata, species_array)
FH = sbioensembleplot(simdata, species_array, time)
```

Arguments

<i>simdata</i>	<i>simdata</i> is a cell array of time series objects, where each time series object holds data for a separate simulation run. The <i>simdata</i> object could change in future versions. Use the <i>simdata</i> object to call only <i>sbioensemblestats</i> and <i>sbioensembleplot</i> .
<i>species_array</i>	Either an array of species objects or a cell array of names of species in the model object that was used to create <i>simdata</i> . If there are species with duplicate names in different submodels, either use fully qualified names or use an array of species objects to identify species correctly. For example, a fully qualified name for a species named sp1 that is in a submodel named sub2, that is inside a submodel sub1, that is parented to the top-level model top1, is top1.sub1.sub2.sp1.
<i>time</i>	Either a single scalar value or a 2-element numeric vector. When it is a single scalar value, it is expected that the data in <i>simdata</i> is interpolated to a common time vector and that the time value in <i>time</i> is a member of the common time vector within a numerical tolerance of 1.0e-4. If <i>time</i> is not found in the common time vector, you see an error. When <i>time</i> is a numeric vector, it must have two elements: [<i>time_lower</i> , <i>time_upper</i>], where <i>time_lower</i> and <i>time_upper</i> are real numbers such that <i>time_lower</i> <= <i>time_upper</i> . The distribution plot are created for data points whose time value lies in the closed interval [<i>time_lower</i> , <i>time_upper</i>]. You can

sbioensembleplot

specify the 2-element numeric vector version of *time* regardless of whether *simdata* is interpolated to a common time vector or not.

FH array of handles to figure windows

Description

`sbioensembleplot(simdata, species_array)` shows a 3-D shaded plot of time-varying distribution of one or more species specified in *species_array*.

`sbioensembleplot(simdata, species_array, time)` shows a 2-D distribution plot of the species specified in *species_array* at time *time*.

FH = `sbioensembleplot(simdata, species_array)` or *FH* = `sbioensembleplot(simdata, species_array, time)` returns an array of handles to the figure windows in *FH*.

Examples

This example shows you how to plot data from an ensemble run without interpolation.

- 1 Load a SimBiology model *m1* from a SimBiology project file.

```
sbioloadproject('radiodecay.sbproj','m1');
```

- 2 Change the solver of the active configuration set to be `.ssa`. Also, adjust the `LogDecimation` property on the `SolverOptions` property of the configuration set to reduce the size of the data generated.

```
cs = getconfigset(m1, 'active');
set(cs, 'SolverType', 'ssa');
so = get(cs, 'SolverOptions');
set(so, 'LogDecimation', 10);
```

- 3 Perform an ensemble of 20 runs with no interpolation.

```
simdata = sbioensemblерun(m1, 20);
```

- 4** Create a 2-D distribution plot of the second species at a time value in the range [1.0 1.05]. Note that since simdata was not interpolated, you should specify a time range here.

```
sbioensembleplot(simdata, m1.species(2), [1.0 1.05] );
```

- 5** Create a 3-D shaded plot of both species.

```
sbioensembleplot(simdata, {'x','z'});
```

See Also

SimBiology functions `sbioensemblerrun`, `sbioensemblestats`, `sbiomodel`

sbioensblerun

Purpose Multiple stochastic ensemble runs of SimBiology model

Syntax `simdata = sbioensblerun(modelObj, numruns, interpolation)`

Arguments

<code>simdata</code>	<code>simdata</code> is a cell array of time series objects, where each time series object holds data for a separate simulation run. The <code>simdata</code> object could change in future versions. Use the <code>simdata</code> object to call only <code>sbioensblestats</code> and <code>sbioensbleplot</code> .
<code>modelObj</code>	Model object to be simulated.
<code>numruns</code>	Integer scalar representing the number of stochastic runs to make.
<code>interpolation</code>	String variable denoting the interpolation scheme to be used if data should be interpolated to get a consistent time vector. Valid values are ' <code>linear</code> ' (linear interpolation), ' <code>zoh</code> ' (zero-order hold), or ' <code>off</code> ' (no interpolation). Default is ' <code>off</code> ' . If interpolation is on, the data is interpolated to match the time vector with the smallest simulation stop time.

Description

`simdata = sbioensblerun(modelObj, numruns, interpolation)`
performs a stochastic ensemble run of the SimBiology model object (`modelObj`) . `modelObj` must be a top-level SimBiology model. A top-level SimBiology model object has its `Parent` property set to the SimBiology root object. The active configset on `modelObj` is used for performing the multiple simulation runs. The `SolverType` property of the active configset must be set to one of the stochastic solvers: '`ssa`', '`expltau`', or '`impltau`' . `sbioensblerun` generates an error if the `SolverType` property of the active configset of `modelObj` is set to any of the deterministic (ODE) solvers.

Examples

This example shows you how to perform an ensemble run and generate a 2D distribution plot.

- 1 Load a SimBiology model `m1` from a SimBiology project file.

```
sbioloadproject('radiodecay.sbproj','m1');
```

- 2 Change the solver of the active configset to be `ssa`. Also, adjust the `LogDecimation` property on the `SolverOptions` property of the configuration set.

```
cs = getconfigset(m1, 'active');
set(cs, 'SolverType', 'ssa');
so = get(cs, 'SolverOptions');
set(so, 'LogDecimation', 10);
```

- 3 Perform an ensemble of 20 runs with linear interpolation to get a consistent time vector for all time series objects.

```
simdata = sbioensblerun(m1, 20, 'linear');
```

- 4 Create a 2D distribution plot of the second species at a time value in the range [1.0 1.05].

```
sbioensembleplot(simdata, m1.species(2), [1.0 1.05] );
```

See Also

SimBiology functions `addconfigset`, `getconfigset`,
`sbioensemblestats`, `sbioensembleplot`, `setactiveconfigset`

sbioensemblestats

Purpose Get statistics from ensemble data created using sbioensemblerun

Syntax

```
[t,m] = sbioensemblestats(simdata)
[t,m,v] = sbioensemblestats(simdata)
[t,m,v,n] = sbioensemblestats(simdata)
[t,m] = sbioensemblestats(simdata, species_array),
[t,m,v] = sbioensemblestats(simdata, species_array)
[t,m,v,n] = sbioensemblestats(simdata, species_array)
[t,m] = sbioensemblestats(simdata, species_array, interpolation),
[t,m,v] = sbioensemblestats(simdata, species_array, interpolation)
[t,m,v,n] = sbioensemblestats(simdata, species_array, interpolation)
```

Arguments

<i>t</i>	Vector of doubles that holds the common time vector after interpolation
<i>m</i>	Matrix of mean values from the ensemble data. The number of rows in <i>m</i> is the length of the common time vector <i>t</i> after interpolation and the number of columns is equal to the number of species. The species order corresponding to the columns of <i>m</i> can be obtained from any of the time series objects in <i>simdata</i> using <i>sbiogetnamedstate</i> .
<i>simdata</i>	<i>simdata</i> is a cell array of time series objects, where each time series object holds data for a separate simulation run. The <i>simdata</i> object could change in future versions. Use the <i>simdata</i> object to call only <i>sbioensemblestats</i> and <i>sbioensembleplot</i> .
<i>v</i>	Matrix of variance obtained from the ensemble data. <i>v</i> has the same dimensions as <i>m</i>

<i>n</i>	Cell array of strings that holds names of species whose mean and variance are returned in <i>m</i> and <i>v</i> , respectively. The number of elements in <i>n</i> is the same as the number of columns of <i>m</i> and <i>v</i> . The order of species names in <i>n</i> corresponds to the species order of columns of <i>m</i> and <i>v</i> .
<i>species_array</i>	Either an array of species objects or a cell array of names of species in the model object that was used to create <i>simdata</i> . If there are species with duplicate names in different submodels, either use fully qualified names or use an array of species objects to identify species correctly. For example, a fully qualified name for a species named sp1 that is in a submodel named sub2, that is inside a submodel sub1, that is parented to the top-level model top1, is top1.sub1.sub2.sp1.
<i>interpolation</i>	String variable denoting the interpolation method to be used if data is to be interpolated to get a consistent time vector. Valid values are 'linear' (linear interpolation), 'zoh' (zero-order hold), or 'off' (no interpolation). Default is 'off'. If interpolation is on, the data is interpolated to match the time vector with the smallest simulation stop time.

Description

$[t, m] = \text{sbioensemblestats}(\text{simdata})$ gets mean information *m* as a function of time for all the species in the model used to generate ensemble data *simdata* by running `sbioensemblerrun`. The species order corresponding to the columns of *m* can be obtained from any of the time series objects in *simdata* by using `sbiogetnamedstate`.

$[t, m, v] = \text{sbioensemblestats}(\text{simdata})$ gets mean *m* and variance *v* as a function of time for all the species in the model used to generate ensemble data *simdata* by running `sbioensemblerrun`.

sbioensemblestats

`[t,m,v,n] = sbioensemblestats(simdata)` gets mean m , variance v , and names n of species corresponding to the columns of m and v .

`[t,m] = sbioensemblestats(simdata, species_array)`, or `[t,m,v]` = `sbioensemblestats(simdata, species_array)`, or `[t,m,v,n] = sbioensemblestats(simdata, species_array)` gets mean m , variance v , and names n only for the species specified in `species_array`.

`[t,m] = sbioensemblestats(simdata, species_array, interpolation)`, or `[t,m,v]` = `sbioensemblestats(simdata, species_array, interpolation)`, or `[t,m,v,n] = sbioensemblestats(simdata, species_array, interpolation)` gets mean m , variance v , and names n only for the species specified in `species_array` using the interpolation method `interpolation`.

Examples

Shows you how to get ensemble run statistics with the default interpolation method.

- 1 Load a SimBiology model `m1` from a SimBiology project file.

```
sbioloadproject('radiodecay.sbproj','m1');
```

- 2 Change the solver of the active configuration set to be `ssa`. Also, adjust the `LogDecimation` property on the `SolverOptions` property of the configuration set.

```
cs = getconfigset(m1, 'active');
set(cs, 'SolverType', 'ssa');
so = get(cs, 'SolverOptions');
set(so, 'LogDecimation', 10);
```

- 3 Perform an ensemble of 20 runs with no interpolation.

```
simdata = sbioensemblerun(m1, 20);
```

- 4 Get ensemble statistics for all species using the default interpolation method.

```
[T,M,V] = sbioensemblestats(simdata);
```

- 5** Get ensemble statistics for a specific species using the default interpolation scheme.

```
[T2,M2,V2] = sbioensemblestats(simdata, {'z'});
```

See Also

[sbioensemblерun](#), [sbioensembleplot](#), [sbiomodel](#), [sbiogetnamedstate](#)

sbiogetmodel

Purpose Get model object that generated simulation data

Syntax `modelObj = sbiogetmodel(timeseriesObj)`

Arguments

`timeseriesObj` Time series object returned by the function `sbiosimulate`.

`modelObj` Model object associated with the time series object.

Description

`modelObj = sbiogetmodel(timeseriesObj)` returns the SimBiology model (`modelObj`) associated with the results from a simulation run (`timeserieObj`). You can use this function to find the model object associated with the specified time series object when you load a project with several model objects and time series objects.

If the SimBiology model used to generate the time series object (`timeseriesObj`) is not currently loaded, `modelObj` is empty.

Example

Retrieve the model object that generated the time series object.

- 1 Create a model object, simulate, and then return the results as a time series object.

```
modelObj = sbmlimport('oscillator');
timeseriesObj = sbiosimulate(modelObj);
```

- 2 Get the model that generated the simulation results.

```
modelObj2 = sbiogetmodel(timeseriesObj)
```

SimBiology Model - Oscillator

Model Components:

Models: 0

```
Parameters:      0
Reactions:      42
Rules:          0
Species:        23
```

3 Check that the two models are the same.

```
modelObj == modelObj2  
ans =  
1
```

See Also

[sbiosimulate](#)

sbiogetnamedstate

Purpose Get state and time data from simulation results

Syntax

```
[t,x]= sbiogetnamedstate(timeseriesObj)
[t,x]= sbiogetnamedstate(timeseriesObj, 'SpeciesName')
[t,x,SpeciesName]= sbiogetnamedstate(...)
```

Description `sbiogetnamedstate` returns state and time data from simulation results. `[t,x]= sbiogetnamedstate(timeseriesObj)` returns the time and state data associated with the simulation run results (`timeseriesObj`) and returns to `t` and `x` respectively. `timeseriesObj` is a time series object returned by the `sbiosimulate` function.

- `t` is a n -by-1 vector, where n is the number of times the reactions fired. `t` defines the time steps of the firing of the reactions.
- `x` is a n -by- m matrix where n is the number of times the reactions fired and m is the number of SimBiology species in the SimBiology model. Each column of `x` defines the variation in the quantity of a species over time.

`[t,x]= sbiogetnamedstate(timeseriesObj, 'SpeciesName')` returns the state data associated with the SimBiology species of name `SpeciesName` from the simulation run results, (`timeseriesObj`) and returns it to `x`. `SpeciesName` can be a cell array of SimBiology species names. If a species with name, `SpeciesName`, does not exist, SimBiology returns a warning. `SpeciesName` must be the fully qualified species name, for example `ModelName.SpeciesName`. If the species is in a submodel, `SpeciesName` must be `ModelName.SubmodelName.SpeciesName`.

`[t,x,SpeciesName]= sbiogetnamedstate(...)` returns the names of the species associated with each column of `x` to `SpeciesName`.

Example This example shows a) how to get the data associated with a specified species and, b) how to get the data and associated names for each state (`x`)

- 1 Import the file for theoscillator model.

```
modelObj = sbmlimport('oscillator.xml');

2 Simulate modelObj.

timeseriesObj = sbiosimulate(modelObj);

3 Get the data for the species namedpol.

[t1, data1] = sbiogetnamedstate(timeseriesObj, 'pol');

4 Get the data and associated names for each x.

[t2, data2, names] = sbiogetnamedstate(timeseriesObj);

5 Plot the results of the simulation.

plot(t3, data3);
legend(names);
```

See Also

[sbiosimulate](#)

sbiogetsensmatrix

Purpose 3-D sensitivity matrix from simulation results

Syntax

```
[T,R,States,Inpfacs] =  
sbiogetsensmatrix(tsObj)  
[T,R,States,Inpfacs] =  
sbiogetsensmatrix(tsObj, StateNames, InpFacNames)
```

Arguments

<i>T</i>	Column vector of length m specifying time points for the sensitivity data in <i>R</i> .
<i>R</i>	m -by- n -by- p array of sensitivity data with times, species states, and input factors labeling its first, second, and third dimensions respectively.
<i>States</i>	Contains names of the species states that label the second dimension of <i>R</i> . $R(:,i,j)$ is the time course for the sensitivity of species <i>States</i> { <i>i</i> } to the input factor <i>Inpfacs</i> { <i>j</i> }.
<i>Inpfacs</i>	Contains names of the input factors that label the third dimension of <i>R</i> . $R(:,i,j)$ is the time course for the sensitivity of species <i>States</i> { <i>i</i> } to the input factor <i>Inpfacs</i> { <i>j</i> }.
<i>tsObj</i>	Time series object returned by <i>sbiosimulate</i> . Contains sensitivity data when sensitivity analysis is enabled.

<i>StateNames</i>	Specify states to get sensitivity data from <i>tsObj</i> . Can be an empty array, or a single name, or a cell array of names. When empty array is specified, returns the sensitivity data on all species states contained in <i>tsObj</i> .
<i>InpFacNames</i>	Specify input factors to get sensitivity data from <i>tsObj</i> . Can be an empty array, or a single name, or a cell array of names. When empty array is specified, returns the sensitivity data for all input factors contained in <i>tsObj</i> .

Description

`[T,R,States,Inpfacs] = sbiogetsensmatrix(tsObj)` gets time and sensitivity data from the time series object *tsObj* generated by simulating a SimBiology model object using `sbiosimulate`. `sbiogetsensmatrix` can only return sensitivity data that is contained in *tsObj*.

The sensitivity data that is logged in *tsObj* is set at simulation time by the active configuration set that is used during the simulation. Note that the sensitivity data *R* returned by `sbiogetsensmatrix` may be normalized, as specified at simulation time.

`[T,R,States,Inpfacs] = sbiogetsensmatrix(tsObj, StateNames, InpFacNames)` gets sensitivity data for the states specified by *StateNames* and the input factors specified by *InpFacNames*.

Examples

Perform sensitivity analysis of the `radiodecay` model.

- Import the radio decay model from SimBiology demos.

```
modelObj = sbmlimport('radiodecay');
```

- Retrieve the configuration set object from `modelObj`.

```
configsetObj = getconfigset(modelObj);
```

- 3** Add a parameter to the ParameterInputFactors property and display. Use the sbioselect function to retrieve the parameter object from the model.

```
set(configsetObj.SensitivityAnalysisOptions, ...
    'ParameterInputFactors', sbioselect(modelObj, 'Type', 'parameter'));
get(configsetObj.SensitivityAnalysisOptions, 'ParameterInputFactors')
```

Parameter Object Array

Index:	Name:	Value:	ValueUnits:
1	c	0.5	1/second

- 4** Add species to the SpeciesInputFactors property and display. Use the sbioselect function to retrieve the species objects from the model.

```
set(configsetObj.SensitivityAnalysisOptions, ...
    'SpeciesInputFactors', modelObj.Species)
get (configsetObj.SensitivityAnalysisOptions, 'SpeciesInputFactors')
```

Species Object Array

Index:	Name:	InitialAmount:	InitialAmountUnits:
1	x	1000	molecule
2	z	0	molecule

- 5** Enable SensitivityAnalysis.

```
set(configsetObj.SolverOptions, 'SensitivityAnalysis', true)
get(configsetObj.SolverOptions, 'SensitivityAnalysis')
```

ans =

1

- 6** Simulate and return the results to a time series object.

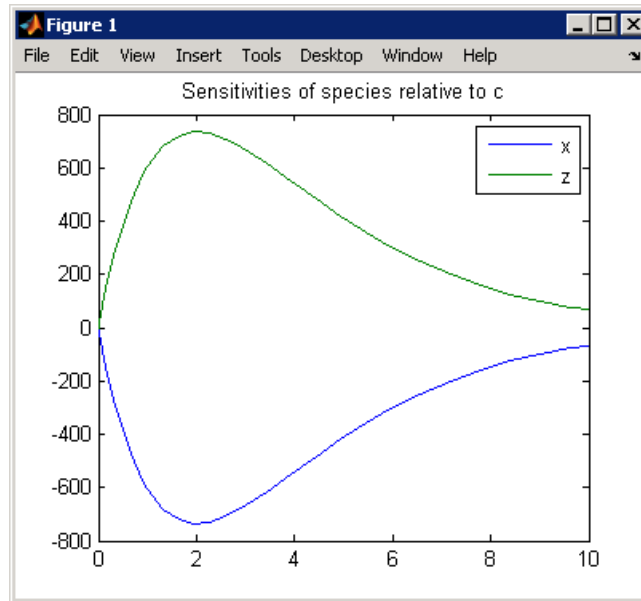
```
tobj = sbiosimulate(modelObj);
```

7 Extract sensitivity data from the time series object.

```
[t R snames ifacs] = sbiogetsensmatrix(tobj);
```

8 Plot the extracted data.

```
plot(t, R(:,:,3));
legend(snames);
title(['Sensitivities of species relative to ' ifacs{3}]);
```



See Also

SimBiology functions `sbiogetnamedstate`, `sbiohelp`, `sbiosimulate`

Purpose	Help for SimBiology functions
Syntax	<pre>sbiohelp('FunctionName') h = sbiohelp ('FunctionName')</pre>
Description	<p><code>sbiohelp('FunctionName')</code> displays information for a SimBiology function <i>FunctionName</i>.</p> <p><code>h = sbiohelp ('FunctionName')</code> returns the help for the SimBiology function <i>FunctionName</i> to <code>h</code>.</p> <p>General information on SimBiology can be returned by specifying <i>FunctionName</i> as '<code>sbio</code>'. General information about a SimBiology object can be returned by specifying <i>FunctionName</i> as one of the following: '<code>AbstractKineticLaw</code>', '<code>KineticLaw</code>', '<code>Model</code>', '<code>Parameter</code>', '<code>Reaction</code>', '<code>Root</code>', '<code>Rule</code>', '<code>Species</code>', '<code>Configset</code>', '<code>CompileOptions</code>', '<code>ExplicitTauSolverOptions</code>', '<code>ImplicitTauSolverOptions</code>', '<code>ODESolverOptions</code>', '<code>RuntimeOptions</code>', or '<code>SSASolverOptions</code>'.</p>
Examples	<pre>sbiohelp('addreaction') sbiohelp addreaction sbiohelp reaction sbiohelp('sbioshowunits')</pre>
See Also	MATLAB function <code>help</code>

Purpose SimBiology last error message

Syntax

```
sbiolasterror
diagstruct = sbiolasterror
sbiolasterror([])
sbiolasterror(diagstruct)
```

Arguments

<i>diagstruct</i>	The diagnostic structure holding Type, Message ID and Message for the errors.
-------------------	---

Description

`sbiolasterror` or `diagstruct = sbiolasterror` return a SimBiology diagnostic structure array containing the last error(s) generated by SimBiology. The fields of the diagnostic structure are:

Type	'error'
MessageID	The message ID for the error (for example, 'SimBiology:ConfigSetNameClash')
Message	Error message

`sbiolasterror([])` resets the SimBiology last error so that it will return an empty array until the next SimBiology error is encountered.

`sbiolasterror(diagstruct)` will set the SimBiology last error(s) to those specified in the diagnostic structure (`diagstruct`).

Example

Shows you an example of using `verify` and `sbiolasterror`.

1 Import a model.

```
a = sbmlimport('radiodecay.xml')
```

SimBiology Model - RadioactiveDecay

Model Components:

Models: 0

```
Parameters:      1
Reactions:      1
Rules:          0
Species:        2
```

2 Change the ReactionRate of a reaction to make the model invalid.

```
a.reactions(1).reactionrate = 'x*y'
```

```
SimBiology Model - RadioactiveDecay
```

```
Model Components:
Models:          0
Parameters:      1
Reactions:       1
Rules:           0
Species:         2
```

3 Use the function verify to validate the model.

```
a.verify
```

```
??? Error using ==> simbio\private\odebuilder>buildPatternSubStrings
The object y does not resolve on reaction with expression 'x*y'.

Error in ==> sbiogate at 22
feval(varargin{:});

??? --> Error reported from Expression Validation :
The object 'y' in reaction 'Reaction1' does not resolve to any in-scope species
or parameters.

--> Error reported from Dimensional Analysis :
Could not resolve species, parameter or model object 'y' during dimensional analysis.

--> Error reported from ODE Compilation:
Error using ==> simbio\private\odebuilder>buildPatternSubStrings
The object y does not resolve on reaction with expression 'x*y'.
```

4 Retrieve the error diagnostic struct.

```
p = sbiolasterror

p =

1x3 struct array with fields:
    Type
    MessageID
    Message
```

5 Display the first error ID and Message.

```
p(1)

ans =

    Type: 'Error'
    MessageID: 'SimBiology:ReactionObjectDoesNotResolve'
    Message: 'The object 'y' in reaction 'Reaction1' does not
              resolve to any in-scope species or parameters.'
```

6 Reset the sbiolasterror.

```
sbiolasterror([])
```

```
ans =
```

```
[]
```

7 Set sbiolasterror to the diagnostic struct.

```
sbiolasterror(p)
```

```
ans =
```

sbiolasterror

```
1x3 struct array with fields:  
    Type  
    MessageID  
    Message
```

See Also [sbiolastwarning](#), [verify](#)

Purpose	SimBiology last warning message						
Syntax	<pre>sbiolastwarning diagstruct = sbiolastwarning sbiolastwarning([]) sbiolastwarning(diagstruct)</pre>						
Arguments	<p><i>diagstruct</i> The diagnostic structure holding Type, Message ID and Message for the warnings.</p>						
Description	<p><code>sbiolastwarning</code> or <code>diagstruct = sbiolastwarning</code> return a SimBiology diagnostic structure array containing the last warnings generated by SimBiology. The fields of the diagnostic structure are:</p> <table><tr><td>Type</td><td>'warning'</td></tr><tr><td>MessageID</td><td>The message ID for the warning (for example, 'SimBiology:DANotPerformedReactionRate')</td></tr><tr><td>Message</td><td>The warning message</td></tr></table> <p><code>sbiolastwarning([])</code> resets the SimBiology last warning so that it will return an empty array until the next SimBiology warning is encountered.</p> <p><code>sbiolastwarning(<i>diagstruct</i>)</code> will set the SimBiology last warnings to those specified in the diagnostic structure (<i>diagstruct</i>).</p>	Type	'warning'	MessageID	The message ID for the warning (for example, 'SimBiology:DANotPerformedReactionRate')	Message	The warning message
Type	'warning'						
MessageID	The message ID for the warning (for example, 'SimBiology:DANotPerformedReactionRate')						
Message	The warning message						
See Also	<code>sbiolasterror</code> , <code>verify</code>						

sbioloadproject

Purpose	Load project from file
Syntax	<pre>sbioloadproject <i>projFilename</i> sbioloadproject <i>projFilename</i> <i>variableName</i> sbioloadproject <i>projFilename</i> <i>variableName1</i> <i>variableName2</i> <i>variableName3</i></pre>
Description	<p><code>sbioloadproject <i>projFilename</i></code> loads a SimBiology project from a project file (<i>projFilename</i>). If no extension is specified SimBiology assumes a default extension of <code>.sbproj</code>.</p> <p><code>sbioloadproject <i>projFilename</i> <i>variableName</i></code> loads only the variable <i>variableName</i> from the project file.</p> <p><code>sbioloadproject <i>projFilename</i> <i>variableName1</i> <i>variableName2</i> <i>variableName3</i></code> loads only the variables <i>variableName1</i>, <i>variableName2</i>, and <i>variableName3</i> from the project. The contents of the project file can be displayed by using the <code>sbiowhos</code> command.</p>
See Also	<code>sbiosaveproject</code> , <code>sbiowhos</code> , <code>sbioaddtolibrary</code> , <code>sbioremovefromlibrary</code>

Purpose	Construct model object
Syntax	<pre>modelObj = sbiomodel('NameValue') modelObj = sbiomodel(...'PropertyName', PropertyValue...)</pre>
Arguments	
<i>NameValue</i>	Required property to specify a unique name for a model object. Enter a character string.
<i>PropertyName</i>	Property name for a Model object from the Property Summary table below.
<i>PropertyValue</i>	Property value. Valid value for the specified property.
Description	<p><code>modelObj = sbiomodel('NameValue')</code> creates a model object and returns the model object (<code>modelObj</code>). In the model object, this method assigns a value (<code>NameValue</code>) to the property Name.</p> <p><code>modelObj = sbiomodel(...'PropertyName', PropertyValue...)</code> defines optional properties. The property name, property value pairs can be in any format supported by the function set (for example, name-value string pairs, structures, and name-value cell array pairs).</p> <p>Simulate <code>modelObj</code> with the function <code>sbiosimulate</code>. Add the model as a submodel to another model object using the model object method <code>addmodel</code> or add as a submodel to a model object with the function <code>copyobj</code>. Add objects to a model object using the methods <code>addkineticlaw</code>, <code>addmodel</code>, <code>addparameter</code>, <code>addrule</code>, and <code>addspecies</code>.</p> <p>All top-level SimBiology model objects can be retrieved from the SimBiology root object. A top-level SimBiology model object has its Parent property set to the SimBiology root object. Submodels have the Parent property set to another SimBiology model and are not stored by the SimBiology root.</p>

Method Summary

addconfigset (model)	Add configuration set object to model object
addmodel (model)	Add submodel object to model object
addparameter (model, kineticlaw)	Add parameter object to model or kinetic law object
addreaction (model)	Add reaction object to model object
addrule (model)	Add rule object to model object
addspecies (model)	Add species object to model object
copyobj (any object)	Copy SimBiology object and its children
delete (any object)	Delete SimBiology object
display (any object)	Display summary of SimBiology object
getadjacencymatrix (model)	Get adjacency matrix from model object
getconfigset (model)	Get configuration set object from model object
getstoichmatrix (model)	Get stoichiometry matrix from model object
removeconfigset (model)	Remove configuration set from model
setactiveconfigset (model)	Set the active configuration set for model object
verify (model)	Validate and verify SimBiology model

Property Summary

Annotation	Property with information about a SimBiology object
Models	Property showing all model objects
Name	Property with name of object
Notes	Property with HTML text describing SimBiology object
Parameters	Property with array of parameter objects
Parent	Property indicating the parent object
Reactions	Property with an array of reaction objects
Rules	Property showing rules in model object
Species	Property showing species in model object
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Examples

- 1 Create a SimBiology model object.

```
modelObj = sbiomodel('cell', 'Tag', 'mymodel');
```

- 2 List all `modelObj` properties and the current values.

```
get(modelObj)
```

MATLAB returns

```
Annotation: ''
    Models: [0x1 double]
        Name: 'cell'
        Notes: ''
    Parameters: [0x1 double]
        Parent: [1x1 SimBiology.Root]
        Species: [0x1 double]
    Reactions: [0x1 double]
        Rules: [0x1 double]
        Tag: 'mymodel'
        Type: 'sbiomodel'
    UserData: []
```

3 Display summary of `modelObj` contents.

```
modelObj

SimBiology Model - cell

Model Components:
    Models:          0
    Parameters:      0
    Reactions:       0
    Rules:           0
    Species:         0
```

See Also

`addconfigset`, `addkineticlaw`, `addmodel`, `addparameter`,
`adddreaction`, `addrule`, `addsspecies`, `sbiорoot`, `copyobj`, `sbiosimulate`
MATLAB functions `set`, `get`

Purpose	Perform parameter estimation
Syntax	<pre>[k, result]= sbioparamestim(modelObj, tspan, xtarget) [...]= sbioparamestim(..., species_array) [...]= sbioparamestim(..., species_array, parameter_array) [...]= sbioparamestim(..., species_array, parameter_array, k0) [...]= sbioparamestim(..., species_array, parameter_array, k0, method)</pre>
Arguments	
<i>k</i>	Vector of estimated parameter values.
<i>result</i>	struct with fields that provide information about the progress of optimization.
<i>tspan</i>	n-by-1 vector representing the time span of the target data <i>xtarget</i> .
<i>xtarget</i>	n-by-m matrix, where n is the number of time samples and m is the number of states you would like to match during the simulation. States can only be species varying with time. You cannot use time varying (non-constant) parameters. The number of rows of <i>xtarget</i> must be the same as the number of rows of <i>tspan</i> .
<i>species_array</i>	Either an array of species objects or a cell array of names of species in <i>modelObj</i> whose amounts should be matched during the estimation process. The length of the <i>species_array</i> must be the same as the number of columns in <i>xtarget</i> . If there are species with duplicate names in different submodels, either use fully qualified names to identify the species correctly or use an array of species objects to identify species correctly. <i>sbioparamestim</i> assumes that order of the species in <i>species_array</i> is the same as the order used to specify columns of <i>xtarget</i> . For example, a fully qualified name for a species named sp1 that is in

sbioparamestim

a submodel named `sub2`, that is inside a submodel `sub1`, that is parented to the top level model `top1`, is `top1.sub1.sub2.sp1`.

`parameter_array` `parameter_array` is either an array of parameter objects or a cell array of names of parameters in `modelObj` whose values should be estimated. If you do not specify `parameter_array`, SimBiology estimates all the parameters in the model. When a vector of parameter initial values, (k_0), is not specified, `sbioparamestim` takes the initial values from `modelObj`. When there are parameters with duplicate names (across submodels or kinetic laws), use either parameter objects or fully qualified parameter names to identify the right parameter object. For a parameter named `param1` that is scoped at a submodel level in submodel `sub1` in the top level model `top1`, the fully qualified name is `top1.sub1.param1`. If a parameter `param2` is scoped to the kinetic law of a reaction named `reaction2` in a top level model `top2`, its fully qualified name is `top2.reaction2.param2`.

`k0` Array of doubles that holds initial values of parameters to be estimated. The length of `k0` is same as that of `parameter_array`. If left unspecified, SimBiology takes initial values for parameters from the model(`modelObj`).

method Either a string or a cell array. If it is a string, it must be the name of the optimization algorithm to be used during the estimation process. Valid values are ‘fminsearch’, ‘lsqcurvefit’ , ‘lsqnonlin’, ‘fmincon’, ‘patternsearch’ , ‘patternsearch_hybrid’. ‘ga’, or ‘ga_hybrid’.

If it is a cell array, it must have two elements: the first one is the name of the optimization method as described before and the second element is a MATLAB struct as returned by either optimset, gaoptimset, or psoptimset.

SimBiology uses the cell array option to specify user-defined optimization options. If you do not specify this argument then it defaults to ‘lsqcurvefit’ if the optimization toolbox is available; otherwise it defaults to ‘fminsearch’.

‘fminsearch’ is a part of basic MATLAB and does not require the optimization toolbox. Note that ‘fminsearch’ is an unconstrained optimization method and this could result in negative values for parameters. In that case, use another optimization method.

interpolation

Description

[*k*, *result*]= sbioparamestim(*modelObj*, *tspan*, *xtarget*) estimates parameters of the SimBiology model object (*modelObj*) to match all its states given by the target state (*xtarget*) whose time variation is given by the time span *tspan*. *modelObj* must be a top-level SimBiology model. A top-level SimBiology model object has its Parent property set to the SimBiology root object.

[...]= sbioparamestim(..., *species_array*) estimates parameters of the SimBiology model object, *modelObj* to match only some of its states given by *species_array*.

sbioparamestim

[...] = sbioparamestim(..., *species_array*, *parameter_array*) estimates only some of the parameters of the SimBiology model object *modelObj*.

[...] = sbioparamestim(..., *species_array*, *parameter_array*, *k0*) lets you specify the initial values of parameters.

[...] = sbioparamestim(..., *species_array*, *parameter_array*, *k0*, *method*) lets you specify the optimization method to use.

Examples

In all the examples below, it is assumed that you have a valid SimBiology model object (*m1*) with species named *sp1*, *sp2*, *sp3* and parameters named *p1* and *p2*.

Example 1

Given a model and some target data, estimate all its parameters without having to specify any initial values. This is the simplest case. Estimate all of its parameters, use default method

```
[k, result] = sbioparamestim(m1, tspan, xtarget);
```

Example 2

Estimate all parameters, but match only some states.

```
% Names of species to match  
sp_array = {sp1', sp2', sp3'};  
  
%Estimate all parameters  
[k, result] = sbioparamestim(m1, tspan, xtarget, sp_array);
```

Example 3

Estimate some parameters, but match all states.

```
% Names of parameters to estimate  
p_array = {p1', p2'};  
% Select all species  
sp_array = sbioselect(m1,'Type','species');  
[k, result] = sbioparamestim(m1, tspan, xtarget, sp_array, p_array);
```

Example 4

Estimate parameters specified in p_array, species specified in sp_array, use different algorithms

```
[k1,r1] = sbioparamestim(m1, tspan, xtarget, sp_array, p_array, ...
    {}, fmincon');
[k2,r2] = sbioparamestim(m1, tspan, xtarget, sp_array, p_array, ...
    {}, patternsearch');
[k3,r3] = sbioparamestim(m1, tspan, xtarget, sp_array, p_array, ...
    {}, ga');
```

Example 5

Estimate parameters specified in p_array, species specified in sp_array, change default optimization options to use user specified options

```
myopt1 = optimset(Display','iter');
[k1,r1] = sbioparamestim(m1, tspan, xtarget, ...
    sp_array, p_array, {},{fmincon', myopt1});

myopt2.Tolmesh = 1.0e-4;
[k2,r2] = sbioparamestim(m1, tspan, xtarget, ...
    sp_array, p_array, {},{patternsearch', myopt2});

myopt3.PopulationSize = 50;
myopt3.Generations = 20;
[k3,r3] = sbioparamestim(m1, tspan, xtarget, ...
    sp_array, p_array, {},{ga', myopt3});
```

See Also

- SimBiology functions-sbiomodel, sbiogetnamedstate
- MATLAB function- optimset
- Genetic Algorithm and Direct Search Toolbox function-gaoptimset, psoptimset

sbioparameter

Purpose	Construct parameter object
Syntax	<pre>parameterObj = sbioparameter(Obj, NameValue) parameterObj = sbioparameter(Obj, NameValue, ValueValue) parameterObj = sbioparameter(...'PropertyName', PropertyValue...)</pre>
Arguments	
<i>Obj</i>	Model object or kinetic law object.
<i>NameValue</i>	Property for a parameter object. Enter a unique character string. Since objects can use this property to reference a parameter, a parameter object must have a unique name at the level it is created. For example, a kinetic law object cannot contain two parameter objects named kappa. However, the model object that contains the kinetic law object can contain a parameter object named kappa along with the kinetic law object. You can use the function <code>sbioselect</code> to find an object with a specific Name property value.
<i>ValueValue</i>	Value of a parameter object. Enter a number.
Description	<p><code>parameterObj = sbioparameter(Obj, NameValue)</code> constructs a SimBiology parameter object, enters a value (<i>NameValue</i>) for the required property Name, and returns the object (<code>parameterObj</code>).</p> <p>To use a parameter object (<code>paramaterObj</code>) in a simulation, you need to add the object to a SimBiology model, or kinetic law object with the method <code>copyobj</code>. You can use the <code>addparameter</code> method to simultaneously create and assign a parameter to a model or kinetic law object. SimBiology objects are constructed with the functions <code>sbiomodel</code>, <code>addmodel</code>, <code>addkineticlaw</code>, and <code>addreaction</code></p> <p><code>parameterObj = sbioparameter(Obj, NameValue, ValueValue)</code> creates a parameter object, assigns a value (<i>NameValue</i>) to the property Name,</p>

assigns the value (*ValueValue*) to the property *Value* and returns the parameter object to a variable (*parameterObj*).

`parameterObj = sbioparameter(...'PropertyName',
 PropertyValue...)` defines optional properties. The property name/property value pairs can be in any format supported by the function set (for example, name-value string pairs, structures, and name-value cell array pairs).

Copy a SimBiology parameter object to a SimBiology model or kinetic law object with the method, `copyobj`. Remove a parameter object from a model or kinetic law object with the method, `delete`.

View additional parameter object properties with the `get` command. Modify additional parameter object properties with the `set` command. You can find help for *parameterObj* properties with the `helpPropertyName` command and help for functions with the `sbiohelpFunctionName` command.

Method Summary

<code>copyobj (any object)</code>	Copy SimBiology object and its children
<code>delete (any object)</code>	Delete SimBiology object
<code>display (any object)</code>	Display summary of SimBiology object

Property Summary

<code>Annotation</code>	Property with information about a SimBiology object
<code>ConstantValue</code>	Property to indicate variable or constant parameter value
<code>Name</code>	Property with name of object
<code>Notes</code>	Property with HTML text describing SimBiology object

sbioparameter

Parent	Property indicating the parent object
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object
Value	Property to assign value to parameter object
ValueUnits	Property with parameter value units

Examples

- 1 Construct a parameter object.

```
parameter)bj = sbioparameter('kappa', 1);
% View the help for the parameter object's Value property.
help(parameterObj, 'Value')
```

- 2 View parameter object properties.

```
get(parameterObj)
```

MATLAB returns

```
Annotation: ''
ConstantValue: 1
Name: 'kappa'
Notes: ''
Parent: [1x1 SimBiology.Reaction]
Tag: ''
Type: 'parameter'
UserData: []
Value: 4
ValueUnits: ''
```

See Also

[addparameter](#), [copyobj](#), [sbiomodel](#)

sbioreaction

Purpose Construct reaction object

Syntax

```
reactionObj = sbioreaction('ReactionValue')
reactionObj = sbioreaction('ReactantsValue',
'ProductsValue')
reactionObj = sbioreaction('ReactantsValue', RStoichCoefficients,
'ProductsValue', PStoichCoefficients)
reactionObj = sbioreaction(...'PropertyName', PropertyValue...)
```

Arguments

<i>ReactionValue</i>	Specify the reaction equation. Enter a character string. A hyphen preceded by a space and followed by a right angle bracket (->) indicate reactants going forward to products. A hyphen with left and right angle brackets (<->) indicate a reversible reaction. Coefficients before reactant or product names must be followed by a space. Examples 'A -> B', 'A + B -> C', '2 A + B -> 2 C', 'A <-> B'.
<i>ReactantsValue</i>	A string defining the species name, a cell array of strings, a species object or an array of species objects.
<i>ProductsValue</i>	A string defining the species name, a cell array of strings, a species object or an array of species objects.
<i>RStoichCoefficients</i>	Stoichiometric coefficients for reactants, length of array equal to length of <i>ReactantsValue</i> .
<i>PStoichCoefficients</i>	Stoichiometric coefficients for products, length of array equal to length of <i>ProductsValue</i> .

Description

`reactionObj = sbioreaction('ReactionValue')` creates a SimBiology reaction object, assigns a value (`ReactionValue`) to the property `Reaction`, and returns the reaction object (`reactionObj`).

To use `reactionObj` in a simulation, you must add `reactionObj` to a SimBiology model object using `copyobj`. You can use `addreaction` to simultaneously create a reaction object and add it to a model object. A SimBiology model object is constructed with the function `sbiomodel`.

`reactionObj = sbioreaction('ReactantsValue', 'ProductsValue')` constructs a SimBiology reaction object that contains reactant species (`Reactants`) and product species (`Products`). The stoichiometric values are assumed to be 1. `Reactants` and `Products` can be a string defining the species name, a cell array of strings, a species object, or an array of species objects.

`reactionObj = sbioreaction('ReactantsValue', RStoichCoefficients, 'ProductsValue', PStoichCoefficients)` adds stoichiometric coefficients (`RStoichCoefficients`) for reactant species, and stoichiometric coefficients (`PStoichCoefficients`) for product species, to the property `Stoichiometry`. The length of `Reactants` and `RCoefficients` must be equal, and the length of `Products` and `PCoefficients` must be equal.

`reactionObj = sbioreaction(...'PropertyName', PropertyValue...)` defines optional properties. The property name/property value pairs can be in any format supported by the function `set` (for example, name-value string pairs, structures, and name-value cell array pairs).

View additional `reactionObj` properties with the `get` command. Modify additional `reactionObj` properties with the `set` command. You can find help for `reactionObj` properties with the `helpPropertyName` command and help for functions with the `sbiohelp functionName` command.

A reaction object that does not have a parent can contain only species objects that do not have a parent. If a parented species object is added to an unparented reaction object, a copy of the species object will be made and added to the reaction as an unparented species.

When an unparented reaction object is added to a model, SimBiology checks the model for the required species. If the model contains the species, the reaction object now uses the model's species object. If the model does not contain the species, the species object is added to the model and the reaction object uses it.

Method Summary

addkineticlaw (reaction)	Add kinetic law object to reaction object
addproduct (reaction)	Add product species object to reaction object
addreactant (reaction)	Add species object as a reactant to reaction object
copyobj (any object)	Copy SimBiology object and its children
delete (any object)	Delete SimBiology object
display (any object)	Display summary of SimBiology object
rmproduct (reaction)	Remove species object from reaction object products
rmreactant (reaction)	Remove species object from reaction object reactants

Property Summary

Active	Property to indicate object use during a simulation
Annotation	Property with information about a SimBiology object
KineticLaw	Property showing kinetic law for ReactionRate
Name	Property with name of object

Notes	Property with HTML text describing SimBiology object
Parent	Property indicating the parent object
Products	Property to indicate reaction products
Reactants	Property to indicate reaction reactants
Reaction	Property to indicate the reaction object reaction
ReactionRate	Property containing the reaction rate equation in reaction object
Reversible	Property to indicate whether a reaction is reversible or irreversible
Stoichiometry	Property that describes species coefficients in a reaction
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Examples

1 Construct reaction objects.

```

reactionObj1 = sbioreaction('a + 3 b -> 2 c');
reactionObj2 = sbioreaction({'a', 'b'}, [1 3], 'c', 2);

% View the help for the reaction object's Reversible property.
help(robj1, 'Reversible')

```

2 View the property summary for reactionObj1.

```
get(reactionObj1)
    Active: 1
    Annotation: ''
    KineticLaw: []
        Name: ''
        Notes: ''
    Parameters: [0x1 double]
        Parent: []
    Products: [1x1 SimBiology.Species]
    Reactants: [2x1 SimBiology.Species]
    Reaction: 'a + 3 b -> 2 c'
    ReactionRate: ''
    Reversible: 0
    Stoichiometry: [-1 -3 2]
        Tag: ''
    Type: 'reaction'
    UserData: []
```

See Also

[addreaction](#), [sbiomodel](#)

Purpose	Create user-defined unit
Syntax	<pre>sbioregisterunit('Name', 'Composition', Multiplier) sbioregisterunit('Name', 'Composition', Multiplier, Offset)</pre>
Description	<p><code>sbioregisterunit('Name', 'Composition', Multiplier)</code> creates a unit with the name <i>Name</i> where the unit is defined as <i>Multiplier</i>*<i>Composition</i> and records the unit in the <code>UserDefinedUnits</code> vector of <code>sbioroot</code> and adds it to the user-defined library.</p> <p><code>sbioregisterunit('Name', 'Composition', Multiplier, Offset)</code> creates a unit with the specified offset. Available units can be listed with the <code>sbioshowunits</code> function.</p> <ul style="list-style-type: none"> • <i>Name</i> is the name of the user-defined unit. <i>Name</i> must begin with characters and can contain characters, underscores or numbers. <i>Name</i> can be any valid MATLAB variable name. • <i>Composition</i> shows the combination of base and derived units that defines the unit <i>Name</i>. For example molarity is <code>mole/liter</code>. Base units are the set of units SimBiology uses to define all unit quantity equations. Derived units are defined using base units or mixtures of base and derived units. • <i>Multiplier</i> is the numerical value that defines the relationship between the unit <i>Name</i> and the base unit as a product of the <i>Multiplier</i> and the base unit. For example 1 mole is <code>6.0221e23*molecule</code>. The <i>Multiplier</i> is <code>6.0221e23</code>. • <i>Offset</i> is the numerical value by which the unit composition is modified from the base unit. For example <code>°Celsius = (5/9) * (°Fahrenheit - 32)</code>; <i>Multiplier</i> is <code>5/9</code> and <i>Offset</i> is <code>32</code>.
Examples	<p>Example 1</p> <pre>sbioregisterunit('pint2','inch^3',28.875); sbioregisterunit('celsius2','fahrenheit',9/5,32);</pre>

sbioregisterunit

Example 2

- 1 Create units for the rate constants of a first order and a second order reaction.

```
sbioregisterunit('firstconstant', '1/second', 1);  
sbioregisterunit('secondconstant', '1/molecule*second', 1);
```

- 2 Display the unit using the command sbiowhos

```
sbiowhos -userdefined -unit
```

```
SimBiology UserDefined Units
```

Index:	Name:	Composition:	Multiplier:	Offset:
1	firstconstant	1/second	1.000000	0.000000
2	secondconstant	1/molecule*second	1.000000	0.000000

See Also

[sbioshowunits](#), [sbioregisterunitprefix](#), [sbiounregisterunit](#)

Purpose	Create user-defined unit prefix						
Syntax	<code>sbioregisterunitprefix('NameValue', Exponent)</code>						
Description	<code>sbioregisterunitprefix('NameValue', Exponent)</code> creates a unit prefix with name <i>NameValue</i> and with a multiplicative factor of 10^{Exponent} , adds it to the <code>UserDefinedUnitPrefixes</code> vector in <code>sbioroot</code> and to the user-defined library. You can see the available unit prefixes with the <code>sbioshowunitprefixes</code> function.						
Example	<p>1 Register a unit prefix.</p> <pre>sbioregisterunitprefix('peta', 15); sbiowhos -userdefined -unitprefix</pre> <p>SimBiology UserDefined Unit Prefixes</p> <table><thead><tr><th>Index:</th><th>Name:</th><th>Multiplier:</th></tr></thead><tbody><tr><td>1</td><td>peta</td><td>1.000000e+015</td></tr></tbody></table> <p>See Also</p> <p><code>sbioshowunits</code>, <code>sbioshowunitprefixes</code>, <code>sbiounregisterunit</code></p>	Index:	Name:	Multiplier:	1	peta	1.000000e+015
Index:	Name:	Multiplier:					
1	peta	1.000000e+015					

sbioremovefromlibrary

Purpose

Remove abstract kinetic law or unit from user-defined library

Syntax

```
sbioremovefromlibrary (abstkineticlawObj)  
sbioremovefromlibrary ('Type', 'Name')
```

Description

`sbioremovefromlibrary` removes an abstract kinetic law or a unit from the user-defined library. `sbioremovefromlibrary (abstkineticlawObj)` removes the abstract kinetic law object (`abstkineticlawObj`) from the user-defined library. `abstkineticlawObj` will no longer be available automatically in future MATLAB sessions.

SimBiology does not remove an abstract kinetic law that is being used in a model.

You can use a built-in or user-defined abstract kinetic law when you construct a kinetic law object with the method `addkineticlaw`.

To retrieve the abstract kinetic law objects from the built-in and user-defined libraries, use the commands `get(sbioroot, 'BuiltInKineticLaws')`, `get(sbioroot, 'UserDefinedKineticLaws')`. To add an abstract kinetic law to the user-defined library, use the method `sbioaddtolibrary`.

`sbioremovefromlibrary ('Type', 'Name')` removes the object of type 'Type' with name 'Name' from the corresponding user-defined library. Type can be 'kineticlaw', 'unit' or 'unitprefix'.

To add a unit to the user-defined library, use the `sbioregisterunit` function. To add a unit prefix to the user-defined library, use the `sbioregisterunitprefix` function.

Example

Shows you how to remove an abstract kinetic law from the user-defined library.

- 1 Create an abstract kinetic law.

```
abstkineticlawObj = sbioabstractkineticlaw('mylaw1', '(k1*s)/(k2+k1*s)');
```

- 2 Add the new abstract kinetic law to the user-defined library.

```
sbioaddtolibrary(abstkineticlawObj);
```

SimBiology adds the abstract kinetic law to the user-defined library.
You can verify this using sbiowhos.

```
sbiowhos -kineticlaw -userdefined
```

```
Abstract Kinetic Law Object Array
```

Index:	Library:	Name:	Expression:
1	UserDefined	mylaw1	(k1*s)/(k2+k1+s)

3 Remove the abstract kinetic law.

```
sbioremovefromlibrary('kineticlaw', 'mylaw1');
```

See Also

[sbioaddtolibrary](#), [sbioabstractkineticlaw](#), [sbioregisterunit](#),
[sbioregisterunitprefix](#), [sbiounregisterunit](#),
[sbiounregisterunitprefix](#)

sbioreset

Purpose Delete all model and simulation objects

Syntax `sbioreset`

Description `sbioreset` delete all SimBiology model and simulation objects at the root level. You cannot use a SimBiology model or simulation object after it is deleted. You should remove objects from the MATLAB workspace with the function `clear`.

The SimBiology root object contains a list of the top-level SimBiology model objects, available units, unit prefixes and kinetic law objects. A top-level SimBiology model object has its `Parent` property set to the SimBiology root object. A SimBiology model object that has its `Parent` property set to another SimBiology model is a submodel and is not stored by the SimBiology root.

To add an abstract kinetic law to the SimBiology root user-defined library, use the `addtolibrary` function. To add a unit to the SimBiology user-defined library on the root, use the `sbioregisterunit` function. To add a unit prefix to the SimBiology user-defined library on the root, use the `sbioregisterunitprefix` function.

Example Shows you the difference between `sbioreset` and `clear all`.

1 Import a model into the workspace.

```
modelObj = sbmlimport('oscillator');
```

Note that the workspace contains `modelObj` and if you query the SimBiology root, there is one model on the root object.

```
rootObj = sbioroot
```

SimBiology Root Contains:

Models:	1
Builtin Abstract Kinetic Laws:	3
User Abstract Kinetic Laws:	0

```
Builtin Units:          54
User Units:            0
Builtin Unit Prefixes: 13
User Unit Prefixes:    0
```

- 2** The command clear all clears the workspace, but the modelObj still exists on the rootObj.

```
clear all
```

```
rootObj
```

```
SimBiology Root Contains:
```

```
Models:                1
Builtin Abstract Kinetic Laws: 3
User Abstract Kinetic Laws: 0
Builtin Units:          54
User Units:            0
Builtin Unit Prefixes: 13
User Unit Prefixes:    0
```

- 3** The command sbioreset deletes the modelObj from the root.

```
sbioreset
rootObj
```

```
SimBiology Root Contains:
```

```
Models:                0
Builtin Abstract Kinetic Laws: 3
User Abstract Kinetic Laws: 0
Builtin Units:          54
User Units:            0
Builtin Unit Prefixes: 13
```

sbioreset

User Unit Prefixes: 0

See Also

[sbioroot](#)

Purpose Return SimBiology root object

Syntax

```
rootObj = sbioroot  
modelObj = sbioroot('modelName')
```

Arguments

<i>rootObj</i>	Return <code>sbioroot</code> to this object.
<i>modelObj</i>	Return the model with name <i>modelName</i> to this object.
<i>modelName</i>	Specify the name of the model that is on the root object.

Description

`rootObj = sbioroot` returns the SimBiology root object to `root`. The SimBiology root object contains a list of the top-level SimBiology model objects, available units, unit prefixes, and available abstract kinetic law objects.

`modelObj = sbioroot('modelName')` returns the top-level SimBiology model with name, `modelName` to `modelObj`. A top-level SimBiology model object has its `Parent` property set to the SimBiology root object. A SimBiology model object that has its `Parent` property set to another SimBiology model is a submodel and is not stored by the SimBiology root.

The units define the set of core units and user-defined units. A user-defined unit can be added with the `sbioregisterunit` function. You can remove user-defined unit with the `sbiounregisterunit` function. The unit prefixes define the set of core unit prefixes and user-defined unit prefixes.

You can add a user-defined unit prefix with the `sbioregisterunitprefix` function. Remove a user-defined unit prefix with the `sbiounregisterunitprefix` function. The abstract kinetic law objects define the core abstract kinetic law objects and user-defined abstract kinetic law objects. SimBiology uses abstract kinetic law objects when configuring a SimBiology reaction object's `KineticLaw` property with the `addkineticlaw` function.

Method Summary		
	copyobj (any object)	Copy SimBiology object and its children
	delete (any object)	Delete SimBiology object
	reset (root)	Delete all model objects from the root object
Property Summary		
	BuiltInKineticLaws	Property containing built-in kinetic laws
	BuiltInUnitPrefixes	Property containing built-in unit prefixes
	BuiltInUnits	Property containing built-in units
	Models	Property showing all model objects
	Type	Property to indicate SimBiology object type
	UserDefinedKineticLaws	Property containing user-defined kinetic laws
	UserDefinedUnitPrefixes	Property containing user-defined unit prefixes
	UserDefinedUnits	Property containing user-defined units

Examples

- 1 Get all SimBiology model objects contained by the root.

```
rootObj = sbioroot;
allmodels = get(rootObj, 'Models');
```

- 2 Get the model with name cell (if model is in root).

```
modelObj = sbioroot('cell');
```

See Also

[addkineticlaw](#), [sbiomodel](#), [sbioregisterunit](#), [sbiounregisterunit](#),
[sbioreset](#)

sbiorule

Purpose Construct rule object

Syntax *ruleObj* = sbiorule('RuleValue')

ruleObj = sbiorule(*RuleValue*,

'RuleTypeValue')

ruleObj = sbiorule(...'PropertyName', *PropertyValue*...)

Arguments

RuleValue Enter a character string within quotes. For example, enter the algebraic rule ' $V_a \cdot E_a + V_i \cdot E_i - K_2$ '.

RuleTypeValue Enter 'algebraic', 'assignment', or 'rate'. An algebraic or rate rule is evaluated at each time step during the simulation. An assignment rule is evaluated once before the simulation starts. Note: if a species or parameter is marked constant, you can still assign an initial value using an assignment rule. The amount or value gets assigned according to the rule and then remains constant during the simulation.

Description

A SimBiology rule is a mathematical expression that modifies a species amount, or a parameter value. A rule is a MATLAB expression that uses species, and parameters.

ruleObj = sbiorule('RuleValue') creates a rule object, assigns a value (*RuleValue*) to the property *Rule*, assigns the value 'algebraic' to the property *RuleType*, and assigns the root object to the property *Parent*.

To use *ruleObj* in a simulation, *ruleObj* must be added to a model object with the function *copyobj*. Note that a rule can also be added to a SimBiology model with the *addrule* function. A model object is constructed with the function *sbiomodel*.

ruleObj = sbiorule(*RuleValue*, 'RuleTypeValue') in addition to the above, this syntax enables you to specify *RuleType*.

`ruleObj = sbiorule(...'PropertyName', PropertyValue...)` defines optional properties. The property name/property value pairs can be in any format supported by the function set (for example, name-value string pairs, structures, and name-value cell array pairs).

View additional rule properties with the function `get`, and modify rule properties with the function `set`. View the rules in a model (`modelObj`) with `get(modelObj, 'Rules')`.

Method Summary

<code>copyobj (any object)</code>	Copy SimBiology object and its children
<code>delete (any object)</code>	Delete SimBiology object
<code>display (any object)</code>	Display summary of SimBiology object

Property Summary

<code>Active</code>	Property to indicate object use during a simulation
<code>Annotation</code>	Property with information about a SimBiology object
<code>Name</code>	Property with name of object
<code>Notes</code>	Property with HTML text describing SimBiology object
<code>Parent</code>	Property indicating the parent object
<code>Rule</code>	Property to define certain species and parameter interactions
<code>RuleType</code>	Property for defining the type of rule for the rule object
<code>Tag</code>	Property to specify a label for a SimBiology object

sbiorule

Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Examples

Example 1

Construct a rule object and copy to a model object.

```
robj = sbiorule('Enzt - Enzi - Enza');
modelObj = sbiomodel('cell')
robj_copy = copyobj(robj, modelObj);
```

Example 2

View the help for the rule object's RuleType property.

```
help(robj, 'RuleType')
```

Example 3

List the properties for a rule.

```
get(robj)

    Active: 1
    Annotation: ''
    Name: ''
    Notes: ''
    Parent: []
        Rule: 'myrule'
    RuleType: 'algebraic'
        Tag: ''
    Type: 'rule'
    UserData: []
```

See Also

addrule, copyobj, sbiomodel

Purpose Save all models in root object

Syntax

```
sbiosaveproject projFilename
sbiosaveproject projFilename variableName
sbiosaveproject projFilename variableName1 variableName2
variableName3
```

Description

`sbiosaveproject projFilename` saves all models in the SimBiology root object to the binary SimBiology project file named `projFilename.sbproj`. The project can be loaded with `sbioloadproject`. SimBiology returns an error if `projFilename.sbproj` is not writable.

`sbiosaveproject` creates the binary SimBiology project file named `simbiology.sbproj`. SimBiology returns an error if this is not writable.

`sbiosaveproject projFilename variableName` saves only `variableName`. `variableName` can be a SimBiology model or any MATLAB variable.

`sbiosaveproject projFilename variableName1 variableName2 variableName3` saves `variableName1, variableName2, and variableName3`.

Use the functional form of `sbiosaveproject` when the filename or variable names are stored in string. For example if the filename is stored in the variable `fileName` and you want to store MATLAB variables `variableName1` and `variableName2`, type `sbiosaveproject(fileName, 'variableName1', 'variableName2')` at the command line.

Examples

1 Import an SBML file and simulate (default configset object is used).

```
modelObj = sbmlimport ('oscillator.xml');
timeseriesObj = sbiosimulate(modelObj);
```

2 Save the model and the simulation results to a project.

```
sbiosaveproject myprojectfile modelObj timeseriesObj
```

See Also

`sbioloadproject`, `sbiowhos`, `sbioaddtolibrary`,
`sbioremovefromlibrary`

Purpose Search for objects with specified constraints

Syntax

```
Out = sbioselect(Obj, 'PropertyName', PropertyValue)
Out =
sbioselect(Obj, 'Type', 'TypeValue', 'PropertyName', PropertyValue)
Out = sbioselect(Obj,
'Where', 'PropertyName', Condition, PropertyValue)
Out = sbioselect(Obj, 'Where', 'PropertyName1', 'Condition1',
PropertyValue1,
'Where', 'PropertyName2', 'Condition2', PropertyValue2, ...)
out= sbioselect(Obj,'depth', Number,...)
```

Arguments

<i>Obj</i>	SimBiology object to search. If an object is not specified, SimBiology searches the root.
<i>PropertyName</i>	Any property of <i>Obj</i>
<i>PropertyValue</i>	Valid <i>PropertyValue</i>
<i>TypeValue</i>	Type of SimBiology object to search, for example <code>sbiomodel</code> , <code>species</code> , <code>reaction</code> , <code>kineticlaw</code>
<i>Condition</i>	Constraint to impose on the search. See table below for a list of Conditions.

Description

`sbioselect` searches for objects with specified constraints.

`Out = sbioselect(Obj, 'PropertyName', PropertyValue)` finds the objects with the property name (*PropertyName*) and property value (*PropertyValue*) contained in any SimBiology object (*Obj*) or an array of SimBiology objects. If an object is not specified, SimBiology searches the root.

`Out =`
`sbioselect(Obj, 'Type', 'TypeValue', 'PropertyName', PropertyValue)` finds the objects of Type, *TypeValue*, with the property name (*PropertyName*) and property value (*PropertyValue*) contained in any SimBiology object (*Obj*) or an array of SimBiology objects. *TypeValue* is the type of SimBiology object for example `species`, `reaction`, or `kineticlaw`.

`Out = sbioselect(Obj, 'Where', 'PropertyName', Condition, PropertyValue)` finds objects that have a property name (*PropertyName*) and value (*PropertyValue*) that matches the condition (*Condition*).

Condition	Example PropertyName	Example PropertyValue
>	InitialAmount	50
>	InitialAmount	50
==	InitialAmount	50
==i	Name	x
~=	InitialAmount	50
~=i	Name	x
>=	InitialAmount	50
<=	InitialAmount	50
between	InitialAmount	[200 300]
~between	InitialAmount	[200 300]
contains	Reactant	Species or species object array
regexp	Name	Value*
~regexp	Name	Value*
regexpi	Name	Value*
~regexpi	Name	Value*

`Out = sbioselect(Obj, 'Where', 'PropertyName1', 'Condition1', PropertyValue1, 'Where', 'PropertyName2', 'Condition2', PropertyValue2, ...)` finds the object contained by Obj that matches all the conditions specified.

Any combination of property name/property value pairs and conditions can be combined in the `sbioselect` command.

`out= sbioselect(Obj, 'depth', Number, ...)` finds objects using a model search depth of *Number*. Valid numbers are positive integer values and `inf`. If *Number* is `inf`, *Obj* and all of its children are searched. If *Number* is 1, children of *Obj* will not be searched. By default, *Number* is `inf`.

- The condition types supported for numeric properties are `>`, `<`, `=`, `~=`, `>=`, `between` and `~between`. Conditions for range are `'between'` and `'~between'`

```
PName= sbioselect (InitialAmount, 'Between', [200, 300])
```

- The condition types supported for string properties are `==`, `==i`, `~=`, `~=i`, `regexp`, `~regexp`, `regexpi` and `~regexpi`. Case conditions are `'CaseSensitive'` and `'CaseInsensitive'`. The `CaseSensitive` and `CaseInsensitive` conditional values can be used only for those properties whose values are strings. If they are used on a property whose value is not a string, it is ignored.

```
Out = sbioselect(Robj, 'Name', 'CaseSensitive', MyModel)
```

- The condition `'Contains'` can be used only for those properties whose values are an array of SimBiology objects. The value for `Contains` is one of the objects that should be in the array. `PValue` is a species or species object array

```
Out = sbioselect(reactionObj, 'Reactant', 'Contains',  
modelObj.Species)
```

- The `regexp` and `regexpi` conditional value supports any of the expressions supported by the functions `regexp` and `regexpi`(`regexp`). When a string property value is searched for without specifying a condition, it must use the same format as `get` returns. For example, if `get` returns the Name as `'MyObject'`, `sbioselect` will not find an object with a Name property value of `'myobject'`.

```
Out = sbioselect(Robj, 'Name', 'RegExp', T*)
```

- Conditions for relational operators: `>`, `<`, `=`, `~=`, `>=`, `<=`.

`PropertyName=InitialAmount`, `Condition` is `>`, and `PropertyValue=50`.

Examples

Find a SimBiology model called `cell`, in the root object.

```
sbioselect(sbioroot, 'Type', 'sbiomodel', 'Name', 'cell')
```

Find all SimBiology reactions of SimBiology model (`modelObj`), that use species A as a reactant.

```
speciesA = sbioselect(modelObj, 'Type', 'species', 'Name', ...
    'A');
out = sbioselect(modelObj, 'Type', 'reaction', 'Where', ...
    'Reactants','contains', speciesA);
```

Find all SimBiology species of SimBiology model (`modelObj`), that have an `InitialAmount` that range between 100 and 300.

```
out = sbioselect(m, 'Type', 'species', 'Where', ...
    'InitialAmount', 'between', [100 300]);
```

See Also

`regexp`

sbioshowunitprefixes

Purpose Information about registered unit prefixes

Syntax

```
Name = sbioshowunitprefixes
[Name, Multiplier] = sbioshowunitprefixes
[Name, Multiplier, Builtin]
= sbioshowunitprefixes
[Name, Multiplier, Builtin]
= sbioshowunitprefixes('Name')
```

Description *sbioshowunitprefixes* returns information about registered unit prefixes.

Name = *sbioshowunitprefixes* returns the names of the registered unit prefixes to *Name* as a cell array of strings.

[Name, Multiplier] = *sbioshowunitprefixes* returns the multiplier for each prefix in *Name* to *Multiplier*. *Multiplier* is a cell array of strings.

[Name, Multiplier, Builtin] = *sbioshowunitprefixes* returns whether the unit prefix is built-in or user-defined for each unit prefix in *Name* to *Builtin*. *Builtin* is an array of logical values. If *Builtin* is true for a unit prefix, the unit prefix is built-in. If *Builtin* is false for a unit prefix, the unit prefix is user-defined.

[Name, Multiplier, Builtin] = *sbioshowunitprefixes('Name')* returns the name, exponent and built-in status for the unit prefix with name *Name*. *Name* can be a cell array of strings.

- *Name* is the name of the prefix. Built-in prefixes are defined based on the International System of Units (SI).
- *Multiplier* shows the value of 10^{Exponent} that defines the relationship of the unit *Name* to the base unit. For example the multiplier in picomole is 10^{-12} .

Examples

```
[name, multiplier] = sbioshowunitprefixes;
[name, multiplier] = sbioshowunitprefixes('nano');
```

See Also *sbioregisterunit*, *sbioshowunits*, *sbioconvertunits*

Purpose

Information about registered units

Syntax

```
Name = sbioshowunits
[Name, Composition] =
sbioshowunits
[Name, Composition, Multiplier]
= sbioshowunits
[Name, Composition, Multiplier, Offset]
= sbioshowunits
[Name, Composition, Multiplier, Offset,
Builtin] = sbioshowunits
[Name, Composition, Multiplier, Offset,
Builtin] = sbioshowunits('Name')
```

Description

Name = *sbioshowunits* returns the names of the registered units to *Name* as a cell array of strings.

[Name, Composition] = *sbioshowunits* returns the composition for each unit in *Name* to *Composition* as a cell array of strings.

[Name, Composition, Multiplier] = *sbioshowunits* returns the multiplier for the unit with name *Name* to *Multiplier*.

[Name, Composition, Multiplier, Offset] = *sbioshowunits* returns the offset for the unit with name *Name* to *Offset*. The unit is defined as *Multiplier*Composition+Offset*.

[Name, Composition, Multiplier, Offset, Builtin] = *sbioshowunits* returns whether the unit is built-in or user-defined for each unit in *Name* to *Builtin*. *Builtin* is an array of logical values. If *Builtin* is true for a unit, the unit is built-in. If *Builtin* is false for a unit, the unit is user-defined.

[Name, Composition, Multiplier, Offset, Builtin] = *sbioshowunits('Name')* returns the name, composition, multiplier, offset and built-in status for the unit with name *Name*. *Name* can be a cell array of strings.

sbioshowunits

- *Name* is the name of the built-in or user-defined unit. *Name* must begin with characters and can contain characters, underscores or numbers.
- *Composition* shows the combination of base and derived units that defines the unit *Name*. For example molarity is mole/liter.
- *Multiplier* is the numerical value that defines the relationship between the unit *Name* and the base or derived unit as a product of the *Multiplier* and the base unit or derived unit. For example 1 mole is $6.0221e23 \text{molecule}$. The *Multiplier* is $6.0221e23$.
- *Offset* is the numerical value by which the unit composition is modified from the base unit. For example ${}^{\circ}\text{Celsius} = (5/9) * ({}^{\circ}\text{Fahrenheit} - 32)$; *Multiplier* is $5/9$ and *Offset* is 32.

Examples

```
[name, composition] = sbioshowunits;
[name, composition] = sbioshowunits('molecule');
```

See Also

[sbioregisterunit](#), [sbioshowunitprefixes](#), [sbioconvertunits](#)

Purpose

Simulate model object

Syntax

```
[T,X] = sbiosimulate(modelObj)
[T,X]
= sbiosimulate(modelObj, configsetObj)
timeseriesObj = sbiosimulate(modelObj)
```

Description

`[T,X] = sbiosimulate(modelObj)` simulates a model object (`modelObj`) using the active configuration set attached to the model (`modelObj`).

- `modelObj` — SimBiology model object. Enter the variable name for a model object.

`[T,X] = sbiosimulate(modelObj, configsetObj)` simulates a model object (`modelObj`) using a configuration set (`configset`) that overrides the active configuration set attached to the model (`modelObj`). After command is executed this override does not exist; the configuration set that is defined as 'active' is reinstated.

- `configsetObj` — A configset object stores simulation specific information. A SimBiology model can contain multiple configuration sets with only one being active at any given time. The active configuration set contains the settings that are used during the simulation.
- `T` — 1-by-n vector where n is the number of times the reactions fired. `T` defines the time steps of the firing of the reactions.
- `X` — n-by-m matrix where n is the number of times the reactions fired and m is the number of species in the model or the number of StatesToLog. Each column of `X` defines the variation in the amount of a species over time. To get the simulation values for the first species logged

```
FirstSpecies = X(:,1)
```

`timeseriesObj = sbiosimulate(modelObj)` simulates a model object (`modelObj`) using the active configuration set attached to the

model (*modelObj*) and returns the results to time series object (*timeseriesObj*).

Use *timeseriesObj*.time to access the time vector from the simulation results. Use *timeseriesObj*.data to access the state matrix from the simulation results. *timeseriesObj* also contains information about the configuration set used to simulate the model. Use *get(timeseriesObj)* to view this information.

To get the configuration sets attached to a model, use *getconfigset*. To attach a new or existing configuration set to a model, use *addconfigset*. To set the active configuration set of a model, use *setactiveconfigset*. To use command line help to get more information on these methods, for example, 'help SimBiology.Model.getconfigset'.

Property Summary	Configuration set property summary
Active	Property to indicate object use during a simulation
CompileOptions	Property holding dimensional analysis and unit conversion information
Name	Property with name of object
Notes	Property with HTML text describing SimBiology object
RuntimeOptions	Property holding options for logged species
SensitivityAnalysisOptions	Hold sensitivity analysis options
SolverOptions	Property holding the model solver options
SolverType	Property to select solver type for simulation
StopTime	Property to set the stop time for a simulation

StopTimeType	Property to specify the type of stop time for a simulation
TimeUnits	Property to show the stop time units for a simulation
Type	Property to indicate SimBiology object type

Example 1

Create a SimBiology model from an SBML file, simulate the model using a solver other than the default solver (default is `ode15s`), and view the results.

- 1** Read the file for theoscillator model.

```
modelObj = sbmlimport('oscillator.xml');
```

- 2** Get the active configset.

```
configsetObj = getconfigset(modelObj, 'active');
```

- 3** Configure the SolverType to `ode45` and set StopTime to 10.

```
set(configsetObj, 'SolverType', 'ode45');
set(configsetObj, 'StopTime', 10);
```

- 4** Simulate `modelObj`.

```
[t,x]= sbiosimulate(modelObj);
```

- 5** Plot the results of the simulation.

```
plot(t, x)
```

Example 2

Simulate the above example with DimensionalAnalysis off (set to `false`).

sbiosimulate

- 1** Repeat steps 1 and 2 above, then set dimensional analysis and unit conversion off in the configset object. DimensionalAnalysis and UnitConversion are properties of the CompileOptions object in the configset object.

```
set(configsetObj.CompileOptions, 'UnitConversion', false);  
set(configsetObj.CompileOptions, 'DimensionalAnalysis', false);
```

- 2** Simulate modelObj.

```
timeseriesObj = sbiosimulate(modelObj);
```

- 3** Plot the results of the simulation.

```
plot(timeseriesObj.Time, timeseriesObj.Data);  
legend(timeseriesObj.SpeciesNames)
```

See Also

SimBiology object constructor `sbiomodel`, model object method
`addconfigset`

Purpose	Construct species object
Syntax	<pre>speciesObj = sbiospecies('NameValue') speciesObj = sbiospecies('NameValue'),InitialAmountValue) speciesObj = sbiospecies(...'PropertyName', PropertyValue...)</pre>
Arguments	
<i>NameValue</i>	Name for a species object. Enter a character string unique to the level of object creation. Species objects are identified by Name within ReactionRate and Rule property strings. You can use the function sbioselect to find an object with a specific Name property value.
<i>InitialAmountValue</i>	Initial amount value for the species object. Enter double. Positive real number, default = 0.
Description	<p><i>speciesObj</i> = <code>sbiospecies('NameValue')</code> constructs a SimBiology.Species object, enters a value (<i>NameValue</i>) for the property Name, and returns the object (<i>speciesObj</i>).</p> <p><i>speciesObj</i> = <code>sbiospecies('NameValue'),InitialAmountValue)</code> in addition to the above, assigns an initial amount (<i>InitialAmountValue</i>) for the species.</p> <p>Species are entities that take part in reactions. A species object represents these entities. There are reserved characters you cannot use in species object name (<i>NameValue</i>)</p> <p>See “Valid Species Names” on page 4-38 for more information on species names.</p> <p>In order for a species object to be used in a simulation, the species object must be added to a SimBiology model object using <code>copyobj</code>. You can use <code>addspecies</code> to simultaneously create a species object and add it to a model object. A SimBiology model object is constructed with the function <code>sbiomodel</code>.</p>

```
speciesObj = sbiospecies(...'PropertyName', PropertyValue...)
```

defines optional properties. The property name/property value pairs can be in any format supported by the function set (for example, name-value string pairs, structures, and name-value cell array pairs).

View species object properties with the function `get`, and change properties with the function `set`. You can find help for `speciesObj` properties with the help `PropertyName` command and help for functions with the `sbiohelp FunctionName` command.

Method Summary

Methods for species objects.

<code>copyobj</code> (any object)	Copy SimBiology object and its children
<code>delete</code> (any object)	Delete SimBiology object
<code>display</code> (any object)	Display summary of SimBiology object

Property Summary

Properties for species object

<code>Annotation</code>	Property with information about a SimBiology object
<code>BoundaryCondition</code>	Property to set a species object to have a boundary condition
<code>ConstantAmount</code>	Property to specify variable or constant species amount
<code>InitialAmount</code>	Property containing initial amount of a species
<code>InitialAmountUnits</code>	Property containing units for species initial amount
<code>Name</code>	Property with name of object
<code>Notes</code>	Property with HTML text describing SimBiology object

Parent	Property indicating the parent object
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Examples

Example 1

Create a species (H_2O) and view properties for the object.

- 1 Create a species object with name H_2O and initial amount 1000.

```
speciesObj = sbiospecies('H2O', 1000);
% View the help for the species object's InitialAmount property.
help(speciesObj, 'InitialAmount')
```

- 2 View properties for the species object.

```
get(sobj)
    Annotation: ''
    ConstantAmount: 0
    InitialAmount: 1000
    InitialAmountUnits: ''
        Name: 'H2O'
        Notes: ''
        Parent: []
        Tag: ''
        Type: 'species'
    UserData: []
```

Example 2

Create two species, one is a reactant and the other is the enzyme catalyzing the reaction.

- 1 Create two species objects with the names glucose_6_phosphate and glucose_6_phosphate_dehydrogenase.

```
speciesObj1 = sbiospecies ('glucose_6_phosphate');
speciesObj2 = sbiospecies ('glucose_6_phosphate_dehydrogenase');
```

- 2 Set initial amount of glucose_6_phosphate to 100 and verify.

```
set(speciesObj1, 'InitialAmount', 100);
get(speciesObj1, 'InitialAmount')
```

MATLAB returns

```
ans =
```

```
100
```

See Also

SimBiology method `addspecies`

MATLAB functions `get` and `set`.

Purpose	Convert value between units
Syntax	<pre>result = sbiounitcalculator('fromUnits', 'toUnits', Value)</pre>
Description	<p><code>result = sbiounitcalculator('fromUnits', 'toUnits', Value)</code> converts the value, <code>Value</code> which is defined in the units, <code>fromUnits</code> to the value, <code>result</code>, which is defined in the units, <code>toUnits</code>.</p>
Example	<pre>result = sbiounitcalculator('mile/hour', 'meter/second', 1)</pre>
See Also	<code>sbioshowunits</code>

sbiounregisterunit

Purpose Remove user-defined unit from root and library

Syntax `sbiounregisterunit('Name')`

Description `sbiounregisterunit('Name')` removes the user-defined unit with the name, *Name* from the user-defined library. You cannot remove a unit from the built-in library. If *Name* is a user-defined unit, then it is removed from the `UserDefinedUnits` vector on the SimBiology root object and also from the user library. Once unregistered, this unit is not available in future MATLAB sessions. You can list the available units and find information on whether the unit is built-in or user-defined using `sbiowhos` or `sbioshowunits`.

Example 1 Create units for the rate constants of a first order and second order reactions.

```
sbioregisterunit('firstconstant', '1/second', 1);
sbioregisterunit('secondconstant', '1/molecule*second', 1);
```

2 Display the unit using the command `sbiowhos`

```
sbiowhos -userdefined -unit
```

SimBiology UserDefined Units

Index:	Name:	Composition:	Multiplier:	Offset:
1	firstconstant	1/second	1.000000	0.000000
2	secondconstant	1/molecule*second	1.000000	0.000000

3 Unregister one of the units and display the user-defined units available.

```
sbiounregisterunit('firstconstant');
sbiowhos -userdefined -unit
```

SimBiology UserDefined Units

Index:	Name:	Composition:	Multiplier:	Offset:
1	secondconstant	1/molecule*second	1.000000	0.000000

See Also

[sbioshowunits](#), [sbioregisterunit](#), [sbiounregisterunitprefix](#),
[sbioroot](#), [sbiowhos](#)

sbiounregisterunitprefix

Purpose Remove user-defined unit prefix from root and library

Syntax `sbiounregisterunitprefix('Name')`

Description `sbiounregisterunitprefix('Name')` removes the user-defined unit prefix with the name, *Name* from the user-defined library. You cannot remove a unit prefix from the built-in library. If *Name* is a user-defined unit prefix, it is removed from the `UserDefinedUnits` vector on the SimBiology root object and also from the user library. Once unregistered, this unit prefix is not available in future MATLAB sessions. You can list the available unit prefixes and find information on whether the unit prefix is built-in or user-defined using `sbiowhos` or `sbioshowunitprefixes`.

Example 1 Register a unit prefix.

```
sbioregisterunitprefix('peta', 15);
sbiowhos -userdefined -unitprefix
```

SimBiology UserDefined Unit Prefixes

Index:	Name:	Multiplier:
1	peta	1.000000e+015

2 Unregister the unit prefix.

```
sbiounregisterunitprefix('peta');
```

See Also `sbioshowunitprefixes`, `sbioshowunits`, `sbioregisterunitprefix`, `sbiounregisterunit`, `sbioroot`, `sbiowhos`

Purpose Show contents of project file, library file or SimBiology root object

Syntax

```
sbiowhos flag
sbiowhos ('flag')
sbiowhos flag1 flag2 ...
sbiowhos FileName
```

Description

sbiowhos shows contents of the SimBiology root object. This includes the built-in and user-defined abstract kinetic laws, units and unit prefixes.

sbiowhos *flag* shows specific information about the SimBiology root object as defined by *flag*. Valid flags are described in the table below:

Flag	Description
-builtin	Built-in abstract kinetic laws, units and unit prefixes.
-data	Data saved in file.
-kineticlaw	Built-in and user-defined abstract kinetic laws
-unit	Built-in and user-defined units.
-unitprefix	Built-in and user-defined unit prefixes.
-userdefined	User-defined abstract kinetic laws, units and unit prefixes.

You can also specify the functional form, sbiowhos ('*flag*')

sbiowhos *flag1 flag2 ...* shows information about the SimBiology root object as defined by *flag1*, *flag2*,...

sbiowhos *FileName* shows contents of SimBiology project or library defined by Name.

Examples

```
% Show contents of the SimBiology root object  
sbiowhos  
  
% Show abstract kinetic laws on the SimBiology root object  
sbiowhos -kineticlaw  
  
% Show the builtin units of the SimBiology root object.  
sbiowhos -builtin -unit  
  
% Show all contents of project file.  
sbiowhos myprojectfile  
  
% Show abstract kinetic laws from a library file.  
sbiowhos -kineticlaw mylibraryfile  
  
% Show all contents of multiple files.  
sbiowhos myfile1 myfile2
```

See Also

MATLAB function whos

Purpose Export SimBiology model to SBML file

Syntax

```
sbmlexport(modelObj)
sbmlexport(modelObj, 'FileName')
```

Arguments

modelObj Model object. Enter a variable name for a model object.

FileName XML file with an Systems Biology Markup Language (SBML) format. Enter either a filename or a path and filename supported by your operating system. If the filename does not have the extension .xml, then .xml is appended to end of the filename.

Description

`sbmlexport(modelObj)` exports a SimBiology model object (*modelObj*) to a file with a Systems Biology Markup Language (SBML) Level 2 format. The default file extension is .xml and the file name matches the model name.

`sbmlexport(modelObj, 'FileName')` exports a SimBiology model object (*modelObj*) to an SBML file named *FileName*. The default file extension is .xml.

A SimBiology model can also be written to a SimBiology project with the `sbiosaveproject` function to save features not supported by SBML.

Functional Characteristics and Limitations

The `sbmlexport` function,

- Exports SBML Level 2 compatible files
- Exports SBML compliant unit definitions
- Does not support submodels
- Does not export features that are not supported by SBML. These are,

sbmlexport

- Abstract kinetic law name and corresponding expression (note that sbmlexport exports the reaction rate equation)
- Configurations sets
- Active property
- UserData
- Tag

Example

Export a model (modelObj) to a file (gene_regulation.xml) in the current working directory.

```
sbmlexport(modelObj, 'gene_regulation.xml');
```

Reference

Finney, A., Hucka, M., (2003), *Systems Biology Markup Language (SBML) Level 2: Structures and facilities for model definitions*. Accessed from SBML.org

See Also

`sbmlimport`, `sbiomodel`, `sbiosaveproject`

Purpose Import SBML formatted file

Syntax `modelObj = sbmlimport('FileName')`

Arguments

FileName XML file with an Systems Biology Markup Language (SBML) format. Enter either a filename or a path and filename supported by your operating system.

Description

`modelObj = sbmlimport('FileName')` imports a SBML formatted file with name *FileName* into MATLAB and creates a model object `modelObj`. *FileName* extensions can be `.sbml` or `.xml`. The `modelObj` properties can be viewed with the `get` command. `modelObj` properties can be modified with the `set` command. At the command line, help for `modelObj` functions can be returned with the `sbiohelp` command.

Functional Characteristics and Limitations

- `sbmlimport` supports SBML Levels 1 and 2.
- If there are MATLAB incompatible variable names in any mathematical expression (for example, rules or rate expressions), SimBiology inserts brackets around those variable names in that expression.
- If the model has a single compartment, the model is read in as a top-level model. If there are multiple compartments, SimBiology returns a warning and does not read the SBML file.
- SimBiology does not support model volume and volume units, function definitions, events, and piecewise kinetics.
- SimBiology does not support brackets in species or parameter names.

Example

```
sbmlobj = sbmlimport('oscillator.xml');
```

sbmlimport

Reference

Finney, A., Hucka, M., (2003), *Systems Biology Markup Language (SBML) Level 2: Structures and facilities for model definitions*. Accessed from SBML.org.

See Also

`sbmlexport`, `sbiosimulate`
MATLAB functions `get` and `set`

Methods — By Category

Abstract Kinetic Laws (p. 3-2)	Methods for abstract kinetic law objects
Configuration Sets (p. 3-3)	Methods for configuration set objects
Kinetic Laws (p. 3-4)	Methods for kinetic law objects.
Models (p. 3-5)	Methods for the model object
Parameters (p. 3-6)	Methods for parameter objects
Reactions (p. 3-7)	Methods for reaction objects
Root (p. 3-8)	Methods for the root object
Rules (p. 3-9)	Methods for rule objects
Species (p. 3-10)	Methods for species objects
Using Object Methods (p. 3-11)	Command-line syntax for using methods with SimBiology objects

Abstract Kinetic Laws

`copyobj (any object)`

Copy SimBiology object and its children

`delete (any object)`

Delete SimBiology object

`display (any object)`

Display summary of SimBiology object

Configuration Sets

copyobj (any object)

Copy SimBiology object and its children

delete (any object)

Delete SimBiology object

display (any object)

Display summary of SimBiology object

Kinetic Laws

<code>addparameter (model, kineticlaw)</code>	Add parameter object to model or kinetic law object
<code>copyobj (any object)</code>	Copy SimBiology object and its children
<code>delete (any object)</code>	Delete SimBiology object
<code>display (any object)</code>	Display summary of SimBiology object
<code>getparameters (kineticlaw)</code>	Get specific parameters in kinetic law object
<code>getspecies (kineticlaw)</code>	Get specific species in kinetic law object
<code>setparameter (kineticlaw)</code>	Specify specific parameters in kinetic law object
<code>setspecies (kineticlaw)</code>	Specify species in kinetic law object

Models

addconfigset (model)	Add configuration set object to model object
addmodel (model)	Add submodel object to model object
addparameter (model, kineticlaw)	Add parameter object to model or kinetic law object
addraction (model)	Add reaction object to model object
addrule (model)	Add rule object to model object
addsspecies (model)	Add species object to model object
copyobj (any object)	Copy SimBiology object and its children
delete (any object)	Delete SimBiology object
display (any object)	Display summary of SimBiology object
getadjacencymatrix (model)	Get adjacency matrix from model object
getconfigset (model)	Get configuration set object from model object
getstoichmatrix (model)	Get stoichiometry matrix from model object
removeconfigset (model)	Remove configuration set from model
setactiveconfigset (model)	Set the active configuration set for model object
verify (model)	Validate and verify SimBiology model

Parameters

copyobj (any object)	Copy SimBiology object and its children
delete (any object)	Delete SimBiology object
display (any object)	Display summary of SimBiology object

Reactions

addkineticlaw (reaction)	Add kinetic law object to reaction object
addproduct (reaction)	Add product species object to reaction object
addreactant (reaction)	Add species object as a reactant to reaction object
copyobj (any object)	Copy SimBiology object and its children
delete (any object)	Delete SimBiology object
display (any object)	Display summary of SimBiology object
rmproduct (reaction)	Remove species object from reaction object products
rmreactant (reaction)	Remove species object from reaction object reactants

Root

copyobj (any object)	Copy SimBiology object and its children
delete (any object)	Delete SimBiology object
reset (root)	Delete all model objects from the root object

Rules

copyobj (any object)	Copy SimBiology object and its children
delete (any object)	Delete SimBiology object
display (any object)	Display summary of SimBiology object

Species

copyobj (any object)	Copy SimBiology object and its children
delete (any object)	Delete SimBiology object
display (any object)	Display summary of SimBiology object

Using Object Methods

Command-line syntax for using methods with SimBiology objects

Constructing (Creating) Objects
(p. 3-11)

Using Object Methods (p. 3-11)

Help for Objects, Methods and
Properties (p. 3-12)

Constructing (Creating) Objects

Create an object that is not referenced by a model using the constructor functions sbioabstractkineticlaw, sbiomodel, sbioparameter, sbioreaction, sbioroot, and sbiospecies.

```
ObjectName = ConstructorFunction(RequiredParameters,...  
                                'PropertyName', 'PropertyValue')
```

To create objects referenced by a model, use the model object methods addconfigset, addmodel, addparameter, addreaction, addrule, and addspecies.

```
ObjectName = ModelName.Method(Arguments)
```

To create objects references by a reaction, us the reaction object methods addkineticlaw, addparemeter, addproduct, and addreactant.

```
ObjectName = ReactionName.Method(Arguments)
```

Note, ObjectName is not a copy of the object but a pointer to the created object.

Using Object Methods

Using MATLAB function notation.

```
MethodName(ObjectName, arguments, ...)
```

Using object dot notation.

```
ObjectName.MethodName(arguments, ...)
```

Help for Objects, Methods and Properties

Display information for SimBiology object methods and properties in the MATLAB Command Window.

<code>help sbio</code>	Display a list of functions and methods.
<code>help FunctionName</code>	Display function information.
<code>sbiohelp('MethodName')</code>	Display method information.
<code>sbiohelp('PropertyName')</code>	Display property information.

Methods — Alphabetical List

The object that the methods apply to are listed in parenthesis after the method name.

addconfigset (model)

Purpose Add configuration set object to model object

Syntax

```
configsetObj = addconfigset(modelObj, 'NameValue')
configsetObj = addconfigset(..., 'PropertyName', PropertyValue, ...)
```

Arguments

<i>modelObj</i>	Model object. Enter a variable name.
<i>NameValue</i>	Descriptive name for a configuration set object. Reserved words 'active' and 'default' are not allowed.
<i>configsetObj</i>	Configuration set object.

Description

configsetObj = *addconfigset(modelObj, 'NameValue')* creates a configuration set object and returns to *configsetObj*.

In the configuration set object, this method assigns a value (*NameValue*) to the property Name.

configsetObj = *addconfigset(..., 'PropertyName', PropertyValue, ...)* constructs a configuration set object, *configsetObj*, and configures *configsetObj* with property value pairs. The property name/property value pairs can be in any format supported by the function set (for example, name-value string pairs, structures, and name-value cell array pairs). The *configsetObj* properties are listed below in the property summary.

A configuration set stores simulation specific information. A model object can contain multiple configuration sets, with one being active at any given time. The active configuration set contains the settings that are used during a simulation. *configsetObj* is not automatically set to active. Use the function *setactiveconfigset* to define the active configset for *modelObj*.

Use the method *copyobj* to copy a configset object and add it to the *modelObj*.

You can additionally view configuration set object properties with the command, `get` . You can modify additional configuration set object properties with the command, `set` .

Method Summary

Methods for configuration set objects

<code>copyobj (any object)</code>	Copy SimBiology object and its children
<code>delete (any object)</code>	Delete SimBiology object
<code>display (any object)</code>	Display summary of SimBiology object

Property Summary

Properties for configuration set objects

<code>Active</code>	Property to indicate object use during a simulation
<code>CompileOptions</code>	Property holding dimensional analysis and unit conversion information
<code>Name</code>	Property with name of object
<code>Notes</code>	Property with HTML text describing SimBiology object
<code>RuntimeOptions</code>	Property holding options for logged species
<code>SensitivityAnalysisOptions</code>	Hold sensitivity analysis options
<code>SolverOptions</code>	Property holding the model solver options
<code>SolverType</code>	Property to select solver type for simulation
<code>StopTime</code>	Property to set the stop time for a simulation

addconfigset (model)

StopTimeType	Property to specify the type of stop time for a simulation
TimeUnits	Property to show the stop time units for a simulation
Type	Property to indicate SimBiology object type

Examples

- 1 Create a model object by reading the file oscillator.xml and add a configuration set that simulates for 3000 seconds.

```
modelObj = sbmlimport('oscillator');
configsetObj = addconfigset(modelObj, 'myset');
```

- 2 Configure the configsetObj StopTime to 3000.

```
set(configsetObj, 'StopTime', 3000)
get(configsetObj)
```

```
Active: 0
CompileOptions: [1x1 SimBiology.CompileOptions]
    Name: 'myset'
    Notes: ''
RuntimeOptions: [1x1 SimBiology.RuntimeOptions]
SolverOptions: [1x1 SimBiology.ODESolverOptions]
    SolverType: 'ode15s'
    StopTime: 3000
StopTimeType: 'simulationTime'
    TimeUnits: 'second'
    Type: 'configset'
```

- 3 Set the new configset to be active, simulate the model using the new configset, and plot the result.

```
setactiveconfigset(modelObj, configsetObj);
```

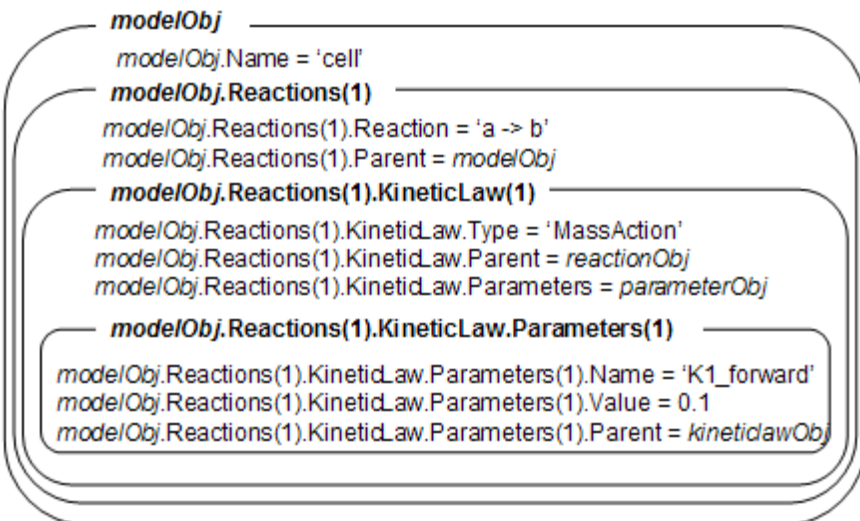
```
[t,x] = sbiosimulate(modelObj);  
plot (t,x)
```

See Also

Model object methods `getconfigset`, `removeconfigset`,
`setactiveconfigset`
MATLAB functions `get` and `set`.

addkineticlaw (reaction)

Purpose	Add kinetic law object to reaction object				
Syntax	<pre>kineticlawObj = addkineticlaw(reactionObj, 'KineticLawNameValue') kineticlawObj= addkineticlaw(..., 'PropertyName', PropertyValue, ...)</pre>				
Arguments	<table><tr><td><i>reactionObj</i></td><td>Reaction object. Enter a variable name for a reaction object.</td></tr><tr><td><i>KineticLawNameValue</i></td><td>Property to select the type of kinetic law object to create. Enter either MassAction or Heneri-Michelis-Menten.</td></tr></table>	<i>reactionObj</i>	Reaction object. Enter a variable name for a reaction object.	<i>KineticLawNameValue</i>	Property to select the type of kinetic law object to create. Enter either MassAction or Heneri-Michelis-Menten.
<i>reactionObj</i>	Reaction object. Enter a variable name for a reaction object.				
<i>KineticLawNameValue</i>	Property to select the type of kinetic law object to create. Enter either MassAction or Heneri-Michelis-Menten.				
Description	<p><i>kineticlawObj</i> = <i>addkineticlaw(reactionObj, 'KineticLawNameValue')</i> creates a kinetic law object and returns the kinetic law object (<i>kineticlawObj</i>).</p> <p>In the kinetic law object, this method assigns a name (<i>KineticLawNameValue</i>) to the property KineticLawName and assigns the reaction object to the property Parent. In the reaction object, this method assigns the kinetic law object to the property KineticLaw.</p> <pre>modelObj = sbiomodel('cell'); reactionObj = addreaction(modelObj, 'a -> b'); kineticlawObj = addkineticlaw(reactionObj, 'MassAction'); parameterObj = addparameter(kineticlawObj, 'K1_forward', 0.1); set(kineticlawObj, ParameterVariableName, 'K1_forward');</pre>				



KineticLawNameValue is any valid abstract kinetic law. See “Abstract Kinetic Law” on page 6-27 for a definition of abstract kinetic laws and more information about how they are used to get the reaction rate expression.

You can find valid *KineticLawNameValues* by querying the SimBiology root object with the commands: `get(sbioroot, 'BuiltInKineticLaws')`, and `get(sbioroot, 'UserDefinedKineticLaws').sbiowhos`. `-kineticlaw` lists `BuiltInKineticLaws` and `UserDefinedKineticLaws` in the SimBiology root. The root contains all `BuiltInKineticLaws` and all `UserDefinedKineticLaws` that are added using `sbioaddtolibrary` or `addtolibrary`.

`kineticlawObj= addkineticlaw(..., 'PropertyName', PropertyValue, ...)` constructs a kinetic law object, `kineticlawObj`, and configures `kineticlawObj` with property value pairs. The property name/property value pairs can be in any format supported by the function set (for example, name-value string pairs, structures, and name-value cell array pairs). The `kineticlawObj` properties are listed below in the property summary.

addkineticlaw (reaction)

You can view additional kinetic law object properties with the command, `get`. You can modify additional kinetic law object properties with the command, `set`. The kinetic law used to determine the `ReactionRate` of the `Reaction` can be viewed with `get(reactionObj, 'KineticLaw')`. Remove a SimBiology kinetic law object from a SimBiology reaction object with the command, `delete`.

Method Summary

Methods for kinetic law objects	
<code>addparameter (model, kineticlaw)</code>	Add parameter object to model or kinetic law object
<code>copyobj (any object)</code>	Copy SimBiology object and its children
<code>delete (any object)</code>	Delete SimBiology object
<code>display (any object)</code>	Display summary of SimBiology object
<code>getparameters (kineticlaw)</code>	Get specific parameters in kinetic law object
<code>getspecies (kineticlaw)</code>	Get specific species in kinetic law object
<code>setparameter (kineticlaw)</code>	Specify specific parameters in kinetic law object
<code>setspecies (kineticlaw)</code>	Specify species in kinetic law object

Property Summary

Properties for kinetic law objects	
<code>Annotation</code>	Property with information about a SimBiology object
<code>Expression</code>	Property containing the expression used to determine the reaction rate equation

KineticLawName	Property showing name of abstract kinetic law
Name	Property with name of object
Notes	Property with HTML text describing SimBiology object
Parameters	Property with array of parameter objects
ParameterVariableNames	Property showing cell array of reaction rate parameters
ParameterVariables	Property showing parameters in abstract kinetic law
Parent	Property indicating the parent object
SpeciesVariableNames	Property showing cell array of species used in reaction rate equation
SpeciesVariables	Property showing species in abstract kinetic law
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Example 1

This example uses the built-in kinetic law Henri-Michaelis-Menten.

- 1 Create a model object, and add a reaction object to the model.

```
modelObj = sbiomodel ('Cell');
reactionObj = addreaction (modelObj, 'Substrate -> Product');
```

addkineticlaw (reaction)

- 2** Define an abstract kinetic law for the reaction object and view the parameters to be set.

```
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');
get (kineticlawObj, 'Expression')

ans =
Vm*S/(Km + S)
```

SimBiology adds an abstract kinetic law expression to the reaction object (*reactionObj*).

The Henri-Michaelis-Menten kinetic law has two parameters (V_m and, K_m) and one species (S). You need to enter values for these parameters by first creating parameter objects, and then adding the parameter objects to the kinetic law object.

- 3** Add parameter objects to a kinetic law object. For example, create a parameter object *parameterObj1* named V_m _d, another parameter *parameterObj2* named K_m _d, and add them to a kinetic law object (*kineticlawObj*).

```
parameterObj1 = addparameter(kineticlawObj, 'Vm_d', 'Value', 6.0);
parameterObj2 = addparameter(kineticlawObj, 'Km_d', 'Value', 1.25);
```

SimBiology creates two parameter objects with concrete values that will be associated with the abstract kinetic law parameters.

- 4** Associate concrete kinetic law parameters with the abstract kinetic law parameters.

```
set(kineticlawObj, 'ParameterVariableNames', {'Vm_d' 'Km_d'});
set(kineticlawObj, 'SpeciesVariableNames', {'Substrate'});
```

SimBiology associates the concrete parameters in the property *ParameterVariableNames* with the abstract parameters in the property *ParameterVariables* using a one-to-one mapping in the order given.

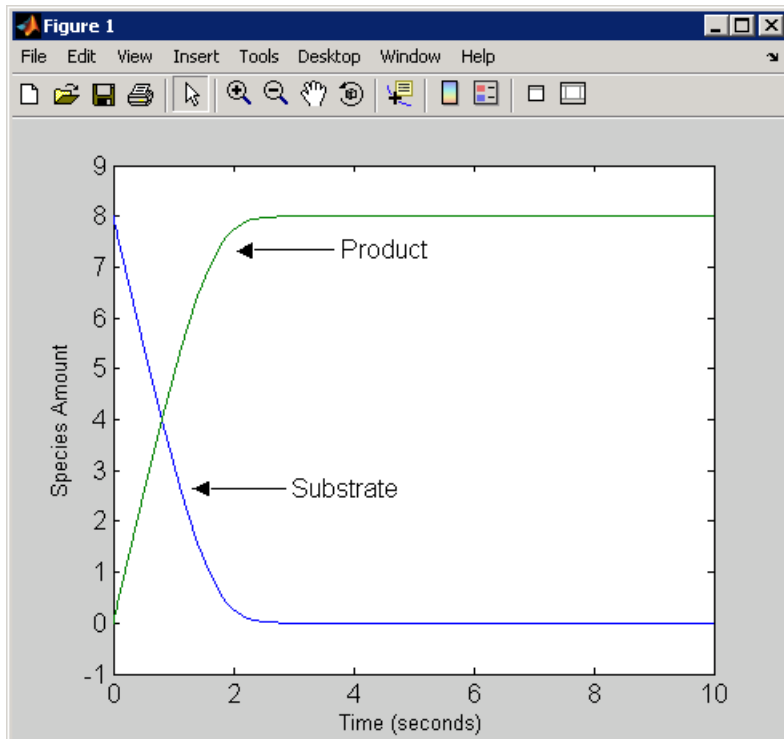
- 5 Verify that the reaction rate is expressed correctly in the reaction object ReactionRate property.

```
get (reactionObj, 'ReactionRate')
```

```
ans =  
Vm_d*Substrate/(Km_d+Substrate)
```

- 6 Enter an initial value for the substrate and simulate.

```
modelObj.Species(1).InitialAmount = 8;  
[T, X] = sbiosimulate(modelObj);  
plot(T,X)
```



addkineticlaw (reaction)

Example2

Example using the built-in kinetic law MassAction.

- 1 Create a model object, and then add a reaction object.

```
modelObj = sbiomodel ('Cell');
reactionObj = addreaction (modelObj, 'a -> b');
```

- 2 Define an abstract kinetic law for the reaction object.

```
kineticlawObj = addkineticlaw(reactionObj, 'MassAction');
get(kineticlawObj, 'Expression')

ans =
    MassAction
```

Notice, the property Expression for an abstract kinetic law with property Type set to MassAction does not show the parameters and species in the reaction rate.

- 3 Assign the rate constant for the reaction.

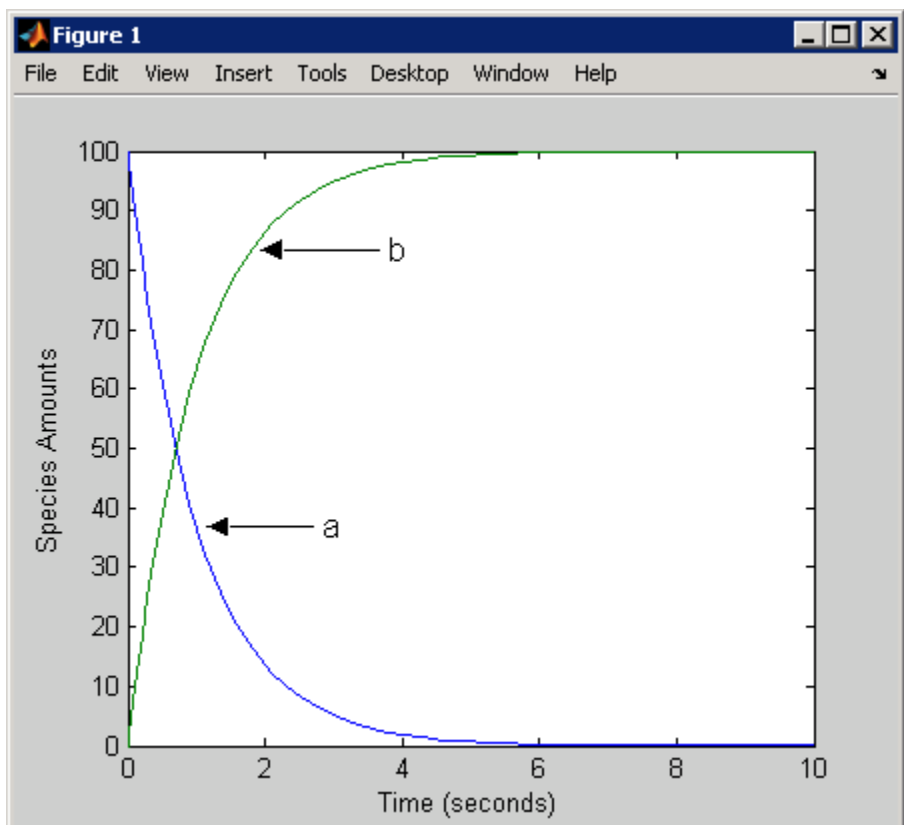
```
parameterObj = addparameter(kineticlawObj, 'k_forward');
set (kineticlawObj, 'ParameterVariablenames', 'k_forward');
get (reactionObj, 'ReactionRate')

ans =
    k_forward*a
```

- 4 Enter an initial value for the substrate and simulate.

```
modelObj.Species(1).InitialAmount = 100;
[T, X] = sbiosimulate(modelObj);
plot(T,X)
```

The value used for k_forward is default value = 1.0.



See Also

[addreaction](#), [setparameter](#)

addmodel (model)

Purpose Add submodel object to model object

Syntax

```
submodelObj = addmodel(modelObj,  
'NameValue')  
submodelObj = addmodel(...'PropertyName', PropertyValue...)
```

Arguments

<i>modelObj</i>	Model object. Enter a name for a model object.
<i>NameValue</i>	Descriptive name for a model object. Enter a unique character string. A model object can be referenced by other objects using this property.
<i>submodelObj</i>	Model object to be added as submodel.

Description

submodelObj = addmodel(*modelObj*, '*NameValue*') creates a submodel object and returns to *submodelObj*. In the submodel object, this method assigns a value (*NameValue*) to the property Name, and assigns the model object (*modelObj*) to the property Parent. In the model object, this method assigns the submodel object to the property Models.

```
modelObj = sbiomodel('cell')  
submodelObj = addmodel('nucleus')
```

modelObj

```
modelObj.Name = 'cell'
```

modelObj.Models(1)

```
modelObj.Models(1).Name = 'nucleus'  
modelObj.Models(1).Parent = modelObj
```

A model object must have a unique name at the level it is created. For example, if you create a model with the name *cell*, you cannot create another model object named *cell1*. However, a model object can contain a submodel object named *cell* which can contain a submodel object named *cell1*.

modelObj does not have access to *submodelObj* parameters. However, *submodelObj* does have access and can use *modelObj* parameters.

submodelObj = addmodel(...'PropertyName', *PropertyValue*...) defines optional property values. The property name/property value pairs can be in any format supported by the function set (for example, name-value string pairs, structures, and name-value cell array pairs).

You can view additional model object properties with the function *get*. You can change additional model object properties with the function *set*. You can view the submodel objects of *modelObj* with the command, *get(modelObj, 'Models')*.

Examples

```
modelObj = sbiomodel('cell');
submodelObj = addmodel(modelObj, 'nucleus');
submodelObj = addmodel(modelObj, 'cytoplasm');
```

See Also

sbiomodel

addparameter (model, kineticlaw)

Purpose

Add parameter object to model or kinetic law object

Syntax

```
parameterObj = addparameter(Obj,  
    'NameValue')  
parameterObj = addparameter(Obj, 'NameValue', ValueValue)  
parameterObj = addparameter(...'PropertyName', PropertyValue...)
```

Arguments

<i>Obj</i>	Model or kinetic law object. Enter a variable name for the object.
<i>NameValue</i>	Property for a parameter object. Enter a unique character string. <i>NameValue</i> can be a cell array of parameter names. Since objects can use this property to reference a parameter, a parameter object must have a unique name at the level it is created. For example, a kinetic law object cannot contain two parameter objects named <i>kappa</i> . However, the model object that contains the kinetic law object can contain a parameter object named <i>kappa</i> along with the kinetic law object.
<i>ValueValue</i>	Property for a parameter object. Enter a number.

Description

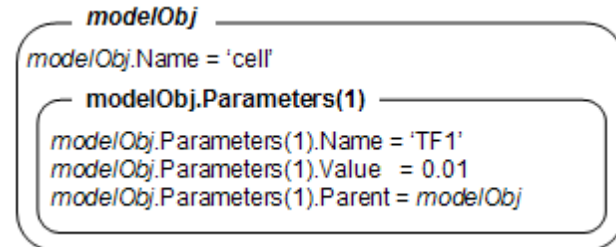
`parameterObj = addparameter(Obj, 'NameValue')` creates a parameter object and returns the object (*parameterObj*). In the parameter object, this method assigns a value (*NameValue*) to the property *Name*, assigns a value 1 to the property *Value*, and assigns the model or kinetic law object to the property *Parent*. In the model or kinetic law object, (*Obj*), this method assigns the parameter object to the property *Parameters*.

A parameter object defines an assignment that a model, or a kinetic law can use. The scope of the parameter is defined by the parameter parent. If a parameter is defined with a kinetic law object, then only the kinetic law object and objects within the kinetic law object can use the parameter. If a parameter object is defined with a model object as its

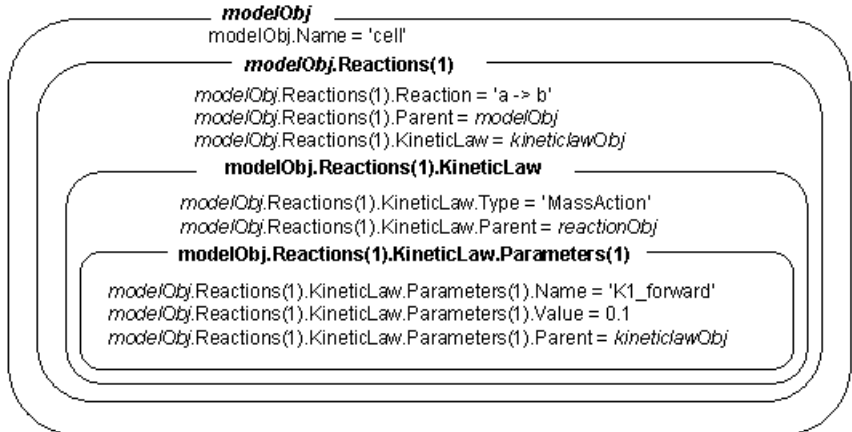
addparameter (model, kineticlaw)

parent, then all objects within the model (including all rules, submodels and kinetic laws) can use the parameter.

```
modelObj = sbiomodel('cell')
parameterObj = addparameter(modelObj, 'TF1', 0.01)
```



```
modelObj = sbiomodel('cell')
reactionObj = addreaction(modelObj, 'a -> b')
kineticlawObj = addkineticlaw(reactionObj, 'MassAction')
parameterObj = addparameter(kineticlawObj, 'K1_forward', 0.1)
```



`parameterObj = addparameter(Obj, 'NameValue', ValueValue)` creates a parameter object, assigns a value (`NameValue`) to the property Name,

addparameter (model, kineticlaw)

assigns the value (*ValueValue*) to the property *Value*, and assigns the model object or the kinetic law object to the property *Parent*. In the model or kinetic law object (*Obj*), this method assigns the parameter object to the property *Parameters*, and returns the parameter object to a variable (*parameterObj*).

`parameterObj = addparameter(...'PropertyName',
PropertyValue...)` defines optional property values. The property name/property value pairs can be in any format supported by the function set (for example, name-value string pairs, structures, and name-value cell array pairs).

Scope of a parameter — A parameter can be *scoped* to either a model or a kinetic law.

- When a kinetic law searches for a parameter in its expression, it first looks in the parameter list of the kinetic law. If the parameter isn't found there it moves to the model that the kinetic law object is in and looks in the model parameter list. If the parameter isn't found there, it moves to the model parent.
- When a rule searches for a parameter in its expression, it looks in the parameter list for the model. If the parameter isn't found there, it moves to the model parent. A rule cannot use a parameter that is scoped to a kinetic law. So for a parameter to be used in both a reaction rate equation and a rule, the parameter should be *scoped* to a model.

Additional parameter object properties can be viewed with the `get` command. Additional parameter object properties can be modified with the `set` command. The parameters of *Obj* can be viewed with `get(Obj, 'Parameters')`

A SimBiology parameter object can be copied to a SimBiology model or kinetic law object with `copyobj`. A SimBiology parameter object can be removed from a SimBiology model or kinetic law object with `delete`.

addparameter (model, kineticlaw)

Method Summary	Methods for parameter objects	
copyobj (any object)		Copy SimBiology object and its children
delete (any object)		Delete SimBiology object
display (any object)		Display summary of SimBiology object
Property Summary	Properties for parameter objects	
Annotation		Property with information about a SimBiology object
ConstantValue		Property to indicate variable or constant parameter value
Name		Property with name of object
Notes		Property with HTML text describing SimBiology object
Parent		Property indicating the parent object
Tag		Property to specify a label for a SimBiology object
Type		Property to indicate SimBiology object type
UserData		Property to specify data to associate with object
Value		Property to assign value to parameter object
ValueUnits		Property with parameter value units

addparameter (model, kineticlaw)

Example

- 1 Create model object, then add a reaction object.

```
modelObj = sbiomodel ('my_model');  
reactionObj = addreaction (modelObj, 'a + b -> c + d');
```

- 2 Define a kinetic law for the reaction object.

```
kineticlawObj = addkineticlaw(reactionObj, 'MassAction');
```

- 3 Add a parameter and assign it to the kinetic law object (kineticlawObj); add another parameter and assign to the model object (modelObj).

```
% Add parameter to kinetic law object  
parameterObj1 = addparameter (kineticlawObj, 'K1');  
  
get (kineticlawObj, 'Parameters')
```

MATLAB returns

```
Parameter Object Array
```

Index:	Name:	Value:	ValueUnits:
1	K1	1	

```
% Add parameter with value 0.9 to model object  
parameterObj1 = addparameter (modelObj, 'K2', 0.9);
```

```
get (modelObj, 'Parameters')
```

MATLAB returns

```
Parameter Object Array
```

Index:	Name:	Value:	ValueUnits:
1	K2	1	

See Also

MATLAB functions—copyobj, delete

Purpose

Add product species object to reaction object

Syntax

```
speciesObj = addproduct(reactionObj, 'NameValue')
speciesObj = addproduct(reactionObj, speciesObj)
speciesObj = addproduct(reactionObj,
    'NameValue', Stoichcoefficient)
speciesObj = addproduct(reactionObj, speciesObj, Stoichcoefficient)
```

Arguments

reactionObj Reaction object. Enter a name for the reaction object.

NameValue Property of a species object that names the object (not the reaction object). Enter a unique character string. For example, 'fructose 6-phosphate'. A species object can be referenced by other objects using this property. You can use the function `sbioselect` to find an object with a specific *NameValue*.

speciesObj Species object.

Stoichcoeffient Stoichiometric coefficients for products, length of array equal to length of *NameValue* or length of *speciesObj*.

Description

speciesObj = `addproduct(reactionObj, 'NameValue')` creates a species object and returns the species object (*speciesObj*). In the species object, this method assigns the value (*NameValue*) to the property `Name`, and assigns the parent object of the *reactionObj* to the property `Parent`. In the reaction object, this method assigns the species object to the property `Products`, modifies the reaction equation in the property `Reaction` to include the new species, and adds the stoichiometric coefficient 1 to the property `Stoichiometry`.

If the parent object (always a model object) of a reaction does not include a species with the specified name ('*NameValue*'), a species object

addproduct (reaction)

is created and assigned to the parent object property `Species`. You can create a species object with the function `sbiospecies`, or create and add a species object to a model object with the method `addspecies`.

`speciesObj = addproduct(reactionObj, speciesObj)`, in the species object (`speciesObj`), assigns the parent object of the `reactionObj` to the species property `Parent`. In the reaction object (`reactionObj`), it assigns the species object to the property `Products`, modifies the reaction equation in the property `Reaction` to include the new species, and adds the stoichiometric coefficient 1 to the property `Stoichiometry`.

`speciesObj = addproduct(reactionObj, 'NameValue', Stoichcoefficient)`, in addition to the description above, this method adds the stoichiometric coefficient (`Stoichcoefficient`) to the property `Stoichiometry`. If `NameValue` is a cell array of species names, then `Stoichcoefficient` must be a vector of doubles with the same length as `NameValue`.

`speciesObj = addproduct(reactionObj, speciesObj, Stoichcoefficient)`, in addition to the description above, this method adds the stoichiometric coefficient (`Stoichcoefficient`) to the property `Stoichiometry`.

Species names are referenced by reaction objects, kinetic law objects, and model objects. If you change the Name of a species SimBiology updates the reaction to use the new name. You must however configure all other applicable elements such as rules that use the species, and the kinetic law object.

See “Valid Species Names” on page 4-38 for more information on species names.

Example

- 1 Create a model object, and then add a reaction object.

```
modelObj = sbiomodel('my_model');
reactionObj = addreaction(modelObj, 'A + C -> U');
```

- 2 Modify the reaction of the `reactionObj` from $A + C \rightarrow U$ to $A + C \rightarrow U + 2 H$.

```
speciesObj = addproduct(reactionObj, 'H', 2);
```

See Also [sbiospecies](#), [addspecies](#)

addreactant (reaction)

Purpose Add species object as a reactant to reaction object

Syntax

```
speciesObj = addreactant(reactionObj, 'NameValue')
addreactant(reactionObj, speciesObj, Stoichcoeffieient)
addreactant(reactionObj,
'NameValue', Stoichcoeffieient)
```

Arguments

<i>reactionObj</i>	Reaction object.
<i>NameValue</i>	Name property of a species object. Enter a unique character string, for example, 'glucose 6-phosphate'. A species object can be referenced by other objects using this property. You can use the function sbioselect to find an object with a specific Name property value.
<i>speciesObj</i>	Species object or cell array of species objects.
<i>Stoichcoeffieient</i>	Stoichiometric coefficients for reactants, length of array equal to length of <i>NameValue</i> or length of <i>speciesObj</i> .

Description

speciesObj = addreactant(*reactionObj*, '*NameValue*') creates a species object and returns the species object (*speciesObj*). In the species object, this method assigns the value (*NameValue*) to the property Name, and assigns the parent object of the *reactionObj* to the property Parent. In the reaction object, this method assigns the species object to the property Reactants, modifies the reaction equation in the property Reaction to include the new species, and adds the stoichiometric coefficient -1 to the property Stoichiometry.

If the parent object (always a model object) of a reaction does not include a species with the specified name ('*NameValue*'), SimBiology creates a species object and assigns the parent object of the *reactionObj* to the parent object property Species. You can create a species object with the

function `sbiospecies`, or create and add a species object to a model object with the method `addsspecies`.

`addreactant(reactionObj, speciesObj, Stoichcoefficient)`, in the species object (`speciesObj`), this method assigns the parent object to the `speciesObj` property `Parent`. In the reaction object (`reactionObj`), it assigns the species object to the property `Reactants`, modifies the reaction equation in the property `Reaction` to include the new species, and adds the stoichiometric coefficient -1 to the property `Stoichiometry`. If `speciesObj` is a cell array of species objects, then `Stoichcoefficient` must be a vector of doubles with the same length as `speciesObj`.

`addreactant(reactionObj, 'NameValue', Stoichcoefficient)`, in addition to the description above, this method adds the stoichiometric coefficient (`Stoichcoefficient`) to the property `Stoichiometry`. If `NameValue` is a cell array of species names, then `Coefficient` must be a vector of doubles with the same length as `NameValue`.

A species object must have a unique name at the level at which it is created. For example, a model object cannot contain two species objects named `H2O`. However, a submodel of the model that contains the species `H2O` can also contain a species named `H2O`.

Species names are referenced by reaction objects, kinetic law objects, and model objects. If you change the Name of a species SimBiology updates the reaction to use the new name. You must however configure all other applicable elements such as rules that use the species, and the kinetic law object.

See “Valid Species Names” on page 4-38 for more information on species names.

Example

- 1 Create a model object, and then add a reaction object.

```
modelObj = sbiomodel('my_model');
reactionObj = addreaction(modelObj, 'A -> U');
```

- 2 Modify the reaction of the `reactionObj` from $A \rightarrow U$ to be $A + 3 C \rightarrow U$.

addreactant (reaction)

```
speciesObj = addreactant(reactionObj, 'C', 3);
```

See Also

[sbiospecies](#), [addspecies](#)

Purpose	Add reaction object to model object
Syntax	<pre>reactionObj = addreaction(modelObj, 'ReactionValue') reactionObj = addreaction(modelObj, 'ReactantsValue', 'ProductsValue') reactionObj = addreaction(modelObj, 'ReactantsValue', RStoichCoefficients, 'ProductsValue', PStoichCoefficients) reactionObj = addreaction(...'PropertyName', PropertyValue...)</pre>
Arguments	
<i>modelObj</i>	SimBiology model object
<i>ReactionValue</i>	Specify the reaction equation. Enter a character string. A hyphen preceded by a space and followed by a right angle bracket (->) indicate reactants going forward to products. A hyphen with left and right angle brackets (<->) indicate a reversible reaction. Coefficients before reactant or product names must be followed by a space. Examples 'A -> B', 'A + B -> C', '2 A + B -> 2 C', 'A <-> B'. Enter reactions with spaces between the species (A + B -> C)
<i>ReactantsValue</i>	A string defining the species name, a cell array of strings, a species object or an array of species objects.
<i>ProductsValue</i>	A string defining the species name, a cell array of strings, a species object or an array of species objects.

addreaction (model)

<i>RStoichCoefficients</i>	Stoichiometric coefficients for reactants, length of array equal to length of <i>ReactantsValue</i> .
<i>PStoichCoefficients</i>	Stoichiometric coefficients for products, length of array equal to length of <i>ProductsValue</i> .

Description

`reactionObj = addreaction(modelObj, 'ReactionValue')` creates a reaction object, assigns a value (*ReactionValue*) to the property `Reaction`, assigns reactant species object(s) to the property `Reactants`, assigns the product species object(s) to the property `Products`, and assigns the model object to the property `Parent`. In the Model object (`modelObj`), this method assigns the reaction object to the property `Reactions`, and returns the reaction object (`reactionObj`).

```
reactionObj = addreaction(modelObj, 'a -> b')
```

```
modelObj
  mode/Obj.Name = 'cell'
  mode/Obj.Reactions(1)
    mode/Obj.Reactions(1).Reaction = 'a -> b'
    mode/Obj.Parameters(1).Parent = mode/Obj
```

If a species specified in a reaction does not exist, a species object is created and assigned to the model object property `Species`. You can manually add a species to a model object with the method `addspecies`.

You can add species to a reaction object using the methods `addreactant` or `addproduct`. You can remove species from a reaction object with the methods `rmreactant` or `rmproduct`. The property `Reaction` is modified by adding or removing species from the reaction equation.

You can copy a SimBiology reaction object can be copied to a SimBiology model object with the function, `copyobj`. You can remove SimBiology reaction object from a SimBiology model object with the function `delete`.

You can view additional reaction object properties with the `get` command, for example, the reaction equation of `reactionObj` can be

viewed with the command, `get(reactionObj, 'Reaction')`. You can modify additional reaction object properties with the command, `set`.

`reactionObj = addreaction(modelObj, 'ReactantsValue', 'ProductsValue')` creates a reaction object, assigns a value to the property `Reaction` using the reactant (`ReactantsValue`) and product (`ProductsValue`) names, assigns the species objects to the properties `Reactants` and `Products`, and assigns the model object to the property `Parent`. In the Model object (`modelObj`), this method assigns the reaction object to the property `Reactions`, and returns the reaction object (`reactionObj`). The stoichiometric values are assumed to be 1.

`reactionObj = addreaction(modelObj, 'ReactantsValue', RStoichCoefficients, 'ProductsValue', PStoichCoefficients)` adds stoichiometric coefficients (`RStoichCoefficients`) for reactant species, and stoichiometric coefficients (`PStoichCoefficients`) for product species to the property `Stoichiometry`. The length of `Reactants` and `RCoefficients` must be equal, and the length of `Products` and `PCoefficients` must be equal.

`reactionObj = addreaction(...'PropertyName', PropertyValue...)` defines optional properties. The property name/property value pairs can be in any format supported by the function `set` (for example, name-value string pairs, structures, and name-value cell array pairs).

Method Summary

Methods for reaction objects

<code>addkineticlaw (reaction)</code>	Add kinetic law object to reaction object
<code>addproduct (reaction)</code>	Add product species object to reaction object
<code>addreactant (reaction)</code>	Add species object as a reactant to reaction object
<code>copyobj (any object)</code>	Copy SimBiology object and its children

addreaction (model)

delete (any object)	Delete SimBiology object
display (any object)	Display summary of SimBiology object
rmproduct (reaction)	Remove species object from reaction object products
rmreactant (reaction)	Remove species object from reaction object reactants

Property Summary

Properties for reaction objects	
Active	Property to indicate object use during a simulation
Annotation	Property with information about a SimBiology object
KineticLaw	Property showing kinetic law for ReactionRate
Name	Property with name of object
Notes	Property with HTML text describing SimBiology object
Parent	Property indicating the parent object
Products	Property to indicate reaction products
Reactants	Property to indicate reaction reactants
Reaction	Property to indicate the reaction object reaction
ReactionRate	Property containing the reaction rate equation in reaction object

Reversible	Property to indicate whether a reaction is reversible or irreversible
Stoichiometry	Property that describes species coefficients in a reaction
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Examples

Create a model, add a reaction object and assign the expression for the reaction rate equation.

- 1 Create a model object, then add a reaction object.

```
modelObj = sbiomodel('my_model');
reactionObj = addreaction(modelObj, 'a -> c + d');
```

- 2 Create a kinetic law object for the reaction object, of the type 'Henri-Michaelis-Menten' .

```
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');
```

reactionObj KineticLaw property is configured to kineticlawObj.

- 3 The 'Henri-Michaelis-Menten' kinetic law has two parameter variables (V_m and K_m) and one species variable (S) that should to be set. To set these variables, first create the parameter variables as parameter objects (parameterObj1, parameterObj2) with names V_m_d , and K_m_d , and assign the objects Parent property value to the kineticlawObj.

```
parameterObj1 = addparameter(kineticlawObj, 'Vm_d');
```

addreaction (model)

```
parameterObj2 = addparameter(kineticlawObj, 'Km_d');
```

- 4** Set the variable names for the kinetic law object.

```
set(kineticlawObj,'ParameterVariableNames', {'Vm_d' 'Km_d'});  
set(kineticlawObj,'SpeciesVariableNames', {'a'});
```

- 5** Verify that the reaction rate is expressed correctly in the reaction object ReactionRate property.

```
get (reactionObj, 'ReactionRate')
```

MATLAB returns

```
ans =
```

```
Vm_d*[a]/(Km_d+[a])
```

See Also

[addkineticlaw](#), [addproduct](#), [addreactant](#), [rmproduct](#), [rmreactant](#)

Purpose Add rule object to model object

Syntax

```
ruleObj = addrule(modelObj,  
'RuleValue')  
ruleObj = addrule(modelObj,  
'RuleValue', 'RuleTypeValue')  
ruleObj = addrule(..., 'PropertyName', PropertyValue,...)
```

Arguments

<i>modelObj</i>	Model object to which to add the rule.
<i>RuleValue</i>	Enter a character string within quotes. For example, enter the algebraic rule ' $V_a * E_a + V_i * E_i - K_2$ '.
<i>RuleTypeValue</i>	Enter 'algebraic', 'assignment', or 'rate'. An algebraic or rate rule is evaluated at each time step during the simulation. An assignment rule is evaluated once before the simulation starts. Note: if a species or parameter is marked constant, you can still assign an initial value using an assignment rule. The amount or value gets assigned according to the rule and then remains constant during the simulation.

Description

A rule is a mathematical expression that changes the amount of a species or the value of a parameter. It also defines how species and parameters interact with one another.

ruleObj = `addrule(modelObj, 'RuleValue')` creates a rule object and returns the rule object (*ruleObj*). In the rule object, this method assigns a value ('*RuleValue*') to the property `Rule`, assigns the value 'algebraic' to the property `RuleType`, and assigns the model object (*modelObj*) to the property `Parent`. In the model object (*modelObj*), this method assigns the rule object to the property `Rules`.

addrule (model)

`ruleObj = addrule(modelObj, 'RuleValue', 'RuleTypeValue')` in addition to the assignments above, assigns a value (`RuleTypeValue`) to the property `RuleType`. For more information on the different types of rules see `RuleType`.

`ruleObj = addrule(..., 'PropertyName', PropertyValue,...)` defines optional properties. The property name/property value pairs can be in any format supported by the function set (for example, name-value string pairs, structures, and name-value cell array pairs).

View additional rule properties with the function `get`, and modify rule properties with the function `set`. Copy a rule object to a model with the function `copyobj`, or delete a rule object from a model with the function `delete`.

Method Summary

Methods for rule objects

<code>copyobj (any object)</code>	Copy SimBiology object and its children
<code>delete (any object)</code>	Delete SimBiology object
<code>display (any object)</code>	Display summary of SimBiology object

Property Summary

Properties for rule objects

<code>Active</code>	Property to indicate object use during a simulation
<code>Annotation</code>	Property with information about a SimBiology object
<code>Name</code>	Property with name of object
<code>Notes</code>	Property with HTML text describing SimBiology object
<code>Parent</code>	Property indicating the parent object

Rule	Property to define certain species and parameter interactions
RuleType	Property for defining the type of rule for the rule object
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Examples

Add a rule with default RuleType.

- 1 Create a model object, and then add a rule object.

```
modelObj = sbiomodel('cell');
ruleObj = addrule(modelObj, '0.1*B-A')
```

- 2 Get a list of properties for a rule object.

```
get(modelObj.Rules(1)) or get(ruleObj)
```

MATLAB displays a list of rule properties.

```
Active: 1
Annotation: ''
Name: ''
Notes: ''
Parent: [1x1 SimBiology.Model]
Rule: '0.1*B-A'
RuleType: 'algebraic'
Tag: ''
Type: 'rule'
UserData: []
```

addrule (model)

Add rule with RuleType property set to rate.

- 1 Create model object, then add a reaction object

```
modelObj = sbiomodel ('my_model');  
reactionObj = addreaction (modelObj, 'a -> b');
```

- 2 Add a rule which defines that the quantity of a species c. In the rule expression k is the rate constant for $a \rightarrow b$.

```
ruleObj = addrule(modelObj, 'c = k*(a+b)')
```

- 3 Change the RuleType from default ('algebraic') to 'rate'. and verify using the get command.

```
set(ruleObj, 'RuleType', 'rate');  
get(ruleObj)
```

MATLAB returns all the properties for the rule object.

```
Active: 1  
Annotation: ''  
Name: ''  
Notes: ''  
Parent: [1x1 SimBiology.Model]  
Rule: 'c = k*(a+b)'  
RuleType: 'rate'  
Tag: ''  
Type: 'rule'  
UserData: []
```

See Also

[copyobj](#), [delete](#), [sbiomodel](#)

Purpose

Add species object to model object

Syntax

```
speciesObj = addspecies(modelObj, NameValue)
speciesObj = addspecies(modelObj, NameValue,
InitialAmountValue)
speciesObj = addspecies(...'PropertyName', PropertyValue...)
```

Arguments

<i>modelObj</i>	Model object.
<i>NameValue</i>	Name for a species object. Enter a character string unique to the level of object creation. Species objects are identified by Name within ReactionRate and Rule property strings. You can use the function sbioselect to find an object with a specific Name property value.
<i>InitialAmountValue</i>	Initial amount value for the species object. Enter double. Positive real number, default = 0.

Description

speciesObj = addspecies(*modelObj*, *NameValue*) creates a species object and returns the species object (*speciesObj*). In the species object, this method assigns a value (*NameValue*) to the property Name, assigns the model object (*modelObj*) to the property Parent. In the model object, this method assigns the species object to the property Species.

speciesObj = addspecies(*modelObj*, *NameValue*, *InitialAmountValue*), in addition to the above, this method assigns an initial amount (*InitialAmountValue*) for the species.

You can also add a species to a reaction using the methods addreactant and addproduct . When a reaction is defined with a species not in the model object (*modelObj*), SimBiology creates a species object.

A species object must have a unique name at the level at which it is created. For example, a model object cannot contain two species objects named H2O. However, a submodel of the model that contains the species H2O can also contain a species named H2O.

addspecies (model)

View properties for a species object with the `get` command, and modify properties for a species object with the `set` command. You can view a summary table of species objects in a model (`Mobj`) with `get(Mobj, 'Species')` or the properties of the first species with `get(Mobj.Species(1))`.

A species in a rule has to be in the model object with the rule. This is different from parameters in a rule, where a parameter can be in the model object or in the kinetic law object or resolve hierarchically.

`speciesObj = addspecies(...'PropertyName', PropertyValue...)` defines optional properties. The property name/property value pairs can be in any format supported by the function `set` (for example, name-value string pairs, structures, and name-value cell array pairs). The property summary on this page shows the list of properties.

Valid Species Names

SimBiology species names can have any number or character, for example, N-acetyl-D-glucosamine.

Species names, however, cannot be left empty and note the following reserved words, characters and constraints:

- The literal words `null` and `time`. Note that you could specify species names with these words contained within the name. For example `nullaminoacids`, or `nullnucleotides`.
- The characters `i`, `j`, `->`, `<>`, `[`, and `]`.
- If you are using a species name that is not a valid MATLAB variable name, do the following:
 - Enclose the name in square brackets when writing a reaction rate equation or a rule.
 - Enter the name without brackets when you are creating the species or when you are adding the reaction.

For example, enclose `[DNA polymerase+]` within brackets in reaction rates and rules; enter `DNA polymerase+` when specifying the name of the species or while writing the reaction.

Method Summary	Methods for species objects
	<code>copyobj (any object)</code>
	<code>delete (any object)</code>
	<code>display (any object)</code>
	Copy SimBiology object and its children
	Delete SimBiology object
	Display summary of SimBiology object
Property Summary	Properties for species objects
	<code>Annotation</code>
	<code>BoundaryCondition</code>
	<code>ConstantAmount</code>
	<code>InitialAmount</code>
	<code>InitialAmountUnits</code>
	<code>Name</code>
	<code>Notes</code>
	<code>Parent</code>
	<code>Tag</code>
	<code>Type</code>
	<code>UserData</code>
	Property with information about a SimBiology object
	Property to set a species object to have a boundary condition
	Property to specify variable or constant species amount
	Property containing initial amount of a species
	Property containing units for species initial amount
	Property with name of object
	Property with HTML text describing SimBiology object
	Property indicating the parent object
	Property to specify a label for a SimBiology object
	Property to indicate SimBiology object type
	Property to specify data to associate with object

addspecies (model)

Examples

Add two species to a model, one is a reactant and the other is the enzyme catalyzing the reaction.

- 1 Create a model object with the name `my_model`.

```
modelObj = sbiomodel ('my_model');
```

- 2 Add two species objects with the names `glucose_6_phosphate` and `glucose_6_phosphate_dehydrogenase`.

```
speciesObj1 = addspecies (modelObj, 'glucose_6_phosphate');
speciesObj2 = addspecies (modelObj, ...
                         'glucose_6_phosphate_dehydrogenase');
```

- 3 Set initial amount of `glucose_6_phosphate` to 100 and verify.

```
set (speciesObj, 'InitialAmount',100);
get (speciesObj, 'InitialAmount')
```

MATLAB returns

```
ans =
```

```
100
```

- 4 Use `get` to note that `modelObj` contains the species object array.

```
get(modelObj, 'Species')
```

MATLAB returns,

Species Object Array

Index: Name:

Initial Initial

Amount: AmountUnits:

1 glucose_6_phosphate

100

```
2      glucose_6_phosphate_dehydrogenase      0
```

5 Retrieve information about the first species in the array.

```
get(modelObj.Species(1))

    Annotation: ''
    ConstantAmount: 'false'
    InitialAmount: 100
    InitialAmountUnits: ''
        Name: 'glucose_6_phosphate'
        Notes: ''
    Parent: [1x1 SimBiology.Model]
        Tag: ''
        Type: 'species'
    UserData: []
```

See Also

[addproduct](#), [addreactant](#), [addreaction](#)

MATLAB functions— [get](#) and [set](#)

copyobj (any object)

Purpose Copy SimBiology object and its children

Syntax

```
copiedObj = copyobj(Obj, parentObj)
copiedObj = copyobj(modelObj)
```

Arguments

<i>Obj</i>	Abstract kinetic law, configuration set, kinetic law, model, parameter, reaction, rule, or species object.
<i>parentObj</i>	If <i>copiedObj</i> is configuration set, reaction, rule or species object, <i>parentObj</i> must be a model object. If <i>copiedObj</i> is a parameter object, <i>ParentObj</i> must be a model or kinetic law object. If <i>copiedObj</i> is a model object, <i>ParentObj</i> must be a model object (for example, in the case of submodels) or sbioroot.
<i>modelObj</i>	Model object to be copied.

Description

copiedObj = *copyobj*(*Obj*, *parentObj*) makes a copy of a SimBiology object (*Obj*) and returns a pointer to the copy (*copiedObj*). In the copied object (*copiedObj*), this method assigns a value (*parentObj*) to the property Parent.

- *Obj*— Can be abstract kinetic law, configuration set, kinetic law, model, parameter, reaction, rule, or species object.
- *parentObj* — If *copiedObj* is configuration set, reaction, rule or species object, *parentObj* must be a model object. If *copiedObj* is a parameter object, *ParentObj* must be a model or kinetic law object. If *copiedObj* is a model object, *ParentObj* must be a model object (for example, in the case of submodels) or sbioroot.

copiedObj = *copyobj*(*modelObj*) makes a copy of a model object (*modelObj*) and returns the copy (*copiedObj*). In the copied model object (*copiedObj*), this method assigns the root object to the property Parent.

Example

Create a reaction object separate from a model object and then add it to a model.

- 1 Create a model object and create a separate reaction object.

```
modelObj = sbiomodel('cell');
reactionObj = sbioreaction('a -> b');
```

- 2 Create a copy of the reaction object and assign it to the model object.

```
reactionObjCopy = copyobj(reactionObj, modelObj);
modelObj.Reactions
```

```
Reaction Object Array
Index:    Reaction:
1          a -> b
```

See Also

[sbiomodel](#), [sbioreaction](#), [sbioroot](#)

delete (any object)

Purpose	Delete SimBiology object
Syntax	<code>delete(<i>Obj</i>)</code>
Arguments	<p><i>Obj</i> SimBiology object: abstract kinetic law, configuration set, kinetic law, model, parameter, reaction, rule, or species.</p>
Description	<p><code>delete(<i>Obj</i>)</code> removes an object (<i>Obj</i>) from its parent.</p> <ul style="list-style-type: none">• If <i>Obj</i> is a species object that is being used by a reaction object, this method returns an error and the species object is not deleted. You need to delete the reaction or remove the species from the reaction before you can delete the species object.• If <i>Obj</i> is a parameter object being used by a kinetic law object, there is no warning when the object is deleted. However, when you try to simulate your model, an error occurs because the parameter cannot be found.• If <i>Obj</i> is a reaction object, this method deletes the object, but the species objects that were being used by the reaction object are not deleted.• If <i>Obj</i> is an abstract kinetic law object and there is a kinetic law object referencing it, this method returns an error.• If <i>Obj</i> is a SimBiology configuration set object, and it is the active configuration set object, this method, after deleting the object, makes the default configuration set object active. Note, you cannot delete the default configuration set.• You cannot delete the SimBiology root.

You can also delete all model objects from the root with one call to the `sbioreset` function.

Examples

Example 1

Delete a reaction from a model. Notice, the species objects are not deleted with the reaction object.

```
modelObj = sbiomodel('cell');
reactionObj = addreaction(modelObj, 'a -> b');
delete(reactionObj)
```

Example 2

Delete a single model from the root object.

```
modelObj1 = sbiomodel('cell');
modelObj2 = sbiomodel('virus');
delete(ModelObj2)
```

See Also

[sbiomodel](#), [sbioreset](#), [sbioroot](#)

display (any object)

Purpose	Display summary of SimBiology object
Syntax	<code>display(Obj)</code>
Arguments	<p><i>Obj</i> SimBiology object: abstract kinetic law, configuration set, kinetic law, model, parameter, reaction, rule, or species.</p>
Description	Display the SimBiology object array. <code>display(Obj)</code> is called for the SimBiology object, <i>Obj</i> when the semicolon is not used to terminate a statement. The display of <i>Obj</i> gives a brief summary of <i>Obj</i> configuration. You can view a complete list of <i>Obj</i> properties with the command <code>get</code> . You can modify all <i>Obj</i> properties that can be changed, with the command <code>set</code> .
Examples	<pre>modelObj = sbiomodel('cell') reactionObj = addreaction(modelObj, 'A + B -> C')</pre>

getadjacencymatrix (model)

Purpose Get adjacency matrix from model object

Syntax

```
M = getadjacencymatrix(modelObj)
M = getadjacencymatrix(modelObj, 'flat')
[M,Headings]
= getadjacencymatrix(modelObj)
[M, Headings, Mask]=getadjacencymatrix(modelObj)
```

Arguments

<i>M</i>	Adjacency matrix for <i>modelObj</i>
<i>modelObj</i>	specify model object <i>modelObj</i>
'flat'	Return adjacency matrix for only specified <i>modelObj</i> not for objects contained in the <i>modelObj</i>
<i>Headings</i>	Return row and column headings to <i>Headings</i>
<i>Mask</i>	Returns 1 for species object 0 for reaction object to <i>Mask</i>

Description

`getadjacencymatrix` returns adjacency matrix for model object.

`M = getadjacencymatrix(modelObj)` returns adjacency matrix for model object, (*modelOBJ*) to *M*.

An adjacency matrix is defined by listing all species contained by *modelObj* and all reactions contained by *modelObj* column-wise and row-wise in a matrix. The reactants of the reactions are represented in the matrix with a 1 at the location of [row of species, column of reaction]. The products of the reactions are represented in the matrix with a 1 at the location of [row of reaction, column of species]. All other locations in the matrix are 0.

`M = getadjacencymatrix(modelObj, 'flat')` returns the adjacency matrix to *M* and defines the adjacency matrix for only *modelObj*. If *modelObj* is a SimBiology model then *M* is the adjacency matrix for the reactions

getadjacencymatrix (model)

and species contained by *modelObj*. *M* does not include any submodel reaction or species information.

[*M*, *Headings*] = getadjacencymatrix(*modelObj*) returns the adjacency matrix to *M* and the row and column headings to *Headings*. *Headings* is defined by listing all Name property values of species contained by *modelObj* and all Name property values of reactions contained by *modelObj*. In the above example, *Headings* would be {'A', 'B', 'C', 'R1'}.

[*M*, *Headings*, *Mask*] = getadjacencymatrix(*modelObj*) returns an array of ones and zeros to *Mask* where a 1 represents a species object and a 0 represents a reaction object. In the above example, *Mask* would be [1 1 1 0].

Examples

- 1 Read in a model using sbmlimport.

```
modelObj = sbmlimport('lotka.xml');
```

- 2 Get the adjacency matrix for the *modelObj*.

```
[M, Headings] = getadjacencymatrix(modelObj)
```

See Also

[getstoichmatrix](#)

Purpose	Get configuration set object from model object						
Syntax	<pre>configsetObj = getconfigset(modelObj, 'NameValue') configsetObj = getconfigset(modelObj) configsetObj = getconfigset(modelObj, 'active')</pre>						
Arguments	<table><tr><td><i>modelObj</i></td><td>Model object. Enter a variable name for a model object.</td></tr><tr><td><i>NameValue</i></td><td>Name of the configset object.</td></tr><tr><td><i>configsetObj</i></td><td>Object holding the simulation specific information.</td></tr></table>	<i>modelObj</i>	Model object. Enter a variable name for a model object.	<i>NameValue</i>	Name of the configset object.	<i>configsetObj</i>	Object holding the simulation specific information.
<i>modelObj</i>	Model object. Enter a variable name for a model object.						
<i>NameValue</i>	Name of the configset object.						
<i>configsetObj</i>	Object holding the simulation specific information.						
Description	<p><i>configsetObj</i> = <code>getconfigset(modelObj, 'NameValue')</code> returns the configuration set attached to <i>modelObj</i> that is named <i>NameValue</i>, to <i>configsetObj</i>.</p> <p><i>configsetObj</i> = <code>getconfigset(modelObj)</code> returns a vector of all attached configuration sets, to <i>configsetObj</i>.</p> <p><i>configsetObj</i> = <code>getconfigset(modelObj, 'active')</code> retrieves the active configuration set.</p> <p>A configuration set object stores simulation specific information. A SimBiology model can contain multiple configsets with one being active at any given time. The active configuration set contains the settings that are used during the simulation.</p> <p>Use the <code>setactiveconfigset</code> function to define the active configset. <i>modelObj</i> always contains at least one configset object with name configured to 'default'. Additional configset objects can be added to <i>modelObj</i> with the method, <code>addconfigset</code> .</p>						
Example	<p>1 Retrieve the defaultconfigset object from the <i>modelObj</i>.</p> <pre>modelObj = sbiomodel('cell');</pre>						

getconfigset (model)

```
configsetObj = getconfigset(modelObj)

    Configuration Settings - default (active)
        SolverType:          ode15s
        StopTime:           10.000000

        SolverOptions:
            AbsoluteTolerance: 1.000000e-006
            RelativeTolerance: 1.000000e-003

        RuntimeOptions:
            StatesToLog:       all

        CompileOptions:
            UnitConversion:    true
            DimensionalAnalysis: true
```

2 Configure the SolverType to ssa.

```
set(configsetObj, 'SolverType', 'ssa')
get(configsetObj)

    Active: 1
    CompileOptions: [1x1 SimBiology.CompileOptions]
        Name: 'default'
        Notes: ''
    RuntimeOptions: [1x1 SimBiology.RuntimeOptions]
        SolverOptions: [1x1 SimBiology.SSASolverOptions]
            SolverType: 'ssa'
            StopTime: 10
            StopTimeType: 'simulationTime'
            TimeUnits: 'second'
            Type: 'configset'
```

See Also

[addconfigset](#), [removeconfigset](#), [setactiveconfigset](#)

getparameters (kineticlaw)

Purpose

Get specific parameters in kinetic law object

Syntax

```
parameterObj = getparameters(kineticlawObj)
parameterObj = getparameters(kineticlawObj,
    'ParameterVariablesValue')
```

Arguments

kineticlawObj Retrieve parameters used by kinetic law object.

ParameterVariablesValue Retrieve parameters used by kinetic law object corresponding to the specified parameter in *ParameterVariables* property of the kinetic law object.

Description

parameterObj = *getparameters(kineticlawObj)* returns the parameters used by the kinetic law object *kineticlawObj* to *parameterObj*.

parameterObj = *getparameters(kineticlawObj,*
'ParameterVariablesValue") returns the parameter in the
ParameterVariableNames property that corresponds to the parameter
specified in the ParameterVariables property of *kineticlawObj*,
to *parameterObj*. ParameterVariablesValue is the name of the
parameter as it appears in the ParameterVariablesproperty of
kineticlawObj. ParameterVariablesValue can be a cell array of
strings.

If you change the name of a parameter you must configure all applicable elements such as rules that use the parameter, any user specified ReactionRate, or the kinetic law object property ParameterVariableNames. Use the method setparameter to configure ParameterVariableNames.

Example

Create a model, add a reaction and assign the ParameterVariableNames for the reaction rate equation.

- 1 Create model object, and then add a reaction object.

getparameters (kineticlaw)

```
modelObj = sbiomodel('my_model');
reactionObj = addreaction(modelObj, 'a -> c + d');
```

- 2** Create a kinetic law object for the reaction object, of the type 'Henri-Michaelis-Menten'.

```
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');
```

- 3** Add two parameter objects.

```
parameterObj1 = addparameter(kineticlawObj, 'Va');
parameterObj2 = addparameter(kineticlawObj, 'Ka');
```

- 4** The 'Henri-Michaelis-Menten' kinetic law has two parameter variables (V_m and K_m) that should to be set. To set these variables,

```
setparameter(kineticlawObj, 'Vm', 'Va');
setparameter(kineticlawObj, 'Km', 'Ka');
```

- 5** To retrieve a parameter variable,

```
parameterObj3 = getparameters(kineticlawObj, 'Vm')
```

MATLAB returns

Parameter Object Array

Index:	Name:	Value:	ValueUnits:
1	Va	1	

```
parameterObj4 = getparameters (kineticlawObj, 'Km')
```

See Also

[addparameter](#), [getspecies](#), [setparameter](#)

getspecies (kineticlaw)

Purpose

Get specific species in kinetic law object

Syntax

```
speciesObj = getspecies(kineticlawObj)
speciesObj = getspecies(kineticlawObj,
    'SpeciesVariablesValue')
```

Arguments

<i>kineticlawObj</i>	Retrieve species used by kinetic law object.
<i>SpeciesVariablesValue</i>	Retrieve species used by kinetic law object corresponding to the specified species in the SpeciesVariables property of the kinetic law object.

Description

speciesObj = *getspecies(kineticlawObj)* returns the species used by the kinetic law object *kineticlawObj* to *speciesObj*.

speciesObj = *getspecies(kineticlawObj, 'SpeciesVariablesValue')* returns the species in the SpeciesVariableNames property to *speciesObj*.

SpeciesVariablesValue is the name of the species as it appears in the SpeciesVariables property of *kineticlawObj*. *SpeciesVariablesValue* can be a cell array of strings.

Species names are referenced by reaction objects, kinetic law objects, and model objects. If you change the name of a species SimBiology updates the reaction to use the new name. You must however configure all other applicable elements such as rules that use the species, and the kinetic law object SpeciesVariableNames. Use the method *setspecies* to configure SpeciesVariableNames.

Example

Create a model, then add a reaction, and assign the SpeciesVariableNames for the reaction rate equation.

- 1 Create model object, then add a reaction object.

getspecies (kineticlaw)

```
modelObj = sbiomodel('my_model');
reactionObj = addreaction(modelObj, 'a -> c + d');
```

- 2** Create a kinetic law object for the reaction object, of the type 'Henri-Michaelis-Menten' .

```
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');
```

reactionObj KineticLaw property is configured to kineticlawObj.

- 3** The 'Henri-Michaelis-Menten' kinetic law has one species variable (S) that should to be set. To set this variable,

```
setspecies(kineticlawObj, 'S', 'a');
```

- 4** Retrieve the species variable using getspecies.

```
speciesObj = getspecies(kineticlawObj, 'S')
```

MATLAB returns

Species Object Array

Index:	Name:	InitialAmount:	InitialAmountUnits:
1	a	0	

See Also

[addspecies](#), [setspecies](#), [getparameters](#), [setparameter](#)

Purpose

Get stoichiometry matrix from model object

Syntax

```
M = getstoichmatrix(modelObj)
M = getstoichmatrix(modelObj,
'flat')
[M,objSpecies]=
getstoichmatrix(modelObj)
[M,objSpecies,objReactions]=
getstoichmatrix(modelObj)
```

Arguments

<i>M</i>	Adjacency matrix for <i>modelObj</i> .
<i>modelObj</i>	Specify model object <i>modelObj</i> .
'flat'	Return stoichiometry matrix for only specified <i>modelObj</i> not for objects contained in the <i>Obj</i> .
<i>objSpecies</i>	Return list of <i>Obj</i> species by Name property of species.
<i>objReactions</i>	Return list of <i>Obj</i> reactions by Name property of reactions.

Description

`getstoichmatrix` returns a stoichiometry matrix for a model object.

`M = getstoichmatrix(modelObj)` returns a stoichiometry matrix for SimBiology a model object, (*modelObj*) to *M*.

A stoichiometry matrix is defined by listing all reactions contained by *modelObj* column-wise and all species contained by *modelObj* row-wise in a matrix. The species of the reaction are represented in the matrix with the stoichiometric value at the location of [row of species, column of reaction]. Reactants have negative values. Products have positive values. All other locations in the matrix are 0.

For example, if *modelObj* is a model object with two reactions with names R1 and R2 and Reaction values of: 2 A + B -> 3 C and B + 3 D -> 4 A, the stoichiometry matrix would be defined as:

getstoichmatrix (model)

	A	B	C	D
R1	-2	-1	3	0
R2	4	-1	0	-3

M = getstoichmatrix(*modelObj*, 'flat') defines the stoichiometry matrix for only *modelObj*. If *Obj* is a SimBiology model then *M* is the stoichiometry matrix for the reactions and species contained by *modelObj*. *M* does not include any submodel reaction or species information.

[*M*,*objSpecies*]= getstoichmatrix(*modelObj*) returns the stoichiometry matrix to *M* and the species to *objSpecies*. *objSpecies* is defined by listing all Name property values of species contained by *Obj*. In the above example, *objSpecies* would be {'A', 'B', 'C', 'D'};

[*M*,*objSpecies*,*objReactions*]= getstoichmatrix(*modelObj*) returns the stoichiometry matrix to *M* and the reactions to *objReactions*. *objReactions* is defined by listing all Name property values of reactions contained by *Obj*. In the above example, *ObjReactions* would be {'R1', 'R2'}.

Example

- 1 Read in a model using sbmlimport.

```
modelObj = sbmlimport('lotka.xml');
```

- 2 Get the stoichiometry matrix for the *modelObj*.

```
[M,objSpecies,objReactions] = getstoichmatrix(modelObj)
```

See Also

[getadjacencymatrix](#)

Purpose

Remove configuration set from model

Syntax

```
removeconfigset(modelObj,  
    'NameValue')  
removeconfigset(modelObj, configsetObj)
```

Arguments

<i>modelObj</i>	Model object from which to remove configuration set.
<i>NameValue</i>	Name of the configuration set.
<i>configsetObj</i>	Configuration set object that is to be removed from model object

Description

`removeconfigset(modelObj, 'NameValue')` removes the configset object with name, *NameValue* from SimBiology model object *modelObj*. A configuration set object stores simulation specific information. A SimBiology model can contain multiple configuration sets with one being active at any given time. The active configuration set contains the settings that are used during the simulation. *modelObj* always contains at least one configuration set object with name configured to 'default'. You cannot remove the default configuration set from *modelObj*. If the active configuration set is removed from *modelObj* then the default configuration set will be made active.

`removeconfigset(modelObj, configsetObj)` removes the configuration set object, *configsetObj* from SimBiology model, *modelObj*. The configuration set is not deleted; if you want to delete *configsetObj* use the `delete` method.

If however, there is no MATLAB variable holding the configset, `removeconfigset(modelObj, 'NameValue')`, removes the configset from the model and deletes it.

Example

- 1 Create a model object by importing the file `oscillator.xml` and add a configset.

removeconfigset (model)

```
modelObj = sbmlimport('oscillator');
configsetObj = addconfigset(modelObj, 'myset');
```

- 2** Remove the configset from modelObj by name or alternatively by indexing.

```
% Remove the configset with name 'myset'.
removeconfigset(modelObj, 'myset');

% Get all configset objects and remove the second.
configsetObj = getconfigset(modelObj);
removeconfigset(modelObj, configsetObj(2));
```

See Also

[addconfigset](#), [getconfigset](#), [setactiveconfigset](#)

Purpose

Delete all model objects from the root object

Syntax

```
reset(sbioroot)
```

Description

`reset(sbioroot)` deletes all SimBiology model objects contained by the SimBiology root. The SimBiology root object is returned with the method, `sbioroot`. This call is equivalent to `sbioreset`.

The SimBiology root object contains a list of the top-level SimBiology model objects, available units, unit prefixes, and abstract kinetic law objects. A top-level SimBiology model object has its `Parent` property set to the SimBiology root object. A SimBiology model object that has its `Parent` property set to another SimBiology model is a submodel and is not stored by the SimBiology root.

To add an abstract kinetic law to the SimBiology root user-defined library, use the `addtolibrary` function. To add a unit to the SimBiology root user-defined library, use the function, `sbioregisterunit`. To add a unit prefix to the SimBiology root user-defined library, use the function, `sbioregisterunitprefix`.

Examples

- 1 Query `sbioroot` that has two model objects.

```
sbioroot
```

SimBiology Root Contains:

Models:	2
Builtin Abstract Kinetic Laws:	3
User Abstract Kinetic Laws:	1
Builtin Units:	54
User Units:	0
Builtin Unit Prefixes:	13
User Unit Prefixes:	0

- 2 Call `reset`.

```
sbioroot
```

reset (root)

SimBiology Root Contains:

Models:	0
Builtin Abstract Kinetic Laws:	3
User Abstract Kinetic Laws:	1
Builtin Units:	54
User Units:	0
Builtin Unit Prefixes:	13
User Unit Prefixes:	0

See Also

[sbioregisterunit](#), [sbioregisterunitprefix](#), [sbioroot](#), [sbioreset](#),
[sbiohelp](#)

Purpose

Remove species object from reaction object products

Syntax

```
rmproduct(reactionObj, SpeciesName)  
rmproduct(reactionObj, speciesObj)
```

Arguments

<i>reactionObj</i>	Reaction object.
<i>SpeciesName</i>	Name for a model object. Enter a species name or cell array of species names.
<i>speciesObj</i>	Species object. Enter a species object or an array of species objects.

Description

`rmproduct(reactionObj, SpeciesName)`, in a reaction object (`reactionObj`), removes a species object with a specified name (`SpeciesName`) from the property `Products`, removes the species name from the property `Reaction`, and updates the property `Stoichiometry` to exclude the species coefficient.

`rmproduct(reactionObj, speciesObj)` removes a species object as described above using a MATLAB variable for a species object.

The species object is not removed from the parent model property `Species`. If the species object is no longer used by any reaction, you can use the function `delete` to remove it from the parent object.

If one of the species specified does not exist as a product, a warning will be returned.

Examples

Example 1

Shows you how to remove a product that was added to a reaction by mistake. You can remove the species object using the species name.

```
modelObj = sbiomodel('cell');  
reactionObj = addreaction(modelObj, 'Phosphocreatine + ADP -> creatine + ATP + Pi');  
rmproduct(reactionObj, 'Pi')
```

rmproduct (reaction)

Reaction Object Array

Index: Reaction:
1 Phosphocreatine + ADP -> creatine + ATP

Example 2

Remove a species object using a model index to a species object.

```
modelObj = sbiomodel('cell');
reactionObj = addreaction(modelObj, 'A -> B + C');
reactionObj.Reaction

ans =
A -> B + C

rmproduct(reactionObj, modelObj.Species(2));
reactionObj.Reaction

ans =
A -> C
```

See Also

[rmreactant](#)

Purpose

Remove species object from reaction object reactants

Syntax

```
rmreactant(reactionObj, SpeciesName)
rmreactant(reactionObj, speciesObj)
```

Arguments

<i>reactionObj</i>	Reaction object.
<i>SpeciesName</i>	Name for a species object. Enter a species name or cell array of species names.
<i>speciesObj</i>	Species object. Enter a species object or an array of species objects.

Description

`rmreactant(reactionObj, SpeciesName)`, in a reaction object (*reactionObj*), removes a species object with a specified name (*SpeciesName*) from the property `Reactants`, removes the species name from the property `Reaction`, and updates the property `Stoichiometry` to exclude the species coefficient.

`rmreactant(reactionObj, speciesObj)` removes a species object as described above using a MATLAB variable for a species object, or a model index for a species object.

The species object is not removed from the parent model property `Species`. If the species object is no longer used by any reaction, you can use the method, `delete` to remove it from the parent object.

If one of the species specified does not exist as a reactant, a warning is returned.

Examples

Example 1

Shows you how to remove a reactant that was added to a reaction by mistake. You can remove the species object using the species name.

```
modelObj = sbiomodel('cell');
reactionObj = addreaction(modelObj, 'Phosphocreatine + ADP + Pi -> creatine + ATP');
rmreactant(reactionObj, 'Pi')
```

rmreactant (reaction)

Reaction Object Array

Index: Reaction:
1 Phosphocreatine + ADP -> creatine + ATP

Example 2

Remove a species object using a model index to a species object.

```
modelObj = sbiomodel('cell');
reactionObj = addreaction(modelObj, 'A -> B + C');

reactionObj.Reaction

ans =
A + B -> C

rmreactant(r, m.Species(2));
reactionObj.Reaction

ans =
A -> C
```

See Also

[rmproduct](#), [delete](#)

Purpose

Set the active configuration set for model object

Syntax

```
configsetObj = setactiveconfigset(modelObj,  
    'NameValue')  
configsetObj2 = setactiveconfigset(modelObj, configsetObj1)
```

Description

configsetObj = *setactiveconfigset(modelObj, 'NameValue')* sets the configuration set *NameValue* to be the active configuration set for the model *modelObj* and returns to *configsetObj*.

configsetObj2 = *setactiveconfigset(modelObj, configsetObj1)* sets the configset *configsetObj1* to be the active configset for *modelObj* and returns to *configsetObj2*. Any change in one of these two configset objects *configsetObj1* and *configsetObj2* is reflected in the other. To copy over a configset object from one model object to another use the *copyobj* method.

The active configuration set contains the settings that are be used during a simulation. A default configuration set is attached to any new model.

Examples

- 1 Create a model object by importing the file *oscillator.xml* and add a configset that simulates for 3000 seconds.

```
modelObj = sbmlimport('oscillator');  
configsetObj = addconfigset(modelObj, 'myset');
```

- 2 Configure the *configsetObj* *StopTime* to 3000.

```
set(configsetObj, 'StopTime', 3000)  
get(configsetObj)
```

```
Active: 0  
CompileOptions: [1x1 SimBiology.CompileOptions]  
Name: 'myset'  
Notes: ''
```

setactiveconfigset (model)

```
RuntimeOptions: [1x1 SimBiology.RuntimeOptions]
SolverOptions: [1x1 SimBiology.ODESolverOptions]
SolverType: 'ode15s'
StopTime: 3000
StopTimeType: 'simulationTime'
TimeUnits: 'second'
Type: 'configset'
```

- 3** Set the new configset to be active, simulate the model using the new configset and plot the result

```
setactiveconfigset(modelObj, configsetObj);
[t,x] = sbiosimulate(modelObj);
plot (t,x)
```

See Also

[addconfigset](#), [getconfigset](#), [removeconfigset](#)

Purpose

Specify specific parameters in kinetic law object

Syntax

```
setparameter(kineticlawObj,  
'ParameterVariablesValue', 'ParameterVariableNamesValue')
```

Arguments

ParameterVariableValue

Specify value of parameter variable in kinetic law object.

ParameterVariableNamesValue

Specify the parameter name with which to configure parameter variable in kinetic law object. Determines parameters in ReactionRate equation.

Description

Configure ParameterVariableNames in kinetic law object.

`setparameter(kineticlawObj, 'ParameterVariablesValue', 'ParameterVariableNamesValue')` configures the `ParameterVariableNames` property of the kinetic law object (`kineticlawObj`). `ParameterVariableValue` corresponds to one of the strings in `kineticlawObj` `ParameterVariables` property. The corresponding element in `kineticlawObj``ParameterVariableNames` property is configured to `ParameterVariableNamesValue`. For example, if `ParameterVariables` is `{'Vm', 'Km'}` and `ParameterVariablesValue` is specified as `Vm`, then the first element of the `ParameterVariableNames` cell array is configured to `ParameterVariableNamesValue`.

Example

Create a model, add a reaction, and assign the `ParameterVariableNames` for the reaction rate equation.

- 1 Create model object, and then add a reaction object.

```
modelObj = sbiomodel('my_model');  
reactionObj = addreaction(modelObj, 'a -> c + d');
```

setparameter (kineticlaw)

- 2** Create a kinetic law object for the reaction object, of the type 'Henri-Michaelis-Menten'.

```
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');
```

reactionObj KineticLaw property is configured to kineticlawObj.

- 3** The 'Henri-Michaelis-Menten' kinetic law has two parameter variables (V_m and K_m) that should be set. To set these variables,

```
setparameter(kineticlawObj, 'Vm', 'Va');  
setparameter(kineticlawObj, 'Km', 'Ka');
```

- 4** Verify that the parameter variables are correct.

```
get (kineticlawObj, 'ParameterVariableNames')
```

MATLAB returns

```
ans =
```

```
'Va'      'Ka'
```

See Also

[addparameter](#), [getspecies](#), [setspecies](#)

Purpose	Specify species in kinetic law object				
Syntax	<pre>setspecies(kineticlawObj, 'SpeciesVariablesValue', 'SpeciesVariableNamesValue')</pre>				
Arguments	<table><tr><td><i>SpeciesVariablesValue</i></td><td>Specify species variable in kinetic law object.</td></tr><tr><td><i>SpeciesVariableNamesValue</i></td><td>Specify the species name with which to configure species variable in kinetic law object. Determines species in ReactionRate equation</td></tr></table>	<i>SpeciesVariablesValue</i>	Specify species variable in kinetic law object.	<i>SpeciesVariableNamesValue</i>	Specify the species name with which to configure species variable in kinetic law object. Determines species in ReactionRate equation
<i>SpeciesVariablesValue</i>	Specify species variable in kinetic law object.				
<i>SpeciesVariableNamesValue</i>	Specify the species name with which to configure species variable in kinetic law object. Determines species in ReactionRate equation				
Description	<p><code>setspecies</code> configures kinetic law object <code>SpeciesVariableNames</code> property.</p> <p><code>setspecies(kineticlawObj, 'SpeciesVariablesValue', 'SpeciesVariableNamesValue')</code> configures the <code>SpeciesVariableNames</code> property of the kinetic law object, <code>KineticlawObj</code>. <code>SpeciesVariablesValue</code> corresponds to one of the strings in <code>SpeciesVariables</code> property of <code>kineticlawObj</code>. The corresponding element in <code>kineticlawObj</code> <code>SpeciesVariableNames</code> property is configured to <code>SpeciesVariableNamesValue</code>.</p> <p>For example, if <code>SpeciesVariables</code> are <code>{'S', 'S1'}</code> and <code>SpeciesVariablesValue</code> is specified as <code>S1</code>, the first element of the <code>SpeciesVariableNames</code> cell array is configured to <code>SpeciesVariableNamesValue</code>.</p>				
Example	Create a model, add a reaction and assign the <code>SpeciesVariableNames</code> for the reaction rate equation.				

- 1 Create model object, and then add a reaction object.

```
modelObj = sbiomodel('my_model');
reactionObj = addreaction(modelObj, 'a -> c + d');
```

setspecies (kineticlaw)

- 2** Create a kinetic law object for the reaction object, of the type 'Henri-Michaelis-Menten'.

```
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');
```

reactionObj KineticLaw property is configured to kineticlawObj.

- 3** The 'Henri-Michaelis-Menten' kinetic law has one species variable (S) that should be set. To set this variable,

```
setspecies(kineticlawObj, 'S', 'a');
```

- 4** Verify that the species variable is correct.

```
get (kineticlawObj, 'SpeciesVariableNames')
```

MATLAB returns

```
ans =  
'a'
```

See Also

[addparameter](#), [getspecies](#), [setparameter](#)

Purpose

Validate and verify SimBiology model

Syntax

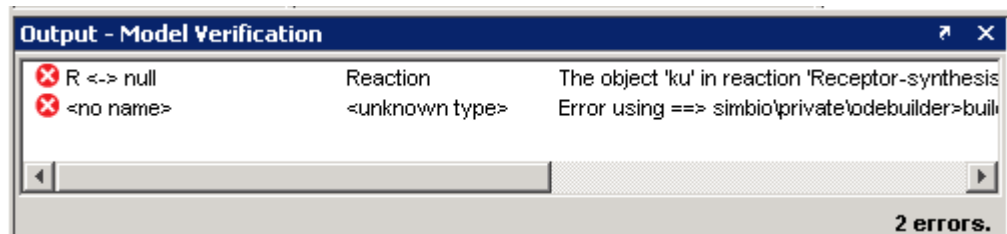
```
verify(modelObj)
```

Description

`verify(modelObj)` performs checks on a model object (*modelObj*) to verify that you can simulate the model. SimBiology generates stacked errors and warnings if any problems are found. To see the entire list of errors and warnings, use `sbiolasterror` and `sbiolastwarning`.

Verification in the SimBiology GUI

While you are building your model in the SimBiology desktop you can click  at any time to generate a list of any errors and warnings in the model. The errors and warnings appear in the **Output** pane. Following is an example of the error generated when the reaction rate of a reaction is set to a parameter that you have not defined in SimBiology.



Double-click the error row to move to the location of the error.

Examples

```
m = sbmlimport('radiodecay.xml');  
verify(m);
```

See Also

`sbiolasterror`, `sbiolastwarning`

Properties — By Category

Abstract Kinetic Law (p. 5-2)	Properties for abstract kinetic law objects
Configuration Sets (p. 5-3)	Properties for configuration set objects
Kinetic Laws (p. 5-4)	Properties for kinetic law objects.
Models (p. 5-5)	Properties for model objects
Parameters (p. 5-6)	Properties for parameter objects
Reactions (p. 5-7)	Properties for reaction objects
Root (p. 5-8)	Properties for the root object
Rules (p. 5-9)	Properties for rule objects
Species (p. 5-10)	Properties for species objects
Using Object Properties (p. 5-11)	Command-line syntax for entering and retrieving property values.

Abstract Kinetic Law

Annotation	Property with information about a SimBiology object
Expression	Property containing the expression used to determine the reaction rate equation
Name	Property with name of object
Notes	Property with HTML text describing SimBiology object
ParameterVariables	Property showing parameters in abstract kinetic law
Parent	Property indicating the parent object
SpeciesVariables	Property showing species in abstract kinetic law
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Configuration Sets

Active	Property to indicate object use during a simulation
CompileOptions	Property holding dimensional analysis and unit conversion information
Name	Property with name of object
Notes	Property with HTML text describing SimBiology object
RuntimeOptions	Property holding options for logged species
SensitivityAnalysisOptions	Hold sensitivity analysis options
SolverOptions	Property holding the model solver options
SolverType	Property to select solver type for simulation
StopTime	Property to set the stop time for a simulation
StopTimeType	Property to specify the type of stop time for a simulation
TimeUnits	Property to show the stop time units for a simulation
Type	Property to indicate SimBiology object type

Kinetic Laws

Annotation	Property with information about a SimBiology object
Expression	Property containing the expression used to determine the reaction rate equation
KineticLawName	Property showing name of abstract kinetic law
Name	Property with name of object
Notes	Property with HTML text describing SimBiology object
Parameters	Property with array of parameter objects
ParameterVariableNames	Property showing cell array of reaction rate parameters
ParameterVariables	Property showing parameters in abstract kinetic law
Parent	Property indicating the parent object
SpeciesVariableNames	Property showing cell array of species used in reaction rate equation
SpeciesVariables	Property showing species in abstract kinetic law
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Models

Annotation	Property with information about a SimBiology object
Models	Property showing all model objects
Name	Property with name of object
Notes	Property with HTML text describing SimBiology object
Parameters	Property with array of parameter objects
Parent	Property indicating the parent object
Reactions	Property with an array of reaction objects
Rules	Property showing rules in model object
Species	Property showing species in model object
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Parameters

Annotation	Property with information about a SimBiology object
ConstantValue	Property to indicate variable or constant parameter value
Name	Property with name of object
Notes	Property with HTML text describing SimBiology object
Parent	Property indicating the parent object
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object
Value	Property to assign value to parameter object
ValueUnits	Property with parameter value units

Reactions

Active	Property to indicate object use during a simulation
Annotation	Property with information about a SimBiology object
KineticLaw	Property showing kinetic law for ReactionRate
Name	Property with name of object
Notes	Property with HTML text describing SimBiology object
Parent	Property indicating the parent object
Products	Property to indicate reaction products
Reactants	Property to indicate reaction reactants
Reaction	Property to indicate the reaction object reaction
ReactionRate	Property containing the reaction rate equation in reaction object
Reversible	Property to indicate whether a reaction is reversible or irreversible
Stoichiometry	Property that describes species coefficients in a reaction
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Root

BuiltInKineticLaws	Property containing built-in kinetic laws
BuiltInUnitPrefixes	Property containing built-in unit prefixes
BuiltInUnits	Property containing built-in units
Models	Property showing all model objects
Type	Property to indicate SimBiology object type
UserDefinedKineticLaws	Property containing user-defined kinetic laws
UserDefinedUnitPrefixes	Property containing user-defined unit prefixes
UserDefinedUnits	Property containing user-defined units

Rules

Active	Property to indicate object use during a simulation
Annotation	Property with information about a SimBiology object
Name	Property with name of object
Notes	Property with HTML text describing SimBiology object
Parent	Property indicating the parent object
Rule	Property to define certain species and parameter interactions
RuleType	Property for defining the type of rule for the rule object
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Species

Annotation	Property with information about a SimBiology object
BoundaryCondition	Property to set a species object to have a boundary condition
ConstantAmount	Property to specify variable or constant species amount
InitialAmount	Property containing initial amount of a species
InitialAmountUnits	Property containing units for species initial amount
Name	Property with name of object
Notes	Property with HTML text describing SimBiology object
Parent	Property indicating the parent object
Tag	Property to specify a label for a SimBiology object
Type	Property to indicate SimBiology object type
UserData	Property to specify data to associate with object

Using Object Properties

Command-line syntax for entering and retrieving property values.

Entering property values (p. 5-11)	Use either MATLAB functions or object dot notation to enter or change property values.
Retrieving property values (p. 5-11)	Use either MATLAB functions or object dot notation to get property values.
Help for Objects, Methods and Properties (p. 5-12)	Use the command <code>sbiohelp</code> to get information about properties.

Entering property values

Enter or change a single property value using dot notation.

```
ObjectName.PropertyName = PropertyValue
```

Enter or change one or more property values using the MATLAB function set.

```
set(ObjectName, 'PropertyName', PropertyValue, ...)
```

Retrieving property values

Retrieve a single property value using dot notation.

```
PropertyValue = ObjectName.PropertyName
```

Retrieve one or more property values using the MATLAB function get.

```
PropertyValue(s) = get(ObjectName, 'PropertyName', ...)
```

Retrieve one or more property values using the object method get.

```
PropertyValue(s) = ObjectName.get('PropertyName', ...)
```

List or retrieve all property values using one of the following commands.

```
get(ObjectName)
AllPropertyValues = get(ObjectName)
```

ObjectName.get

Help for Objects, Methods and Properties

Display information for SimBiology object methods and properties in the MATLAB Command Window.

<code>help sbio</code>	Display a list of functions and methods.
<code>help FunctionName</code>	Display function information.
<code>sbiohelp('MethodName')</code>	Display method information.
<code>sbiohelp('PropertyName')</code>	Display property information.

Properties — Alphabetical List

AbsoluteTolerance

Purpose

Property to specify largest allowable absolute error

Description

AbsoluteTolerance specifies the largest allowable absolute error at any step in simulation. It is a property of SolverOptions object. SolverOptions is a property of the configset object. AbsoluteTolerance is available for the ode solvers ('ode45', 'ode23', 'ode113', 'ode15s', 'ode23s', 'ode23t', and 'ode23tb').

At each simulation step, the solver estimates the local error e_i in the i^{th} state vector y . Simulation converges at that time step if e_i satisfies the following equation:

$$|e_i| \leq \max(\text{RelativeTolerance} * |y_i|, \text{AbsoluteTolerance})$$

Thus at higher state values, convergence is determined by RelativeTolerance. As the state values approach zero, convergence is controlled by AbsoluteTolerance. The choice of values for RelativeTolerance and AbsoluteTolerance will vary depending on the problem. The default values should work for first trials of the simulation; however if you want to optimize the solution, consider that there is a trade-off between speed and accuracy. If the simulation takes too long, you can increase the values of RelativeTolerance and AbsoluteTolerance at the cost of some accuracy. If the results appear to be inaccurate you can decrease the tolerance values but this will slow down the solver. If the magnitude of the state values is high, you can try to decrease the relative tolerance to get more accurate results.

This may be important for reactions where species values tend to zero. Even if you are not interested in the value of a state $y(i)$ when it is small, you may have to specify AbsoluteTolerance small enough to get some correct digits in $y(i)$ so that you can accurately compute more interesting state values.

Characteristics

Applies to

Object: SolverOptions

Data type

double

Data values	>0, <1; default is 1e-6
Access	Read/Write

Example

This example shows how to change AbsoluteTolerance.

- 1 Retrieve the configset object from the modelObj.

```
modelObj = sbiomodel('cell');
configsetObj = getconfigset(modelObj)
```

- 2 Change the AbsoluteTolerance to 1e-8.

```
set(configsetObj.SolverOptions, 'AbsoluteTolerance', 1.0e-8)
get(configsetObj.SolverOptions, 'AbsoluteTolerance')

ans =
1.0000e-008
```

See Also

[RelativeTolerance](#)

Active

Purpose	Property to indicate object use during a simulation
Description	Indicates whether a simulation is using a SimBiology object. A SimBiology model is organized into a hierarchical group of objects. Use the Active property to include or exclude objects during a simulation. When a reaction or rule object Active property is set to be <code>false</code> , the simulation does not include the reaction or rule. This is a convenient way to test a model with and without a reaction or rule. For configset object, use the method <code>setactiveconfigset</code> , to set the object Active property to <code>true</code> .

Characteristics

Applies to	Objects: configset, reaction, rule
Data type	<code>boolean</code>
Data values	<code>true</code> or <code>false</code> . Default value is <code>true</code> . For default configset object default is <code>true</code> , for added configset object default is <code>false</code> .
Access	Read/Write

Example

- 1 Create a model object.

```
modelObj = sbiomodel ('my_model');
```

- 2 Add reaction object and verify that the Active property setting is '`true`' or 1.

```
reactionObj = addreaction (modelObj, 'a + b -> c + d');
get (reactionObj, 'Active')
```

MATLAB returns

```
ans =
```

```
1
```

- 3 Set Active property to '`false`' and verify.

```
set (reactionObj, 'Active', false);
get (reactionObj, 'Active')
```

MATLAB returns

```
ans =
0
```

See Also

[addrule](#), [addreaction](#), [setactiveconfigset](#), [addconfigset](#)

Annotation

Purpose Property with information about a SimBiology object

Description URL or filename linking to information about a model.

Characteristics

Applies to	Objects: kineticlaw, model, parameter, reaction, root, rule, species
Data type	char string, URL
Data values	Character string with a directory path and filename or a URL.
Access	Read/Write

Example 1 Create a model object

```
modelObj = sbiomodel ('my_model');
```

2 Set annotation for model object

```
set (modelObj, 'annotation', 'www.reactome.org')
```

3 Verify the assignment.

```
get (modelObj, 'annotation')
```

MATLAB returns

```
ans =
```

```
www.reactome.org
```

See Also sbiomodel, addkineticlaw, addparameter, addreaction, addrule, addspecies, sbioroot

Purpose

Property to set a species object to have a boundary condition

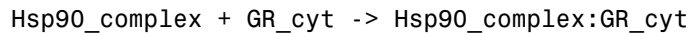
Description

Indicates whether a species object has a boundary condition. If BoundaryCondition is true, the species quantity is determined by InitialAmount and/or a rule object, and not by the reaction rate equation. In Simbiology, all species are state variables regardless of BoundaryCondition or ConstantAmount property.

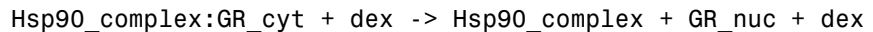
By default BoundaryCondition is false and Simbiology uses reaction rate equations to determine the rate of change of a species quantity in the model. Boundary condition is used when a species is modeled as a participant of reactions but the species quantity is not determined by a reaction rate equation. Consider the following two use cases of boundary conditions:

- Modeling receptor-ligand interactions that affect the rate of change of the receptor but not the ligand. For example, in response to hormone, steroid receptors such as the glucocorticoid receptor (GR) translocate from the cytoplasm (cyt) to the nucleus (nuc). The hsp90/hsp70 chaperone complex directs this nuclear translocation [Pratt 2004]. The natural ligand for GR is cortisol; the synthetic hormone dexamethasone (dex) is used in place of cortisol in experimental systems. In this system dexamethasone participates in the reaction but the quantity of dexamethasone in the cell is regulated using a rule. To simply model translocation of GR you could use the following reactions:

Formation of the chaperone–receptor complex,



In response to the synthetic hormone dexamethasone (dex), GR moves from the cytoplasm to the nucleus.



For dex,

```
BoundaryCondition = true; ConstantAmount = false
```

BoundaryCondition

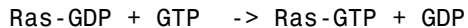
In this example dex is modeled as a boundary condition with a rule to regulate the rate of change of dex in the system. Here, the quantity of dex is not determined by the rate of the second reaction but by a rate rule such as

$$\frac{ddex}{dt} = 0.001$$

which is specified in SimBiology as

$$dex = 0.001$$

- Modeling the role of nucleotides (for example, GTP, ATP, cAMP) and cofactors (for example, Ca^{++} , NAD^+ , coenzyme A). Consider the role of GTP in the activation of Ras by receptor tyrosine kinases.



For GTP, BoundaryCondition = true; ConstantAmount = true

Model GTP and GDP with boundary conditions, thus making them *boundary species*. In addition you can set the ConstantAmount property of these species to true to indicate that their quantity does not vary during a simulation.

Characteristics

Applies to	Object: species
Data type	boolean
Data values	true or false. The default value is false.
Access	Read/Write

Example

- 1 Create a model object

```
modelObj = sbiomodel ('my_model');
```

- 2** Add a species object and verify that boundary condition property setting is 'false' or 0.

```
speciesObj = addspecies(modelObj, 'glucose');
get(speciesObj, 'BoundaryCondition')
```

MATLAB returns

```
ans =
0
```

- 3** Set boundary condition to 'true' and verify

```
set(speciesObj, 'BoundaryCondition', true);
get(speciesObj, 'BoundaryCondition')
```

MATLAB returns

```
ans =
1
```

References

Pratt, W.B., Galigniana, M.D., Morishima, Y., Murphy, P.J. (2004), Role of molecular chaperones in steroid receptor action, *Essays Biochem*, 40:41-58.

See Also

addrule, addspecies, ConstantAmount, InitialAmount

BuiltInKineticLaws

Purpose Property containing built-in kinetic laws

Description BuiltInKineticLaws is a SimBiology root object property showing all abstract kinetic laws that are shipped with SimBiology. Use the command `sbiowhos -builtin -kineticlaw` to see the list of built-in kinetic laws. You can use built-in kinetic laws when you use the command `addkineticlaw` to create a kinetic law object for a reaction object. Refer to the kinetic law by name when you create the kinetic law object, for example:

```
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');
```

You cannot add, modify, or delete BuiltInKineticLaws.

See “Abstract Kinetic Law” on page 6-27 for a definition and more information.

Characteristics

Applies to	Object: root
Data type	char string of valid abstract kinetic law name.
Data values	Valid kinetic laws
Access	Read-only

Examples

Example 1

This example uses the command `sbiowhos` to show the current list of built-in kinetic laws.

```
sbiowhos -builtin -kineticlaw
```

Abstract Kinetic Law Object Array

Index:	Library:	Name:	Expression:
1	BuiltIn	Unknown	Unknown
2	BuiltIn	MassAction	MassAction

3	BuiltIn	Henri-Michaelis-Menten	$V_m * S / (K_m + S)$
---	---------	------------------------	-----------------------

Example 2

This example shows the current list of built-in kinetic laws by accessing the root object.

```
rootObj = sbioroot;
get(rootObj, 'BuiltInKineticLaws')
```

Abstract Kinetic Law Object Array

Index:	Library:	Name:	Expression:
1	BuiltIn	Unknown	Unknown
2	BuiltIn	MassAction	MassAction
3	BuiltIn	Henri-Michaelis-Menten	$V_m * S / (K_m + S)$

See Also

[UserDefinedKineticLaws](#), [BuiltInUnits](#), [BuiltInUnitPrefixes](#)
MATLAB functions `get` and `set`

BuiltInUnitPrefixes

Purpose Property containing built-in unit prefixes

Description BuiltInUnitPrefixes is a SimBiology root object property showing all unit prefixes that are shipped with SimBiology. You can specify units with prefixes for species amounts and parameter values, because, SimBiology enables you to do dimensional analysis and unit conversion during simulation. The valid units and unit prefixes are either built-in or user-defined. You can display the built-in unit prefixes either by using the command sbiowhos, or by accessing the root object. Both methods are illustrated in the examples below.

You cannot add, modify, or delete BuiltInUnitsPrefixes.

Characteristics

Applies to	Object: root
Data type	char string
Data values	Valid units
Access	Read-only

Examples

Example 1

This example uses the command sbiowhos to show the current list of built-in unit prefixes.

```
sbiowhos -builtin -unitprefix
```

Example 2

This example shows the current list of built-in unit prefixes by accessing the root object.

```
rootObj = sbioroot;
get(rootObj, 'BuiltInUnitPrefixes')
```

See Also

[BuiltInUnits](#), [UserDefinedUnits](#), [BuiltInKineticLaws](#)

MATLAB functions `get` and `set`.

BuiltInUnits

Purpose Property containing built-in units

Description BuiltInUnits is a SimBiology root object property showing all units that are shipped with SimBiology. You can specify units for species amounts and parameter values, because, SimBiology enables you to do dimensional analysis and unit conversion during simulation. The valid units are either built-in or user-defined. You can display the built-in units either by using the command sbioswhos, or by accessing the root object. Both methods are illustrated in the examples below.

You cannot add, modify, or delete BuiltInUnits.

Characteristics

Applies to	Object: root
Data type	char string
Data values	Valid units.
Access	Read-only

Examples

Example 1

This example shows the current list of built-in units using the sbiowhos command.

```
sbiowhos -builtin -units
```

Example 2

This example shows the current list of built-in units by accessing the root object.

```
rootObj = sbioroot;
get(rootObj, 'BuiltInUnits')
```

See Also

[BuiltInUnitPrefixes](#), [UserDefinedUnits](#), [BuiltInKineticLaws](#)

MATLAB functions `get` and `set`.

CompileOptions

Purpose Property holding dimensional analysis and unit conversion information

Description The SimBiology `CompileOptions` object defines the compile options available for simulation; you can specify whether dimensional analysis and unit conversion is necessary for simulation. Compile options are checked during compile time. The compile options object can be accessed through the `CompileOptions` property of the `configset` object. Retrieve `CompileOptions` object properties with the `get` function and configure the properties with the `set` function.

Property Summary	DimensionalAnalysis	Property to indicate whether to perform dimensional analysis
	Type	Property to indicate SimBiology object type
	UnitConversion	Indicate whether to perform unit conversion

Characteristics

Applies to	Object: <code>configset</code> object
Data type	Object
Data values	Compile time options
Access	Read-only

Example

1 Retrieve the `configset` object of `modelObj`

```
modelObj = sbiomodel('cell');
configsetObj = getconfigset(modelObj);
```

2 Retrieve the `CompileOptions` object (`optionsObj`) from the `configsetObj`

```
optionsObj = get(configsetObj, 'CompileOptions');
```

```
DimensionalAnalysis: 1
Type: 'compileoptions'
UnitConversion: 1
```

See Also

MATLAB functions `get`, `set`

ConstantAmount

Purpose

Property to specify variable or constant species amount

Description

ConstantAmount indicates whether the quantity of the species object can vary during the simulation. ConstantAmount can be either true or false. If ConstantAmount is true, the quantity of the species cannot vary during the simulation. By default, ConstantAmount is false and the quantity of the species can vary during the simulation. If ConstantAmount is false, the quantity of the species can be determined by reactions and rules.

The following is example of modeling species as constant amounts:

Modeling the role of nucleotides (GTP, ATP, cAMP) and cofactors (Ca^{++} , NAD^+ , coenzyme A). Consider the role of GTP in the activation of Ras by receptor tyrosine kinases.



Model GTP and GDP with constant amount set to true. In addition, you can set the BoundaryCondition of these species to true, thus making them *boundary species*.

The property ConstantAmount is for species objects; the property ConstantValue is for parameter objects.

Characteristics

Applies to	Object: species
Data type	boolean
Data values	true or false. The default value is false.
Access	Read/Write

Example

- 1 Create a model object with name my_model.

```
modelObj = sbiomodel ('my_model');
```

- 2 Add a species object and verify that the ConstantAmount property setting is 'false' or 0

```
speciesObj = addspecies (modelObj, 'glucose');
get (speciesObj, 'ConstantAmount')
```

MATLAB returns

```
ans =
```

```
0
```

3 Set constant amount to 'true' and verify

```
set (speciesObj, 'ConstantAmount', true);
get (speciesObj, 'ConstantAmount')
```

MATLAB returns

```
ans =
```

```
1
```

See Also

[addspecies](#), [BoundaryCondition](#)

ConstantValue

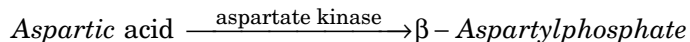
Purpose

Property to indicate variable or constant parameter value

Description

Indicates whether the value of a parameter can change during a simulation. Enter either true (value is constant) or false (value can change).

You can allow the value of the parameter to change during a simulation by specifying a rule that changes the Value property of the parameter object. For example, consider feedback inhibition of an enzyme such as aspartate kinase by threonine. Aspartate kinase has three isozymes that are independently inhibited by the products of downstream reactions (threonine, homoserine, and lysine). Although threonine is made through a series of reactions in the synthesis pathway, for illustration the reactions are simplified as follows:



To model inhibition of aspartate kinase by threonine you could use a rule like the algebraic rule below to vary the rate of the above reaction and simulate inhibition. In the rule, the rate constant for the above reaction is denoted by k_aspartate_kinase and the quantity of threonine is threonine.

```
k_aspartate_kinase - (1/threonine)
```

The property ConstantValue is for parameter objects; the property ConstantAmount is for species objects.

Characteristics

Applies to	Object: parameter
Data type	boolean
Data values	true or false. Default value is 'true'.
Access	Read/Write

Example

- 1 Create a model object.

```
modelObj = sbiomodel ('my_model');
```

- 2 Add parameter object.

```
parameterObj = addparameter (modelObj, 'kf');
```

- 3 Change the ConstantValue property of the parameter object from default (true) to false and verify.

MATLAB returns 1 for true and 0 for false.

```
set (parameterObj, 'ConstantValue', false)
get(parameterObj, 'ConstantValue')
```

MATLAB returns

```
ans =
```

```
0
```

See Also

[addparameter](#)

DimensionalAnalysis

Purpose

Property to indicate whether to perform dimensional analysis

Description

DimensionalAnalysis specifies whether to perform dimensional analysis on the model before simulation. It is a property of the CompileOptions object. CompileOptions holds the model's compile time options and is the object property of the configset object. When DimensionalAnalysis is set to true, SimBiology checks whether the physical quantities of the units involved in reactions and rules, match and are applicable.

For example, consider a reaction $a + b \rightarrow c$. Using mass action kinetics, the reaction rate is defined as $a*b*k$ where k is the rate constant of the reaction. If you specify that initial amounts of a and b are 0.01M and 0.005M respectively, then units of k are $1/(M\cdot second)$. If you specify k with another equivalent unit definition, for example $1/[(moles/liter)\cdot second]$, DimensionalAnalysis checks whether the physical quantities match. If the physical quantities do not match, you see an error and the model is not simulated, UnitConversion is the next step after DimensionalAnalysis.

Valid physical quantities for reaction rates are amount/time, mass/time or concentration/time.

Characteristics

Applies to	Object: CompileOptions (in configset object)
Data type	boolean
Data values	true or false. Default value is true.
Access	Read/Write

Example

Shows how to retrieve and set DimensionalAnalysis from the default true to false in the default configuration set in a model object.

- 1 Import a model.

```
modelObj = sbmlimport('oscillator')
```

```
SimBiology Model - Oscillator
```

Model Components:

Models:	0
Parameters:	0
Reactions:	42
Rules:	0
Species:	23

2 Retrieve the configset object of the model object.

```
configsetObj = getconfigset(modelObj)

Configuration Settings - default (active)
    SolverType:          ode15s
    StopTime:            10.000000

    SolverOptions:
        AbsoluteTolerance: 1.000000e-006
        RelativeTolerance: 1.000000e-003

    RuntimeOptions:
        StatesToLog:       all

    CompileOptions:
        UnitConversion:    true
        DimensionalAnalysis: true
```

3 Retrieve the CompileOptions object.

```
optionsObj = get(configsetObj,'CompileOptions')

Compile Settings:
    UnitConversion:    true
    DimensionalAnalysis: true
```

4 Assign a value of false to DimensionalAnalysis.

Dimensional Analysis

```
set(optionsObj,'DimensionalAnalysis' false)
```

See Also

[getconfigset](#), [sbiosimulate](#)
MATLAB functions `get` and `set`.

Purpose

Property specifies explicit or implicit tau error tolerance

Description

ErrorTolerance specifies the error tolerance for the explicit tau and implicit tau stochastic solvers. It is a property of the SolverOptions object. SolverOptions is a property of the configset object. The explicit and implicit tau solvers automatically chooses a time interval (tau) such that the relative change in the propensity function for each reaction is less than the user-specified error tolerance.

A propensity function describes the probability that the reaction will occur in the next smallest time interval, given the conditions and constraints.

If the error tolerance is too large, there may not be a solution to the problem and that could lead an error. If the error tolerance is small, the solver will take more steps than when the error tolerance is large leading to longer simulation times. The error tolerance should be adjusted depending upon the problem, but a good value for the error tolerance is between 1 % to 5 %.

Characteristics

Applies to	Object: SolverOptions
Data type	double
Data values	>0, <1; default is 3e -2
Access	Read/Write

Example

Shows how to change ErrorTolerance settings.

- 1 Retrieve the configset object from the modelObj and change the SolverType to expltau.

```
modelObj = sbiomodel('cell');
configsetObj = getconfigset(modelObj)
set(configsetObj, 'SolverType', 'expltau')
```

- 2 Change the ErrorTolerance to 1e-8.

ErrorTolerance

```
set(configsetObj.SolverOptions, 'ErrorTolerance', 5.0e-2)
get(configsetObj.SolverOptions, 'ErrorTolerance')

ans =
5.000000e-002
```

See Also

[LogDecimation](#), [RandomState](#)

Purpose	Property containing the expression used to determine the reaction rate equation
Description	Indicates the expression that is used to determine the <code>ReactionRate</code> property of the reaction object. <code>Expression</code> is a reaction rate expression assigned by the abstract kinetic law used by the kinetic law object. The abstract kinetic law being used is indicated by the property <code>KineticLawName</code> . You can configure <code>Expression</code> for user-defined abstract kinetic laws but not for builtin abstract kinetic laws. <code>Expression</code> is read-only for kinetic law objects.

Abstract Kinetic Law

The **abstract kinetic law** provides a mechanism for applying a specific rate law to multiple reactions. It acts as a mapping template for the reaction rate. The abstract kinetic law is defined by a reaction rate expression, which is defined in the property `Expression`, and the species and parameter variables used in the expression. The species variables are defined in the `SpeciesVariables` property, and the parameter variables are defined in the `ParameterVariables` property of the kinetic law object.

If a reaction is using an abstract kinetic law, the `ReactionRate` property of the reaction object shows the result of a mapping from an abstract kinetic law. To determine `ReactionRate` the species variables and parameter variables that participate in the reaction rate should be clearly mapped in the kinetic law for the reaction. In this case SimBiology determines the `ReactionRate` by using the `Expression` property of the abstract kinetic law object, and by mapping `SpeciesVariableNames` to `SpeciesVariables` and `ParameterVariableNames` to `ParameterVariables`.

For example, the abstract kinetic law `Henri-Michaelis-Menten` has the `Expression` $V_m \cdot [S] / (K_m + [S])$, where V_m and K_m are defined as parameters in the `ParameterVariables` property of the abstract kinetic law object, and S is defined as a species in the `SpeciesVariable` property of the abstract kinetic law object.

Expression

By applying the abstract kinetic law Henri-Michaelis-Menten to a reaction $A \rightarrow B$ with V_a mapping to V_m and A mapping to S the rate equation for the reaction becomes $V_a * [A] / (K_a + [A])$.

The exact expression of a reaction using MassAction kinetic law varies depending upon the number of reactants. Thus, for mass action kinetics the Expression property is set to MassAction because In general for mass action kinetics the reaction rate is defined as

$$r = k \prod_{i=1}^{n_r} [S_i]^{m_i}$$

where $[S_i]$ is the concentration of the i^{th} reactant, m_i is the stoichiometric coefficient of $[S_i]$, n_r is the number of reactants and k is the mass action reaction rate constant.

SimBiology comes with some built-in kinetic laws. Users can also define their own abstract kinetic laws. To find the list of available kinetic laws, use the `sbiowhos -kineticlaw` command (`sbiowhos`). You can create an abstract kinetic law with the function `sbioabstractkineticlaw` and add it to the library using `sbioaddtolibrary`.

Characteristics

Applies to	Objects: kineticlaw, abstractkineticlaw
Data type	char string
Data values	Defined by abstract kinetic law
Access	Read-only in kinetic law object. Read/Write in user-defined abstract kinetic law.

Examples

Example with Henri-Michaelis-Menten kinetics

- 1 Create a model object, and add a reaction object to the model.

```
modelObj = sbiomodel ('my_model');
```

```
reactionObj = addreaction (modelObj, 'a + b -> c + d');
```

2 Define a kinetic law for the reaction object

```
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');
```

3 Verify that the Expression property for the kinetic law object is Henri-Michaelis-Menten

```
get (kineticlawObj, 'Expression')
```

MATLAB returns

```
ans =
```

```
Vm*S/(Km + S)
```

4 The 'Henri-Michaelis-Menten' kinetic law has two parameter variables (V_m and K_m) and one species variable (S) that you should set. To set these variables, first create the parameter variables as parameter objects (parameterObj1, parameterObj2) with names V_m_d , K_m_d , and assign the objects Parent property value to the kineticlawObj. The species object with Name,a is created when reactionObj is created and need not be redefined.

```
parameterObj1 = addparameter(kineticlawObj, 'Vm_d');  
parameterObj2 = addparameter(kineticlawObj, 'Km_d');
```

5 Set the variable names for the kinetic law object

```
set(kineticlawObj, 'ParameterVariableNames', {'Vm_d' 'Km_d'});  
set(kineticlawObj, 'SpeciesVariableNames', {'a'});
```

6 Verify that the reaction rate is expressed correctly in the reaction object ReactionRate property

```
get (reactionObj, 'ReactionRate')
```

MATLAB returns

Expression

```
ans =  
Vm_d*a/(Km_d+a)
```

Example with Mass Action kinetics.

- 1 Create a model object, then add a reaction object

```
modelObj = sbiomodel ('my_model');  
reactionObj = addreaction (modelObj, 'a + b -> c + d');
```

- 2 Define a kinetic law for the reaction object

```
kineticlawObj = addkineticlaw(reactionObj, 'MassAction');  
get(kineticlawObj, 'Expression')
```

MATLAB returns

```
ans =  
MassAction
```

- 3 Assign the rate constant for the reaction.

```
set (kineticlawObj, 'ParameterVariblenames', 'k');  
get (reactionObj, 'ReactionRate')
```

MATLAB returns

```
ans =  
k*a*b
```

See Also

Abstract and kinetic law object properties: `SpeciesVariables`, `ParameterVariables`

Kinetic law object properties: `KineticLawName`, `Parameters`, `SpeciesVariableNames`, `ParameterVariableNames`

Reaction object property: ReactionRate

Functions: sbioaddtolibrary, sbiowhos

InitialAmount

Purpose Property containing initial amount of a species

Description Indicates the initial quantity of the SimBiology species object. InitialAmount is the quantity of the species before the simulation starts.

Characteristics

Applies to	Object: species
Data type	double
Data values	Positive real number. Default value is 0.
Access	Read/Write

Example Add a species with name and value to a model object.

1 Create a model object with named my_model.

```
modelObj = sbiomodel ('my_model');
```

2 Add the species object with the name glucose.

```
speciesObj = addspecies (modelObj, 'glucose');
```

3 Set the initial amount to 100 and verify.

```
set (speciesObj, 'InitialAmount', 100);  
get (speciesObj, 'InitialAmount')
```

MATLAB returns

```
ans =
```

```
100
```

See Also addspecies, InitialAmountUnits

Purpose

Property containing units for species initial amount

Description

Indicates the unit definition for the InitialAmount property of a species object. InitialAmountUnits can be one of the builtin units. To get a list of the defined units use the sbioshowunits function. If InitialAmountUnits changes from one unit definition to another, the InitialAmount does not automatically convert to the new units. The sbioconvertunits function does this conversion. To add a user-defined unit to the list see sbioregisterunit.

Characteristics

Applies to	object: species
Data type	char string
Data values	unit from Units list; Default = '' (None)
Access	Read/Write

Example

- 1 Create a model object named my_model.

```
modelObj = sbiomodel ('my_model');
```

- 2 Add a species object with the name glucose.

```
speciesObj = addspecies (modelObj, 'glucose');
```

- 3 Set the initial amount to 100, InitialAmountUnits to molecule, and verify.

```
set (speciesObj, 'InitialAmount',100, ...
      'InitialAmountUnits', 'molecule');
get (speciesObj,'InitialAmountUnits')
```

MATLAB returns

```
ans =
```

InitialAmountUnits

molecule

See Also

[InitialAmount](#), [sbioshowunits](#), [sbioconvertunits](#),
[sbioregisterunit](#)

Purpose	Property showing kinetic law for ReactionRate								
Description	<p>KineticLaw defines the kinetics used to determine the reaction rate that is specified in the ReactionRate property of the reaction object. This property shows the kinetic law used to define ReactionRate.</p> <p>KineticLaw can be configured with the addkineticlaw method. The addkineticlaw function configures the ReactionRate based on the KineticLaw and the species and parameters specified in the kinetic law object properties SpeciesVariableNames and ParameterVariableNames. SpeciesVariableNames are determined automatically for mass action kinetics.</p> <p>If the reaction is updated, the ReactionRate is automatically updated only for mass action kinetics. For all other kinetics the SpeciesVariableNames property of the kinetic law object should be reconfigured.</p>								
Characteristics	<table border="0"> <tr> <td>Applies to</td><td>Object: reaction</td></tr> <tr> <td>Data type</td><td>Kinetic law object</td></tr> <tr> <td>Data values</td><td>Kinetic law object. Default is empty ([]). </td></tr> <tr> <td>Access</td><td>Read-only</td></tr> </table>	Applies to	Object: reaction	Data type	Kinetic law object	Data values	Kinetic law object. Default is empty ([]).	Access	Read-only
Applies to	Object: reaction								
Data type	Kinetic law object								
Data values	Kinetic law object. Default is empty ([]).								
Access	Read-only								
Example	<p>Example with Henri-Michaelis-Menten kinetics</p> <p>1 Create a model object, then add a reaction object</p> <pre>modelObj = sbiomodel ('my_model'); reactionObj = addreaction (modelObj, 'a + b -> c + d');</pre> <p>2 Define a kinetic law for the reaction object</p> <pre>kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');</pre>								

- 3 Verify that the KineticLaw property for the reaction object is Henri-Michaelis-Menten

```
get (reactionObj, 'KineticLaw')
```

MATLAB returns

```
Kinetic Law Object Array
```

Index:	KineticLawName:
1	Henri-Michaelis-Menten

See Also

Kinetic law object properties: KineticLawName, Parameters, SpeciesVariableNames, ParameterVariableNames

Reaction object property: ReactionRate

Purpose

Property showing name of abstract kinetic law

Description

Indicates the name of the abstract kinetic law in the kinetic law object. KineticLawName can be any valid name from the builtin or user-defined abstract kinetic law library. See “Abstract Kinetic Law” on page 6-27 for a definition and more information.

You can find the KineticLawName list in the abstract kinetic law library by using the command `sbiowhos -kineticlaw` (`sbiowhos`). You can create an abstract kinetic law with the function `sbioabstractkineticlaw` and add it to the library using `sbioaddtolibrary`.

Characteristics

Applies to	Object: kineticlaw
Data type	char string
Data values	char string defined by abstract kinetic law
Access	Read-only

Examples

- 1 Create a model object, add a reaction object, and define a kinetic law for the reaction object.

```
modelObj = sbiomodel ('my_model');
reactionObj = addreaction (modelObj, 'a + b -> c + d');
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');
```

- 2 Verify KineticLawName of kineticlawObj

```
get (kineticlawObj, 'KineticLawName')
```

MATLAB returns

```
ans =
```

```
Henri-Michaelis-Menten
```

KineticLawName

See Also

Abstract and kinetic law object properties: Expression,
SpeciesVariables, ParameterVariables

Kinetic law object properties: Parameters, SpeciesVariableNames,
ParameterVariableNames

Reaction object property: ReactionRate

Functions: sbioaddtolibrary, sbiowhos

Purpose

Property to specify recorded simulation output frequency

Description

LogDecimation defines how often the simulation data is recorded as output. It is a property of the SolverOptions object. SolverOptions is a property of the configset object. LogDecimation is available for ssa, expltau, and inmpltau solvers.

Use LogDecimation to specify how frequently you want to record the output of the simulation. For example, if the LogDecimation is set to 1, for the command `(t, x) = sbiosimulate(modelObj)`, at each simulation step the time will be logged in t and the quantity of each logged species will be logged as a row in x. If LogDecimation is 10, then every 10th simulation step will be logged in t and x.

Characteristics

Applies to	Object: SolverOptions
Data type	int
Data values	>0 default is 1.
Access	Read/Write

Example

Shows how to change LogDecimation settings.

- 1 Retrieve the configset object from the modelObj and change the SolverType to expltau

```
modelObj = sbiomodel('cell');
configsetObj = getconfigset(modelObj)
set(configsetObj, 'SolverType', 'expltau')
```

- 2 Change the LogDecimation to 10.

```
set(configsetObj.SolverOptions, 'LogDecimation', 10)
get(configsetObj.SolverOptions, 'LogDecimation')
```

LogDecimation

```
ans =
```

```
10
```

See Also

ErrorTolerance, RandomState

Purpose Property to specify nonlinear solver maximum iterations in implicit tau

Description MaxIterations specifies the maximum number of iterations for the nonlinear solver in impltau. It is a property of the SolverOptions object. SolverOptions is a property of the configset object.

The implicit tau solver in SimBiology internally uses a nonlinear solver to solve a set of algebraic nonlinear equations at every simulation step. Starting with an initial guess at the solution, the nonlinear solver iteratively tries to find the solution to the algebraic equations. The closer the initial guess is to the solution, the fewer the iterations the nonlinear solver will take before it finds a solution. MaxIterations specifies the maximum number of iterations the nonlinear solver should take before it issues a “failed to converge” error. If you get this error, during simulation try increasing MaxIterations. The default value of MaxIterations is 15.

Characteristics

Applies to	Object: SolverOptions
Data type	int
Data values	>0 default is 15.
Access	Read/Write

Example Shows how to change MaxIterations settings.

- 1 Retrieve the configset object from the modelObj and change the SolverType to impltau.

```
modelObj = sbiomodel('cell');
configsetObj = getconfigset(modelObj)
set(configsetObj, 'SolverType', 'impltau')
```

- 2 Change the MaxIterations to 25.

```
set(configsetObj.SolverOptions, 'MaxIterations', 25)
get(configsetObj.SolverOptions, 'MaxIterations')
```

MaxIterations

```
ans =
```

```
25
```

See Also

ErrorTolerance, LogDecimation, RandomState

Purpose Specify upper bound on solver step size

Description The MaxStep property specifies the size of the bounds on the size of the time steps. If the differential equation has periodic coefficients or solutions, it might be a good idea to set MaxStep to some fraction (such as 1/4) of the period. This guarantees that the solver does not enlarge the time step too much and step over a period of interest. For more information on MaxStep, see odeset in the MATLAB documentation.

Characteristics

Applies to	Object: SolverOptions
Data type	Positive scalar
Data values	{0.1*abs(t0-tf)} default is []
Access	Read/Write

See Also SimBiology property RelativeTolerance

MATLAB function odeset

Models

Purpose	Property showing all model objects
Description	<p>Indicates the models in a Model object or in the SimBiology root. Read-only array of Model objects. SimBiology has a hierarchical organization. A top-level model object has the SimBiology root as its Parent. Model objects with another model object as Parent are submodels. For a model object to access configset, kinetic law, reaction, rule and species objects, you must assign the model object as Parent in these objects. Parameter objects can have a model object or kinetic law object as Parent. You can display all the component objects with <code>modelObj.Models</code> or <code>get (modelObj, 'Models')</code>.</p> <p>The components of a submodel are contained within the submodel. In addition, a submodel object can reference parameter variables that have been assigned to the model object. For example, a parameter defined within a submodel cannot be used by the parent model or another model object. A submodel object however, can use the parameters assigned to the model object.</p> <p>You can add a submodel to a model object with the method <code>addmodel</code> and removed from its parent with the method <code>delete</code>.</p>
Characteristics	
Applies to	
Data type	
Data values	
Access	

Example

1 Create a model object

```
modelObj = sbiomodel ('cell');
```

2 Add submodels to model object and verify

```
submodelObj1 = addmodel (modelObj, 'nucleus');
submodelObj2 = addmodel (modelObj, 'mitochondrion');
```

```
get (modelObj, 'Models')
```

MATLAB returns

```
SimBiology Model Object Array
```

Index:	Name:
1	nucleus
2	mitochondrion

See Also

[sbiomodel](#), [addmodel](#)

Name

Purpose	Property with name of object
Description	<p>Identifies a SimBiology object. Species, parameter, and model objects can be referenced by other objects using the object property <code>Name</code>, therefore <code>Name</code> must be unique for these objects.</p> <p>Use the function <code>sbioselect</code> to find an object with the same <code>Name</code> property value.</p> <p>There are reserved characters that cannot be used in object names:</p> <ul style="list-style-type: none">• Models cannot have an empty in <code>Name</code>.• Species names cannot be empty and note the following reserved words, characters and constraints:<ul style="list-style-type: none">▪ The literal words <code>null</code> and <code>time</code>. Note that you can specify species names with these words contained within the name. For example <code>nullaminoacids</code>, or <code>nullnucleotides</code>.▪ The characters <code>i</code>, <code>j</code>, <code>-></code>, <code><></code>, <code>[</code>, and <code>]</code>.▪ If you are using a species name that is not a valid MATLAB variable name, do the following:<ul style="list-style-type: none">• Enclose the name in square brackets when writing a reaction rate equation or a rule.• Enter the name without brackets when you are creating the species or when you are adding the reaction. <p>For example, enclose <code>[DNA polymerase+]</code> within brackets in reaction rates and rules; enter <code>DNA polymerase+</code> when specifying the name of the species or while writing the reaction.</p> <p>For more information on valid MATLAB variable names see <code>genvarname</code>.</p> <ul style="list-style-type: none">▪ The literal words <code>null</code> and <code>time</code>. Note that you could specify species names with these words contained within the name. For example <code>nullaminoacids</code>, or <code>nullnucleotides</code> .▪ The characters <code>i</code>, <code>j</code>, <code>-></code>, <code><></code>, <code>[</code>, and <code>]</code>.

- Parameters cannot have an empty name or have the name time.

Characteristics

Applies to	Objects: kineticlaw, model, parameter, reaction, rule, species
Data type	char string
Data values	Any char string except reserved words and characters.
Access	Read/Write

Example

- 1 Create a model object with the name my_model.

```
modelObj = sbiomodel ('my_model');
```

- 2 Add a reaction object to the model object

```
reactionObj = addreaction(modelObj, 'Aspartic acid -> beta-Aspartyl-P04')
```

MATLAB returns

Reaction Object Array

Index:	Reaction:
1	Aspartic acid -> beta-Aspartyl-P04

- 3 Set reaction Name and verify

```
set (reactionObj, 'Name', 'Aspartate kinase reaction');
get (reactionObj, 'Name')
```

MATLAB returns

```
ans =
Aspartate kinase reaction
```

Name

See Also

`addkineticlaw`, `addmodel`, `addparameter`, `addrule`,
`addspecies`, `sbiomodel`

Purpose

Specify normalization type for sensitivity analysis

Description

Normalization is a property of the `SensitivityAnalysisOptions` object. `SensitivityAnalysisOptions` is a property of the configuration set object. Use `Normalization` to specify the normalization for the computed sensitivities.

The following values let you specify the type of normalization; the examples show you how sensitivities of a species x with respect to a parameter k are calculated for each normalization type:

- 'None' specifies no normalization.

$$dx(t)/dk$$

- 'Half' specifies normalization relative to the numerator (species quantity) only.

$$(1/x(t))(dx(t)/dk)$$

- 'Full' specifies that the data should be made dimensionless.

$$(k/x(t))(dx(t)/dk)$$

Characteristics

Applies to	Object: <code>SensitivityAnalysisOptions</code>
Data type	enum
Data values	'None', 'Half', 'Full'. Default is 'None'.
Access	Read/Write

See Also

`ParameterInputFactors`, `SensitivityAnalysis`,
`SensitivityAnalysisOptions`, `SpeciesInputFactors`

Notes

Purpose	Property with HTML text describing SimBiology object
Description	Contains user-specified comments about a SimBiology object
Characteristics	
Applies to	objects: kinetic law, model, parameter, reaction, rule, species
Data type	char string
Data values	Any char string
Access	Read/Write
Example	<ol style="list-style-type: none">1 Create a model object.<pre>modelObj = sbiomodel ('my_model');</pre>2 Write notes for the model object.<pre>set (modelObj, 'notes', '09/01/05 experimental data')</pre>3 Verify the assignment <pre>get (modelObj, 'notes')</pre>MATLAB returns <pre>ans = 09/01/05 experimental data</pre>
See Also	addkineticlaw, addmodel, addparameter, addreaction, addrule, addspecies, sbiomodel

Purpose

Hold parameter input factors for sensitivity analysis

Description

ParameterInputFactors is a property of the SensitivityAnalysisOptions object. SensitivityAnalysisOptions is a property of the configuration set object. Use ParameterInputFactors to specify the parameters with respect to which you want to compute the sensitivities of the species states in your model. When you simulate a model with SensitivityAnalysis enabled in the active configuration set object, SimBiology returns the computed sensitivities of the species specified in StatesToLog. For a description of the output, see the SensitivityAnalysisOptions property description.

Characteristics

Applies to	Object: SensitivityAnalysisOptions
Data type	parameter object or array of parameter objects
Data values	Parameter object array. Default is [].
Access	Read/Write

Examples

This example shows how to set ParameterInputFactors for sensitivity analysis.

- 1 Import the radio decay model from SimBiology demos.

```
modelObj = sbmlimport('radiodecay');
```

- 2 Retrieve the configuration set object from modelObj.

```
configsetObj = getconfigset(modelObj);
```

- 3 Add a parameter to the ParameterInputFactors property and display. Use the sbioselect function to retrieve the parameter object from the model.

ParameterInputFactors

```
set(configsetObj.SensitivityAnalysisOptions,'ParameterInputFactors', ...
    sbioselect(modelObj, 'Type', 'parameter', 'Name', 'c'));
get (configsetObj.SensitivityAnalysisOptions, 'ParameterInputFactors')
```

Parameter Object Array

Index:	Name:	Value:	ValueUnits:
1	c	0.5	1/second

See Also

SimBiology function `sbioselect`

SimBiology properties `SensitivityAnalysis`,
`SensitivityAnalysisOptions`, `SpeciesInputFactors`

Purpose	Property with array of parameter objects
Description	Indicates the parameters in a Model, or KineticLaw object. Read-only array of Parameter objects. Display with <code>modelObj.Parameters</code> or <code>get(modelObj, 'Parameters')</code> .
	The scope of a parameter object is hierarchical and is defined by the parameter's parent. If a parameter is defined with a kinetic law object as its parent, then only the kinetic law object can use the parameter. If a parameter object is defined with a model object as its parent, then all components within the model (including all rules, submodels and kinetic laws (reaction rate equations) can use the parameter.
	You can add a parameter to a model object, or kinetic law object with the method <code>addparameter</code> and delete it with the method <code>delete</code> .
	You can view parameter object properties with the <code>get</code> command and configure properties with the <code>set</code> command.

Characteristics

Applies to	Objects: <code>model</code> , <code>kineticlaw</code>
Data type	array of parameter objects
Data values	Parameter objects; Default value is empty ([]).
Access	Read-only

Example

- 1 Create a model object, then add a reaction object

```
modelObj = sbiomodel ('my_model');
reactionObj = addreaction (modelObj, 'a + b -> c + d');
```

- 2 Define a kinetic law for the reaction object

```
kineticlawObj = addkineticlaw (REACTIONObj, 'MassAction');
```

- 3 Add a parameter and assign it to the kinetic law object
(`kineticlawObj`);

Parameters

```
parameterObj1 = addparameter (kineticlawObj, 'K1');
get (kineticlawObj, 'Parameters')
```

Parameter Object Array

Index:	Name:	Value:	ValueUnits:
1	K1	1	

4 Add a parameter and assign it to the model object (modelObj);

```
parameterObj1 = addparameter (modelObj, 'K2');
get (modelObj, 'Parameters')
```

Parameter Object Array

Index:	Name:	Value:	ValueUnits:
1	K2	1	

See Also

[addparameter](#), [delete](#), [sbioparameter](#)

MATLAB functions [get](#) and [set](#)

Purpose

Property showing cell array of reaction rate parameters

Description

ParameterVariableNames shows the parameters used by the kinetic law object to determine the ReactionRate equation in the reaction object. Use setparameter to assign ParameterVariableNames. When you assign species to ParameterVariableNames, SimBiology maps these parameter names to ParameterVariables in the kinetic law object.

If the reaction is using a kinetic law the ReactionRate property of a reaction object shows the result of a mapping from an abstract kinetic law. The ReactionRate is determined by the kinetic law object Expression property by mapping ParameterVariableNames to ParameterVariables and SpeciesVariableNames to SpeciesVariables.

Characteristics

Applies to	Object: kineticlaw
Data type	Cell array of strings
Data values	Cell array of parameters
Access	Read/Write

Example

Create a model, add a reaction, and assign the SpeciesVariableNames for the reaction rate equation.

- 1 Create a model object, and then add a reaction object.

```
modelObj = sbiomodel('my_model');
reactionObj = addreaction(modelObj, 'a -> c + d');
```

- 2 Create a kinetic law object for the reaction object, of type 'Henri-Michaelis-Menten'

```
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');

reactionObj KineticLaw property is configured to kineticlawObj.
```

ParameterVariableNames

- 3** The 'Henri-Michaelis-Menten' kinetic law has two parameter variables (V_m and K_m) that should to be set. To set these variables,

```
setparameter(kineticlawObj, 'Vm', 'Va');  
setparameter(kineticlawObj, 'Km', 'Ka');
```

- 4** Verify that the parameter variables are correct.

```
get (kineticlawObj, 'ParameterVariableNames')
```

MATLAB returns

```
ans =  
'Va'      'Ka'
```

See Also

Reaction object property: ReactionRate

Abstract kinetic law object and kinetic law object properties:
Expression, SpeciesVariables, ParameterVariables

Kinetic law object property: SpeciesVariableNames

Method: setparameter

Purpose

Property showing parameters in abstract kinetic law

Description

Description

Indicates the parameter variables that are used in the Expression property of the abstract kinetic law object. Used to determine the ReactionRate equation in the reaction object. Use the MATLAB function set to assign ParameterVariables to an abstract kinetic law. For more information see abstract kinetic law.

Characteristics

Applies to	Objects: abstract kinetic law, kineticlaw
Data type	Cell array of strings
Data values	Defined by abstract kinetic law
Access	Read/Write in abstract kinetic law. Read-only in kinetic law.

Example

Create a model, add a reaction and assign the SpeciesVariableNames for the reaction rate equation.

- 1 Create a model object, then add a reaction object

```
modelObj = sbiomodel('my_model');
reactionObj = addreaction(modelObj, 'a -> c + d');
```

- 2 Create a kinetic law object for the reaction object, of the type 'Henri-Michaelis-Menten'

```
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');

reactionObj KineticLaw property is configured to kineticlawObj.
```

- 3 The 'Henri-Michaelis-Menten' kinetic law has two parameter variables.

ParameterVariables

```
get (kineticlawObj, 'ParameterVariables')
```

MATLAB returns

```
ans =
```

```
'Vm' 'Km'
```

See Also

Reaction object property: ReactionRate

Abstract kinetic law object and kinetic law object properties:
Expression, SpeciesVariables

Kinetic law object property: SpeciesVariableNames,
ParameterVariableNames

Method: setparameter

MATLAB function set

Purpose	Property indicating the parent object
Description	Indicates the parent object for a SimBiology object (read-only). The Parent property indicates accessibility of the object. The object is accessible to the Parent object and other objects within the Parent object. The value of Parent depends on the type of object and how it was created. <ul style="list-style-type: none">• The top level model always has the SimBiology root as the Parent• A model object can have another model object as Parent; this is the case for submodels.• Reaction and species objects, are limited to a model object or [] as Parent.• Parameter objects, are limited to a model object or a kinetic law object as Parent.• Rule object, are limited to a model object or [] as Parent• An abstract kinetic law object has [] as Parent until it has been added to the library, then has the SimBiology root as Parent
Characteristics	
Applies to	Object: abstractkineticlaw, kineticlaw, model, parameter, reaction, rule, species
Data type	Object
Data values	SimBiology component object or empty []. Default value is run-time.
Access	Read-only
See Also	sbiomodel , addkineticlaw , addmodel , addparameter , addreaction

Products

Purpose Property to indicate reaction products

Description Array of SimBiology.Species objects.

Products is a 1-by-n species object array that indicates the species that are changed by the reaction. If the Reaction property is modified to use a different species, the Products property is updated accordingly.

You can add product species to the reaction with addproduct function. You can remove product species from the reaction with rmproduct. You can also update reaction products by setting the Reaction property with the functionset.

Characteristics

Applies to	Object: reaction
Data type	Array of objects
Data values	Species objects. Default is [].
Access	Read-only

Example

1 Create a model object

```
modelObj = sbiomodel ('my_model');
```

2 Add reaction objects

```
reactionObj = addreaction (modelObj, 'a + b -> c + d');
```

3 Verify assignment.

```
productsObj = get(reactionObj, 'Products')
```

MATLAB returns

Species Object Array

Index:	Name:	InitialAmount:	InitialAmountUnits:
1	c	0	

2 d 0

See Also

`addkineticlaw`, `addspecies`, `addproduct`, `rmproduct`

RandomState

Purpose	Property to set random number generator
Description	RandomState sets the random number generator for the stochastic solvers. It is a property of the SolverOptions object. SolverOptions is a property of the configset object. SimBiology uses a pseudorandom number generator. The sequence of numbers generated is determined by the state of the generator, which can be specified by the integer RandomState. If RandomState is set to integer J, the random number generator is initialized to its J th state. The random number generator can generate all the floating-point numbers in the closed interval [2 ⁻⁵³ , 1-2 ⁻⁵³]. Theoretically, it can generate over 2 ¹⁴⁹² values before repeating itself. But for a given state, the sequence of numbers generated will be the same. To change the sequence, change RandomState. SimBiology resets the state at startup. The default value of RandomState is [].
Characteristics	
Applies to	Object: SolverOptions for SSA, explttau, impltau
Data type	int
Data values	Default is [].
Access	Read/Write
Example	Shows how to change RandomState settings.
1 Retrieve the configset object from the modelObj and change the SolverType to explttau.	
<pre>modelObj = sbiomodel('cell'); configsetObj = getconfigset(modelObj) set(configsetObj, 'SolverType', 'explttau')</pre>	
2 Change the Randomstate to 5.	
<pre>set(configsetObj.SolverOptions, 'RandomState', 5)</pre>	

```
get(configsetObj.SolverOptions, 'RandomState')
```

```
ans =
```

```
5
```

See Also

ErrorTolerance, LogDecimation, MaxIterations

Reactants

Purpose	Property to indicate reaction reactants								
Description	<p>Reactants is a 1-by-n species object array with species in the reaction. If the Reaction property is modified to use a different reactant, the Reactants property will be updated accordingly.</p> <p>You can add reactant species to the reaction with the addreaction method.</p> <p>You can remove reactant species from the reaction with the rmreactant method. You can also update reactants by setting the Reaction property with the function set.</p>								
Characteristics									
Applies to	Objects: reaction								
Data type	Species object or array of species objects								
Data values	Species objects, default is []								
Access	Read-only								
Example	<p>1 Create a model object</p> <pre>modelObj = sbiomodel ('my_model');</pre> <p>2 Add reaction objects</p> <pre>reactionObj = addreaction (modelObj, 'a + b -> c + d');</pre> <p>3 View the reactants for reactionObj.</p> <pre>get(reactionObj, 'Reactants')</pre> <p>MATLAB returns</p> <p>Species Object Array</p> <table><thead><tr><th>Index:</th><th>Name:</th><th>InitialAmount:</th><th>InitialAmountUnits:</th></tr></thead><tbody><tr><td>1</td><td>a</td><td>0</td><td></td></tr></tbody></table>	Index:	Name:	InitialAmount:	InitialAmountUnits:	1	a	0	
Index:	Name:	InitialAmount:	InitialAmountUnits:						
1	a	0							

2 b 0

See Also

[addreaction](#), [addspecies](#), [addreactant](#), [rmreactant](#)

Reaction

Purpose	Property to indicate the reaction object reaction
Description	<p>Property to indicate the reaction represented in the reaction object. Indicates the chemical reaction that can change the amount of one or more species, for example: 'A + B > C'. This property is different from the model object property Reactions.</p> <p>If the Reaction property value is modified, the Species property value of the reaction object's parent is updated. If applicable, the Reactant and/or Product properties of the reaction object are also updated.</p> <p>For example, if an additional species is added to the reaction, the species object is added to the model object Species property value. The species is also added to either the Reactant or Product property value. If a species is removed from a reaction, the species object is not removed from the Species property value. However, it is removed from the Reactant or Product property. The delete function can be used to remove the species object from the Species property value.</p>

While the following are valid reactions,

```
A > null  
null -> B
```

reactions that combine species with null are invalid.

```
A + null -> B  
A -> B + null
```

Note the use of spaces around species names and stoichiometric values.

```
glucose + 2 ADP + 2 Pi -> 2 lactic acid + 2 ATP + 2 H2O
```

Characteristics

Applies to	Object: reaction
Data type	char string
Data values	Valid reaction string, default is ''
Access	Read/Write

Example

- 1 Create a model object, then add a reaction object.

```
modelobj = sbiomodel ('my_model');  
reactionObj = addreaction (modelobj, 'a + b -> c + d');
```

- 2 Verify that the reaction property records the input.

```
get (reactionObj, 'Reaction')
```

MATLAB returns

```
ans =  
  
a + b -> c + d
```

See Also

[sbioreaction](#), [addreaction](#)

ReactionRate

Purpose	Property containing the reaction rate equation in reaction object
Description	<p>Defines the reaction rate equation. You can define a ReactionRate with or without the KineticLaw property. KineticLaw defines the type of reaction rate. The addkineticlaw function configures the ReactionRate based on the KineticLaw and the species and parameters specified in the kinetic law object properties SpeciesVariableNames and ParameterVariableNames.</p> <p>The reaction takes place in the reverse direction if the Reversible property is true. This is reflected in ReactionRate. The ReactionRate includes the forward and reverse rate if reversible</p> <p>You can specify ReactionRate without KineticLaw. Use the set function to specify the reaction rate equation. SimBiology adds species variables while creating reactionObj using the addreaction method. You must add the parameter variables (to the modelObj in this case). See the example below.</p> <p>Once you have specified the ReactionRate without KineticLaw, if you later configure the reactionObj to use KineticLaw the ReactionRate is unset until you specify SpeciesVariableNames and ParameterVariableNames.</p>
Characteristics	
Applies to	Object: reaction
Data type	char string
Data values	Reaction rate string. Default is ''
Access	Read/Write

Examples

Example 1

Create a model, add a reaction, and assign the expression for the reaction rate equation.

- 1 Create a model object, then add a reaction object.

```
modelObj = sbiomodel('my_model');
reactionObj = addreaction(modelObj, 'a -> c + d');
```

- 2** Create a kinetic law object for the reaction object, of the type 'Henri-Michaelis-Menten'.

```
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');
```

reactionObj KineticLaw property is configured to kineticlawObj.

- 3** The 'Henri-Michaelis-Menten' kinetic law has two parameter variables (V_m and K_m) and one species variable (S) that you should set. To set these variables, first create the parameter variables as parameter objects (parameterObj1, parameterObj2) with a names V_m_d , K_m_d and assign them to kineticlawObj.

```
parameterObj1 = addparameter(kineticlawObj, 'Vm_d');
parameterObj2 = addparameter(kineticlawObj, 'Km_d');
```

- 4** Set the variable names for the kinetic law object.

```
set(kineticlawObj, 'ParameterVariableNames', {'Vm_d' 'Km_d'});
set(kineticlawObj, 'SpeciesVariableNames', {'a'});
```

- 5** Verify that the reaction rate is expressed correctly in the reaction object ReactionRate property.

```
get (reactionObj, 'ReactionRate')
```

MATLAB returns

```
ans =
```

```
Vm_d*[a]/(Km_d+[a])
```

Example 2

Create a model, add a reaction, and specify ReactionRate without a kinetic law.

ReactionRate

- 1 Create a model object, then add a reaction object.

```
modelObj = sbiomodel('my_model');  
reactionObj = addreaction(modelObj, 'a + b -> c + d');
```

- 2 Specify ReactionRate and verify the assignment.

```
set(reactionObj, 'ReactionRate', 'k*a');  
get(reactionObj, 'ReactionRate')
```

MATLAB returns

```
ans =  
k*a
```

- 3 You cannot simulate the model until you add the parameter k to the modelObj.

```
parameterObj = addparameter(modelObj, 'k');
```

SimBiology adds the parameter to the modelObj with default Value = 1.0 for the parameter.

See Also

[sbioreaction](#), [addreaction](#), [sbioparameter](#), [addparameter](#), [Reversible](#)

Purpose	Property with an array of reaction objects								
Description	<p>Property to indicate the reactions in a Model object. Read-only array of reaction objects.</p> <p>A reaction object defines a chemical reaction that occurs between species. The species for the reaction are defined in the Model object property Species.</p> <p>You can add a reaction to a model object with the method addreaction and you can remove a reaction from the model object with the method delete.</p>								
Characteristics	<table><tr><td>Applies to</td><td>Objects: model</td></tr><tr><td>Data type</td><td>Array of reaction objects</td></tr><tr><td>Data values</td><td>Reaction object</td></tr><tr><td>Access</td><td>Read-only</td></tr></table>	Applies to	Objects: model	Data type	Array of reaction objects	Data values	Reaction object	Access	Read-only
Applies to	Objects: model								
Data type	Array of reaction objects								
Data values	Reaction object								
Access	Read-only								
Example	<p>1 Create a model object, then add a reaction object</p> <pre>modelObj = sbiomodel ('my_model'); reactionObj = addreaction (modelObj, 'a + b -> c + d');</pre> <p>2 Verify that the reactions property records the input</p> <pre>get (modelObj, 'Reactions')</pre> <p>MATLAB returns</p> <pre>Reaction Object Array</pre> <table><tr><td>Index:</td><td>Reaction:</td></tr><tr><td>1</td><td>a + b -> c + d</td></tr></table>	Index:	Reaction:	1	a + b -> c + d				
Index:	Reaction:								
1	a + b -> c + d								
See Also	sbioreaction , addreaction , delete								

RelativeTolerance

Purpose

Property to specify allowable error relative to component

Description

RelativeTolerance specifies the allowable error tolerance relative to the state vector at each simulation step. The state vector contains values for all the state variables, for example species amounts for all the species.

RelativeTolerance is a property of SolverOptions object.

SolverOptions is a property of the configset object.

RelativeTolerance is available for the ode solvers ('ode45', 'ode23', 'ode113', 'ode15s', 'ode23s', 'ode23t', and 'ode23tb').

If you set the RelativeTolerance at $1e-2$ you are specifying that an error of 1% relative to each state value is acceptable at each simulation step.

At each simulation step, the solver estimates the local error e_i in the i^{th} state vector y . Simulation converges at that time step if e_i satisfies the following equation:

$$|e_i| \leq \max(\text{RelativeTolerance} * |y_i|, \text{AbsoluteTolerance})$$

Thus at higher state values, convergence is determined by RelativeTolerance. As the state values approach zero, convergence is controlled by AbsoluteTolerance. The choice of values for RelativeTolerance and AbsoluteTolerance will vary depending on the problem. The default values should work for first trials of the simulation; however if you want to optimize the solution, consider that there is a trade-off between speed and accuracy. If the simulation takes too long, you can increase the values of RelativeTolerance and AbsoluteTolerance at the cost of some accuracy. If the results appear to be inaccurate, you can decrease the tolerance values but this will slow down the solver. If the magnitude of the state values is high, you can try to decrease the relative tolerance to get more accurate results.

Characteristics

Applies to	Object: SolverOptions
Data type	double
Data values	>0, <1; default is 1e-3.
Access	Read/Write

Example

Shows how to change AbsoluteTolerance.

- 1 Retrieve the configset object from the modelObj.

```
modelObj = sbiomodel('cell');
configsetObj = getconfigset(modelObj)
```

- 2 Change the AbsoluteTolerance to 1e-8.

```
set(configsetObj.SolverOptions, 'RelativeTolerance', 1.0e-6)
get(configsetObj.SolverOptions, 'RelativeTolerance')
```

```
ans =
1.0000e-006
```

See Also

AbsoluteTolerance

Reversible

Purpose Property to indicate whether a reaction is reversible or irreversible

Description Defines whether a reaction is reversible or irreversible. The rate of the reaction is defined by the ReactionRate property. For a reversible reaction the reaction rate equation is the sum of the rate of the forward and reverse reactions. The type of reaction rate is defined by the KineticLaw property. If a reaction is changed from reversible to irreversible or vice versa after KineticLaw is assigned, the new ReactionRate is determined only if Type is MassAction.. All other Types result in unchanged ReactionRate. For MassAction the first parameter specified is assumed to be the rate of the forward reaction.

Characteristics

Applies to	Object: reaction
Data type	boolean
Data values	true, false. Default value is false
Access	Read/Write

Example

Create a model, add a reaction, and assign the expression for the reaction rate equation.

- 1 Create model object, then add a reaction object.

```
modelObj = sbiomodel('my_model');
reactionObj = addreaction(modelObj, 'a -> c + d');
```

- 2 Set the Reversible property for the reactionObj to true and verify this setting.

```
set (reactionObj, 'Reversible', true)
get (reactionObj, 'Reversible')
```

MATLAB returns

```
ans =
```

1

MATLAB returns 1 for true and 0 for false.

In the next steps the example illustrates how the reaction rate equation is assigned for reversible reactions.

- 3 Create a kinetic law object for the reaction object, of the type 'MassAction'.

```
kineticlawObj = addkineticlaw(reactionObj, 'MassAction');
```

reactionObj KineticLaw property is configured to kineticlawObj.

- 4 The 'MassAction' kinetic law for reversible reactions has two parameter variables ('Forward Rate Parameter' and 'Reverse Rate Parameter') that you should set. The species variables for MassAction are automatically determined. To set the parameter variables, first create the parameter variables as parameter objects (parameterObj1, parameterObj2) with names Kf, Kr and assign the object to kineticlawObj.

```
parameterObj1 = addparameter(kineticlawObj, 'Kf');  
parameterObj2 = addparameter(kineticlawObj, 'Kr');
```

- 5 Set the variable names for the kinetic law object.

```
set(kineticlawObj, 'ParameterVariableNames', {'Kf' 'Kr'});
```

- 6 Verify that the reaction rate is expressed correctly in the reaction object ReactionRate property.

```
get (reactionObj, 'ReactionRate')
```

MATLAB returns

```
ans =
```

```
Kf*a*b - Kr*c*d
```

Reversible

See Also

[sbioreaction](#), [addreaction](#), [addparameter](#), [addreactant](#),
[ParameterVariableNames](#), [ReactionRate](#)

Purpose	Property to define certain species and parameter interactions
----------------	---

Description	A rule defines how certain species and parameters should interact with one another. For example, a rule could state that the total number of species A and species B must be some value. Rule is a MATLAB expression that defines the change in the species object quantity or a parameter object Value when the rule is evaluated.
--------------------	---

You can add a rule to a model object with the `addrule` method and remove the rule with the `delete` method. For more information on rules see `addrule`, and `RuleType`.

Characteristics

Applies to	Object: rule
Data type	char string
Data values	char string defined as species or parameter objects. Default is empty.
Access	Read/write

Example

- 1 Create a model object, then add a reaction object

```
modelObj = sbiomodel ('my_model');
reactionObj = addreaction (modelObj, 'a + b -> c + d');
```

- 2 Add a rule

```
ruleObj = addrule(modelObj, '10-a+b')
```

MATLAB returns

Rule Object Array

Index:	RuleType:	Rule:
1	algebraic	10-a+b

See Also

`addrule`, `delete`, `sbiorule`

RuleType

Purpose

Property for defining the type of rule for the rule object

Description

RuleType indicates the type of rule defined by the rule object. A Rule object defines how certain species and parameters should interact with one another. For example, a rule could state that the total number of species A and species B must be some value. Rule is a MATLAB expression that defines the change in the species object quantity or a parameter object Value when the rule is evaluated.

You can add a rule to a model object with the `addrule` method and remove the rule with the `delete` method. For more information on rules see `addrule`, and `sbiorule`

The three rule types defined are `algebraic`, `assignment`, and `rate`:

- **Algebraic** — Algebraic rules are evaluated continuously during a simulation. An algebraic rule takes the form `0 = Expression`, and the rule is specified as the `Expression`. For example, a mass conservation expression such as `species_total = species1 + species2`, where `species_total` is the independent variable, would be written as

```
species1 + species2 - species_total
```

- **Assignment** — Assignment rules are evaluated once at the beginning of a simulation. Assignment rules are expressed as `Variable = Expression`. For example write an assignment rule to set the amount of `species1` to be proportional to `species2`;

```
species1 = k/species2  
(where k is a known constant with units = concentration^2)
```

- **Rate** — Rate rules are evaluated continuously during a simulation. Rate rules are determined by `dVariable/dt = Expression`, which is expressed in SimBiology as `Variable = Expression`. For example, write a rate rule to define the rate of change in the quantity of a new species, `species3`, using the expression.

```
dspecies1/dt = k * (species1 + species2)
```

Write the rule in SimBiology as

```
species3 = k * (species1 + species2)
```

Characteristics

Applies to	Object: rule
Data type	char string
Data values	'algebraic', 'assignment', 'rate'. Default value is 'assignment'.
Access	Read/write

Example

- 1 Create a model object, then add a reaction object

```
modelObj = sbiomodel ('my_model');  
reactionObj = addreaction (modelObj, 'a -> b');
```

- 2 Add a rule that specifies the quantity of a species c. In the rule expression k is the rate constant for a -> b

```
ruleObj = addrule(modelObj, 'c = k*(a+b)')
```

- 3 Change the RuleType from the default ('algebraic') to 'rate'. and verify using the get command

```
set(ruleObj, 'RuleType', 'rate');  
get(ruleObj)
```

MATLAB returns all the properties for the rule object

```
Active: 1  
Annotation: ''  
Name: ''  
Notes: ''  
Parent: [1x1 SimBiology.Model]  
Rule: 'c = k*(a+b)'  
RuleType: 'rate'
```

RuleType

```
    Tag: ''
    Type: 'rule'
    UserData: []
```

See Also [sbiorule](#), [addrule](#), [delete](#)

Purpose Property showing rules in model object

Description Indicates the rules in a Model object. Read-only array of SimBiology.Rule objects.

A **rule** is a mathematical expression that modifies a species amount or a parameter value. A rule defines how certain species and parameters should interact with one another. For example, a rule could state that the total number of species A and species B must be some value.

You can add a rule to a model object with the `addrule` method and remove the rule with the `delete` method. For more information on rules see `addrule`, and `RuleType`.

Characteristics

Applies to	Object: model
Data type	Array of rule objects
Data values	Rule object
Access	Read-only

Example

- 1 Create a model object, then add a reaction object

```
modelObj = sbiomodel ('my_model');
reactionObj = addreaction (modelObj, 'a + b -> c + d');
```

- 2 Add a rule

```
ruleObj = addrule(modelObj, '10-a+b')
```

MATLAB returns

Rule Object Array

Index:	RuleType:	Rule:
1	algebraic	10-a+b

Rules

See Also

[addrule](#), [delete](#), [sbiorule](#)

Purpose	Property holding options for logged species					
Description	<p>RuntimeOptions holds options for species that will be logged during the simulation run. The runtime options object can be accessed through this property.</p> <p>The LogDecimation property of the configuration set object defines how often data is logged.</p>					
Property Summary	<table><tr><td>StatesToLog</td><td>Property to specify species data recorded</td></tr><tr><td>Type</td><td>Property to indicate SimBiology object type</td></tr></table>		StatesToLog	Property to specify species data recorded	Type	Property to indicate SimBiology object type
StatesToLog	Property to specify species data recorded					
Type	Property to indicate SimBiology object type					
Characteristics	Applies to	Object: configset				
	Data type	Object				
Example	Data values	Run time options				
	Access	Read-only				
Example	<p>1 Create a model object and retrieve its configuration set.</p>					
	<pre>modelObj = sbiomodel('cell'); configsetObj = getconfigset(modelObj);</pre>					
<p>2 Retrieve the RuntimeOptions object from the configset object.</p>						
<pre>runtimeObj = get(configsetObj, 'RunTimeOptions')</pre>						
<p>Runtime Settings:</p>						
<pre>StatesToLog: all</pre>						

RuntimeOptions

See Also

MATLAB functions `get`, `set`

Purpose	Enable or disable sensitivity analysis
Description	<p>The <code>SensitivityAnalysis</code> property lets you compute the time-dependent sensitivities of all the species states defined by the <code>StatesToLog</code> property with respect to the <code>SpeciesInputFactors</code> and the <code>ParameterInputFactors</code> that you specify in the <code>SensitivityAnalysisOptions</code> property of the configuration set object.</p> <p><code>SensitivityAnalysis</code> is a property of the <code>SolverOptions</code> object. <code>SolverOptions</code> is a property of the configuration set object. <code>SensitivityAnalysis</code> is available for the ode solvers ('ode45', 'ode23', 'ode113', 'ode15s', 'ode23s', 'ode23t', and 'ode23tb').</p> <p>While computing sensitivities, SimBiology uses the "complex-step approximation" to calculate derivatives of reaction rates. This technique yields accurate results for the vast majority of typical reaction kinetics, which involve only simple mathematical operations and functions. When a reaction rate involves a non-analytic function, this technique can lead to inaccurate results. An example of such a non-analytic function is the MATLAB function <code>abs</code>. If SimBiology sensitivity analysis gives questionable results on a model whose reaction rates contain unusual functions, you may be running into limitations of the complex-step method. Contact the MathWorks Technical Support group for additional information.</p> <p>See <code>SensitivityAnalysisOptions</code> for more information on sensitivity analysis.</p>
Characteristics	
Applies to	
Object: <code>SolverOptions</code>	
Data type	
<code>logical</code>	
Data values	
1, 0, <code>true</code> , <code>false</code> . Default is <code>false</code> .	
Access	
Read/Write	

SensitivityAnalysis

Examples

This example shows how to enable SensitivityAnalysis.

- 1 Retrieve the configset object from the modelObj.

```
modelObj = sbiomodel('cell');
configsetObj = getconfigset(modelObj)
```

- 2 Enable SensitivityAnalysis.

```
set(configsetObj.SolverOptions, 'SensitivityAnalysis', true)
get(configsetObj.SolverOptions, 'SensitivityAnalysis')
```

```
ans =
```

```
on
```

See Also

[SensitivityAnalysisOptions](#), [SolverOptions](#), [SolverType](#),
[StatesToLog](#)

Purpose	Hold sensitivity analysis options
Description	<p>The <code>SensitivityAnalysisOptions</code> property is an object that holds the sensitivity analysis options in the configuration set object. Sensitivity analysis is only supported for deterministic (ODE) simulations.</p> <p>Properties of <code>SensitivityAnalysisOptions</code> are summarized in “Property Summary” on page 6-88.</p> <p>When sensitivity analysis is enabled, the following command</p> <pre>[t,x,names] = sbiosimulate(modelObj)</pre> <p>returns <code>[t,x,names]</code>, where</p> <ul style="list-style-type: none">• <code>t</code> is a n-by-1 vector, where n is the number of steps taken by the ode solver and <code>t</code> defines the time steps of the solver.• <code>x</code> is a n-by-m matrix, where n is the number of steps taken by the ode solver and m is <pre>Number of states specified in StateToLog + (Number of species specified in StatesToLog * Number of input factors)</pre> <p>A SimBiology state includes species and non-constant parameters.</p> <ul style="list-style-type: none">• <code>names</code> is the list of states logged and the list of sensitivities of the species specified in <code>StatesToLog</code> with respect to the input factors. <p>For an example of the output see Examples.</p> <p>You can add a number of configuration set objects with different <code>SensitivityAnalysisOptions</code> to the model object with the <code>addconfigset</code> method. Only one configuration set object in the model object can have the <code>Active</code> property set to true at any given time.</p>

SensitivityAnalysisOptions

Property Summary

Normalization	Specify normalization type for sensitivity analysis
ParameterInputFactors	Hold parameter input factors for sensitivity analysis
SpeciesInputFactors	Hold species input factors for sensitivity analysis

Characteristics

Applies to	Object: configuration set object
Data type	Object
Data values	SensitivityAnalysisOptions properties as summarized in “Property Summary” on page 6-88 .
Access	Read-only

Examples

This example shows how to set SensitivityAnalysisOptions.

- 1 Import the radio decay model from SimBiology demos.

```
modelObj = sbmlimport('radiodecay');
```

- 2 Retrieve the configset object from the modelObj.

```
configsetObj = getconfigset(modelObj);
```

- 3 Add a parameter to the ParameterInputFactors property and display. Use the sbioselect function to retrieve the parameter object from the model.

```
set(configsetObj.SensitivityAnalysisOptions,'ParameterInputFactors', ...
    sbioselect(modelObj, 'Type', 'parameter', 'Name', 'c'));
get (configsetObj.SensitivityAnalysisOptions, 'ParameterInputFactors')
```

SensitivityAnalysisOptions

Parameter Object Array

Index:	Name:	Value:	ValueUnits:
1	c	0.5	1/second

- 4** Add a species to the `SpeciesInputFactors` property and display. Use the `sbioselect` function to retrieve the species object from the model.

```
set(configsetObj.SensitivityAnalysisOptions, 'SpeciesInputFactors', ...
    sbioselect(modelObj, 'Type', 'species', 'Name', 'z'));
get (configsetObj.SensitivityAnalysisOptions, 'SpeciesInputFactors')
```

Species Object Array

Index:	Name:	InitialAmount:	InitialAmountUnits:
1	z	0	molecule

- 5** Enable SensitivityAnalysis.

```
set(configsetObj.SolverOptions, 'SensitivityAnalysis', true)
get(configsetObj.SolverOptions, 'SensitivityAnalysis')
```

ans =

1

- 6** Simulate and return the results to three output variables. See Description for more information.

```
[t,x,names] = sbiosimulate(modelObj);
```

- 7** Display names.

names

names =

'x'

SensitivityAnalysisOptions

```
'z'  
'd[x]/d[z]_0'  
'd[z]/d[z]_0'  
'd[x]/d[c]'  
'd[z]/d[c]'
```

8 Display state values x .

x

SimBiology follows the column order shown in `names` for the values in x . The rows correspond to t .

See Also

[addconfigset](#), [getconfigset](#)

Purpose	Property holding the model solver options	
Description	<p>SolverOptions is an object that holds the model solver options in the configset object. Changing the property SolverType changes the options specified in the SolverOptions object.</p> <p>Properties of SolverOptions are summarized in the property summary on this page.</p>	
Property Summary		
AbsoluteTolerance	Property to specify largest allowable absolute error	
ErrorTolerance	Property specifies explicit or implicit tau error tolerance	
LogDecimation	Property to specify recorded simulation output frequency	
MaxIterations	Property to specify nonlinear solver maximum iterations in implicit tau	
MaxStep	Specify upper bound on solver step size	
RandomState	Property to set random number generator	
RelativeTolerance	Property to specify allowable error relative to component	
SensitivityAnalysis	Enable or disable sensitivity analysis	
Type	Property to indicate SimBiology object type	

SolverOptions

Characteristics

Applies to	Object: configset
Data type	Object
Data values	Solver options depending on SolverType. Default is SolverOptions for default SolverType (ode15s).
Access	Read-only

Example

Illustrates the changes in SolverOptions for various SolverType settings.

- 1 Retrieve the configset object from the modelObj.

```
modelObj = sbiomodel('cell');
configsetObj = getconfigset(modelObj)
```

- 2 Configure the SolverType to ode45.

```
set(configsetObj, 'SolverType', 'ode45')
get(configsetObj, 'SolverOptions')
```

Solver Settings: (ode)

```
AbsoluteTolerance: 1.000000e-006
RelativeTolerance: 1.000000e-003
```

- 3 Configure the SolverType to ssa.

```
set(configsetObj, 'SolverType', 'ssa')
get(configsetObj, 'SolverOptions')
```

Solver Settings: (ssa)

```
LogDecimation:      1
RandomState:        []
```

4 Configure the SolverType to impltau.

```
set(configsetObj, 'SolverType', 'impltau')
get(configsetObj, 'SolverOptions')
```

Solver Settings: (impltau)

```
ErrorTolerance:    3.000000e-002
LogDecimation:      1
AbsoluteTolerance:  1.000000e-002
RelativeTolerance: 1.000000e-002
MaxIterations:      15
RandomState:         []
```

5 Configure the SolverType to explttau.

```
set(configsetObj, 'SolverType', 'explttau')
get(configsetObj, 'SolverOptions')
```

Solver Settings: (explttau)

```
ErrorTolerance:    3.000000e-002
LogDecimation:      1
RandomState:         []
```

See Also [addconfigset](#), [getconfigset](#)

SolverType

Purpose Property to select solver type for simulation

Description SolverType selects a solver for a simulation. The valid SolverType values are 'ssa', 'expltau', 'impltau', 'ode45', 'ode23', 'ode113', 'ode15s', 'ode23s', and 'ode23t'. The default solver is ode15s. For a discussion about these solver types, see "Selecting a Solver".

Changing the solver type changes the options (properties) specified in the SolverOptions property of the configset object. If you change any SolverOptions these changes are persistent when you switch SolverType. For example if you set the ErrorTolerance for the expltau solver and then change to impltau when you switch back to expltau the ErrorTolerance will have the number you assigned.

Characteristics

Applies to	Object: configset
Data type	enum
Data values	'ssa', 'expltau', 'impltau', 'ode45', 'ode23', 'ode113', 'ode15s', 'ode23s', 'ode23t'. Default is ode15s.
Access	Read/Write

Example 1 Retrieve the configset object from the modelObj.

```
modelObj = sbiomodel('cell');
configsetObj = getconfigset(modelObj)
```

```
Configuration Settings - default (active)
SolverType:          ode15s
StopTime:            10.000000

SolverOptions:
AbsoluteTolerance:   1.000000e-006
RelativeTolerance:   1.000000e-003
```

```
RuntimeOptions:  
    StatesToLog:      all  
  
CompileOptions:  
    UnitConversion:   true  
    DimensionalAnalysis: true
```

2 Configure the SolverType to ode45.

```
set(configsetObj, 'SolverType', 'ode45')  
configsetObj  
  
Configuration Settings - default (active)  
    SolverType:      ode45  
    StopTime:        10.000000  
  
SolverOptions:  
    AbsoluteTolerance: 1.000000e-006  
    RelativeTolerance: 1.000000e-003  
  
RuntimeOptions:  
    StatesToLog:      all  
  
CompileOptions:  
    UnitConversion:   true  
    DimensionalAnalysis: true
```

See Also

[getconfigset](#)

MATLAB function set

Species

Purpose	Property showing species in model object
Description	<p>Indicates the species in a Model object. Read-only array of SimBiology species objects.</p> <p>Species are entities that take part in reactions. A species object is added to the Species property when a reaction is added to the model object with the method addreaction. A species object can also be added to the Species property with the method addspecies.</p> <p>If you remove a reaction with the method delete, and a species is no longer being used by any of the remaining reactions, the species object is <i>not</i> removed from the Species property. You have to use the delete method to remove species.</p> <p>There are reserved characters that cannot be used in species object names:</p> <p>Species names cannot be empty, and note the following reserved words, characters and constraints:</p> <ul style="list-style-type: none">• The literal words null and time. Note that you could specify species names with these words contained within the name. For example nullaminoacids, or nullnucleotides.• The characters i, j, -> <>, [, and].• If you are using a species name that is not a valid MATLAB variable name, do the following:<ul style="list-style-type: none">▪ Enclose the name in square brackets when writing a reaction rate equation or a rule.▪ Enter the name without brackets when you are creating the species or when you are adding the reaction. <p>For example, enclose [DNA polymerase+] within brackets in reaction rates and rules; enter DNA polymerase+ when specifying the name of the species or while writing the reaction.</p>

Characteristics

Applies to	Object: model
Data type	Array of species objects
Data values	Species object, default is empty []
Access	Read-only

See Also

[sbiospecies](#), [addreaction](#), [addsspecies](#), [delete](#)

SpeciesInputFactors

Purpose	Hold species input factors for sensitivity analysis
Description	SpeciesInputFactors is a property of the SensitivityAnalysisOptions object. SensitivityAnalysisOptions is a property of the configuration set object. Use SpeciesInputFactors to specify the species with respect to which you want to compute the sensitivities of the species states in your model. SimBiology calculates sensitivities with respect to the initial amounts of the species specified in this property. When you simulate a model with SensitivityAnalysis enabled in the active configuration set object, SimBiology returns the computed sensitivities of the species specified in StatesToLog. For a description of the output see the SensitivityAnalysisOptions property description.

Characteristics

Applies to	Object: SensitivityAnalysisOptions
Data type	Species object or array of species objects
Data values	Species object array. Default is [].
Access	Read/Write

Examples

This example shows how to set SpeciesInputFactors for sensitivity analysis.

- 1 Import the radio decay model from SimBiology demos.

```
modelObj = sbmlimport('radiodecay');
```

- 2 Retrieve the configuration set object from modelObj.

```
configsetObj = getconfigset(modelObj);
```

- 3 Add a species to the SpeciesInputFactors property and display. Use the sbioselect function to retrieve the species object from the model.

```
set(configsetObj.SensitivityAnalysisOptions,'SpeciesInputFactors', ...
    sbioselect(modelObj, 'Type', 'species', 'Name', 'z'));
get (configsetObj.SensitivityAnalysisOptions, 'SpeciesInputFactors')
```

Species Object Array

Index:	Name:	InitialAmount:	InitialAmountUnits:
1	z	0	molecule

See Also

SimBiology function `sbioselect`

SimBiology properties `SensitivityAnalysis`,
`SensitivityAnalysisOptions`, `ParameterInputFactors`

SpeciesVariableNames

Purpose Property showing cell array of species used in reaction rate equation

Description SpeciesVariableNames shows the species used by the kinetic law object to determine the ReactionRate equation in the reaction object. Use setspecies to assign SpeciesVariableNames. When you assign species to SpeciesVariableNames, SimBiology maps these species names to SpeciesVariables in the kinetic law object.

The ReactionRate property of a reaction object shows the result of a mapping from an abstract kinetic law. The ReactionRate is determined by the kinetic law object Expression property by mapping ParameterVariableNames to ParameterVariables and SpeciesVariableNames to SpeciesVariables.

Characteristics

Applies to	Object: kinetic law
Data type	Cell array of strings
Data values	Cell array of species names
Access	Read/Write

Example

Create a model, add a reaction, and assign the SpeciesVariableNames for the reaction rate equation.

- 1 Create a model object, and then add a reaction object.

```
modelObj = sbiomodel('my_model');  
reactionObj = addreaction(modelObj, 'a -> c + d');
```

- 2 Create a kinetic law object for the reaction object, of the type 'Henri-Michaelis-Menten'

```
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');
```

The reactionObj KineticLaw property is configured to kineticlawObj.

- 3** The 'Henri-Michaelis-Menten' kinetic law has one species variable (S) that you should set. To set this variable,

```
setspecies(kineticlawObj, 'S', 'a');
```

- 4** Verify that the species variable is correct.

```
get(kineticlawObj, 'SpeciesVariableNames')
```

MATLAB returns

```
ans =
```

```
'a'
```

See Also

Reaction object property: ReactionRate

Abstract kinetic law object and kinetic law object properties:
Expression, SpeciesVariables, ParameterVariables

Kinetic law object property: ParameterVariableNames

Method: setparameter

SpeciesVariables

Purpose

Property showing species in abstract kinetic law

Description

Property showing species variables that are used in the Expression property of the kinetic law object to determine the ReactionRate equation in the reaction object. Use the MATLAB function set to assign SpeciesVariables to an abstract kinetic law. For more information see abstract kinetic law.

Characteristics

Applies to	Objects: abstract kinetic law, kineticlaw
Data type	Cell array of strings
Data values	Defined by abstract kinetic law
Access	Read/Write in abstract kinetic law. Read-only in kinetic law.

Example

Create a model, add a reaction, and assign the SpeciesVariableNames for the reaction rate equation.

- 1 Create a model object, then add a reaction object

```
modelObj = sbiomodel('my_model');  
reactionObj = addreaction(modelObj, 'a -> c + d');
```

- 2 Create a kinetic law object for the reaction object, of the type 'Henri-Michaelis-Menten'

```
kineticlawObj = addkineticlaw(reactionObj, 'Henri-Michaelis-Menten');
```

reactionObj KineticLaw property is configured to kineticlawObj.

- 3 View the species variable for 'Henri-Michaelis-Menten' kinetic law.

```
get (kineticlawObj, 'SpeciesVariables')
```

MATLAB returns

```
ans =  
'S'
```

See Also

Reaction object property: ReactionRate

Abstract kinetic law object and kinetic law object properties:
Expression, ParameterVariables

Kinetic law object property: ParameterVariableNames,
SpeciesVariableNames

Method: setparameter

MATLAB function set

StatesToLog

Purpose Property to specify species data recorded

Description StatesToLog indicates the species data to log during a simulation. This is the data returned in x during execution of $(t, x) = sbiosimulate(modelObj)$. By default all species are logged.

Characteristics

Applies to	Object: RunTimeOptions
Data type	Object or vector of objects
Data values	Species objects to log. Default is All.
Access	Read/Write

Example Illustrates how to assign species to StatesToLog.

1 Create a model object by importing the file oscillator.xml.

```
modelObj = sbmlimport('oscillator');
```

2 Retrieve the first and second species in the modelObj.

```
speciesObj1 = modelObj.Species(1);  
speciesObj2 = modelObj.Species(2);
```

3 Retrieve the configsetObj of modelObj.

```
configsetObj = getconfigset(modelObj);
```

4 Set the StatesToLog to record three species; two using the retrieved species objects and one using indexing and view the species in StatesToLog.

```
set (configsetObj.RuntimeOptions, 'StatesToLog', ...  
[speciesObj1, speciesObj2, modelObj.Species(3)]);  
get(configsetObj.RuntimeOptions, 'StatesToLog')
```

Species Object Array

Index:	Name:	InitialAmount:	InitialAmountUnits:
1	pA	100	
2	pB	0	
3	pC	0	

Stoichiometry

Purpose	Property that describes species coefficients in a reaction
Description	<p>Specifies the species coefficients in a reaction. Enter an array of doubles indicating the stoichiometry of reactants (negative value) and products (positive value). Example: [-1 -1 2].</p> <p>The double specified cannot be 0. The reactants of the reaction are defined with a negative number. The products of the reaction are defined with a positive number. For example, the reaction 3 H + A-> 2 C + F has the Stoichiometry value of [-3 -1 2 1].</p> <p>When this property is configured the Reaction property updates accordingly. In the above example, if the Stoichiometry value was set to [-2 -1 2 3], the Reaction is updated to 2H + A -> 2C + 3F.</p> <p>The length of the Stoichiometry array is the sum of the Reactants array and the Products array. To remove a product or reactant from a reaction use the rmproduct or rmreactant functions. Add a product or reactant and set stoichiometry with methods addproduct and addreactant</p> <p>ODE solvers support double stoichiometry values such as 0.5. Stochastic solvers and dimensional analysis currently only support integers in Stoichiometry, therefore you must balance the reaction equation and specify integer values for these two cases.</p> <p>A -> null has a stoichiometry value of [-1]. null -> B has a stoichiometry value of [1].</p>
Characteristics	
Applies to	Object: reaction
Data type	Double array
Data values	1-by-n double, where n is length (products) + length (reactants). Default [] (empty)
Access	Read/Write

Example

- 1 Create a reaction object

```
reactionObj = sbioreaction('2 a + 3 b -> d + 2 c');
```

- 2 Verify the Reaction and Stoichiometry properties for reactionObj.

```
get(reactionObj, 'Stoichiometry')
```

MATLAB returns

```
ans =  
-2     -3      1      2
```

- 3 Set stoichiometry to [-1 -2 2 2].

```
set (reactionObj, 'Stoichiometry', [-1 -2 2 2]);  
get (reactionObj, 'Stoichiometry')
```

MATLAB returns

```
ans =  
-1     -2      2      2
```

- 4 Note with get that the Reaction property updates automatically.

```
get (reactionObj, 'Reaction')
```

MATLAB returns

```
ans =  
a + 2 b -> 2 d + 2 c
```

See Also

[sbioreaction](#), [addreaction](#), [addproduct](#), [addreactant](#), [rmproduct](#), [rmreactant](#), [Reaction](#)

StopTime

Purpose Property to set the stop time for a simulation

Description StopTime sets the stop time for a simulation. The type of StopTime is specified in the property StopTimeType.

Characteristics

Applies to	Object: configset
Data type	double
Data values	Enter a positive number. Default is 10.
Access	Read/Write

Example 1 Retrieve the configset object from the modelObj.

```
modelObj = sbiomodel('cell');
configsetObj = getconfigset(modelObj)
```

2 Configure the StopTime to 20.

```
set(configsetObj, 'StopTime', 20)
get(configsetObj, 'StopTime')
```

```
ans =
```

```
20
```

See Also StopTimeType, TimeUnits

Purpose

Property to specify the type of stop time for a simulation

Description

StopTimeType sets the type of stop time for a simulation. The stop time is specified in the StopTime property of the configset object. Valid types are approxWallTime, numberOfLogs, and simulationTime. The default is simulationTime.

- simulationTime— specify the stop time for the simulation. The solver determines and sets the time steps and the simulation stops when it reaches the specified StopTime.
- approxWallTime— specify the approximate stop time according to the clock. For example, 10s of approxWallTime is approximately 10s of real time.
- numberOfLogs— specify the total number of simulation steps to be recorded during the simulation. For example if you want to log three simulation steps, the numberOfLogs is 3. The simulation will stop after the specified numberOfLogs.

You can change the StopTimeType setting with the set function.

Characteristics

Applies to	Object: configset
Data type	enum
Data values	approxWallTime, numberOfLogs, and simulationTime
Access	Read/Write

Example

1 Retrieve the configset object from the modelObj.

```
modelObj = sbiomodel('cell');
configsetObj = getconfigset(modelObj)
```

StopTimeType

2 Configure the StopTimeType to approxWallTime.

```
set(configsetObj, 'StopTimeType', 'approxWallTime')
get(configsetObj, 'StopTimeType')
```

```
ans =
```

```
approxWallTime
```

See Also [StopTime](#), [StatesToLog](#), [TimeUnits](#)
MATLAB function set

Purpose

Property to specify a label for a SimBiology object

Description

Specifies a label associated with a SimBiology object. Use this property to group objects and then use `sbioselect` to retrieve. For example, use the Tag property in reaction objects to group synthesis or degradation reactions. You can then retrieve all synthesis reactions using `sbioselect`. Similarly, for species objects you can enter and store classification information. For example, membrane protein, transcription factor, enzyme classifications, or whether a species is an independent variable. You can also enter the full form of the name of the species. This is useful when viewing the model in the Block Diagram Explorer. For example, the species object Name could be G6P for convenience, but in the Tag you should enter the full name, Glucose-6phosphate. The graphical representation of the model in the Block Diagram Explorer (available in `sbiodesktop`) can be sorted by the Tag field, and this feature provides a method to view the full name.

Characteristics

Applies to	Objects: abstract kinetic law, kinetic law, model, parameter, reaction, rule, species
Data type	char string
Data values	Any char string
Access	Read/Write

Example

- 1 Create a model object.

```
modelObj = sbiomodel ('my_model');
```

- 2 Add reaction object and set Tag property to 'Synthesis Reaction'.

```
reactionObj = addreaction (modelObj, 'a + b -> c + d');
set (reactionObj, 'Tag', 'Synthesis Reaction')
```

- 3 Verify Tag assignment.

```
get (reactionObj, 'Tag');
```

MATLAB returns

```
ans =  
    'Synthesis Reaction'
```

See Also

[sbiomodel](#), [sbioabstractkineticlaw](#), [addkineticlaw](#), [addparameter](#),
[addreaction](#), [addrule](#), [addsspecies](#), [sbioroot](#)

Purpose

Property to show the stop time units for a simulation

Description

TimeUnits shows units for the stop time for a simulation. The type of StopTime is specified in the property StopTimeType. Unit is seconds.

Characteristics

Applies to	Object: configset
Data type	string
Data values	Default value is second.
Access	Read-only

See Also

StopTimeType, StopTime

Type

Purpose	Property to indicate SimBiology object type
Description	<p>Indicates a SimBiology object type. When you create an object in SimBiology, the value of Type is automatically defined.</p> <p>For example, when a Species object is created, the value of Type is automatically defined as 'species'.</p>
Characteristics	
Applies to	Objects: abstract kinetic law, configuration set, CompileOptions, kinetic law, model, parameter, reaction, root, rule, species, RuntimeOptions, SolverOptions.
Data type	char string
Data values	abstract_kinetic_law, configset, compileoptions, kineticlaw, parameter, reaction, root, rule, runtimeoptions, sbiomodel, species, solveroptions.
Access	Read-only
See Also	<code>sbiomodel</code> , <code>sbioparameter</code> , <code>sbioreaction</code> , <code>sbioroot</code> , <code>sbiorule</code> , <code>sbiospecies</code>

Purpose

Indicate whether to perform unit conversion

Description

The `UnitConversion` property specifies whether to perform unit conversion for the model before simulation. It is a property of the `CompileOptions` object. `CompileOptions` holds the model's compile time options and is the object property of the `configset` object.

When `UnitConversion` is set to `true`, SimBiology converts the matching physical quantities to one consistent unit system in order to resolve them. This conversion is in preparation for correct simulation, but species amounts are returned in the user-specified units.

For example, consider a reaction $a + b \rightarrow c$. Using mass action kinetics the reaction rate is defined as $a*b*k$ where k is the rate constant of the reaction. If you specify that initial amounts of a and b are $0.01M$ and $0.005M$ respectively, then units of k are $1/(M*second)$. If you specify k with another equivalent unit definition, for example, $1/((molecules/liter)*second)$, `UnitConversion` occurs after `DimensionalAnalysis`.

If `UnitConversion` fails, then you see an error when you simulate (`sbiosimulate`).

If `UnitConversion` is set to `false`, SimBiology uses the given object values.

Characteristics

Applies to	Object: <code>CompileOptions</code> (in <code>configset</code> object)
Data type	<code>boolean</code>
Data values	<code>true</code> or <code>false</code> . Default value is <code>false</code> .
Access	Read/Write

Example

Shows how to retrieve and set `unitconversion` from the default `true` to `false` in the default configuration set in a model object

- 1 Import a model.

UnitConversion

```
modelObj = sbmlimport('oscillator')
```

```
SimBiology Model - Oscillator
```

```
Model Components:
```

```
    Models:          0
    Parameters:      0
    Reactions:       42
    Rules:           0
    Species:         23
```

2 Retrieve the configset object of the model object.

```
configsetObj = getconfigset(modelObj)
```

```
Configuration Settings - default (active)
```

```
    SolverType:        ode15s
    StopTime:         10.000000
```

```
SolverOptions:
```

```
    AbsoluteTolerance: 1.000000e-006
    RelativeTolerance: 1.000000e-003
```

```
RuntimeOptions:
```

```
    StatesToLog:      all
```

```
CompileOptions:
```

```
    UnitConversion:    false
    DimensionalAnalysis: true
```

3 Retrieve the CompileOptions object.

```
optionsObj = get(configsetObj,'CompileOptions')
```

```
Compile Settings:
```

```
    UnitConversion:    false
```

DimensionalAnalysis: true

4 Assign a value of false to UnitConversion.

```
set(optionsObj, 'UnitConversion', true)
```

See Also

[getconfigset](#), [sbiosimulate](#).

MATLAB functions [get](#) and [set](#).

UserData

Purpose	Property to specify data to associate with object
Description	Property to specify data that you want to associate with a SimBiology object. The object does not use this data directly, but you can access it using the function get or dot notation.
Characteristics	
Applies to	Objects: abstract kinetic law, kinetic law, model, parameter, reaction, rule, species
Data type	Any
Data values	Any. Default is empty
Access	Read/Write
See Also	<code>sbiomodel</code> , <code>sbioabstractkineticlaw</code> , <code>sbioparameter</code> , <code>sbioreaction</code> , <code>sbioroot</code> , <code>sbiorule</code> , <code>sbiospecies</code>

Purpose

Property containing user-defined kinetic laws

Description

UserDefinedKineticLaws is a SimBiology root object property showing all user-defined abstract kinetic laws. Use the command `sbiowhos -userdefined -kineticlaw` to see the list of user-defined kinetic laws. You can use user-defined kinetic laws when you use the command `addkineticlaw` to create a kinetic law object for a reaction object. Refer to the kinetic law by name when you create the kinetic law object, for example:

```
kineticlawObj = addkineticlaw(reactionObj, 'my_kinetic_law');
```

You can add, modify, or delete UserDefinedKineticLaws. Create an abstract kinetic law with the command `sbioabstractkineticlaw` and add it to the user-defined kinetic law library with the command `sbioaddtolibrary`. `sbioaddtolibrary` also updates the UserDefinedKineticLaws property of the root object.

See “Abstract Kinetic Law” on page 6-27 for a definition and more information.

Characteristics

Applies to	Object: root
Data type	char string
Data values	Valid kinetic laws
Access	Read/Write

Examples

Example 1

This example shows the current list of user-defined kinetic laws, using the command `sbiowhos`.

```
sbiowhos -userdefined -kineticlaw
```

Abstract Kinetic Law Object Array

UserDefinedKineticLaws

Index:	Library:	Name:	Expression:
1	UserDefined	AKL1	S+P-S*P
2	UserDefined	AKL2	P+S*k
3	UserDefined	AKL3	P-S*k
4	UserDefined	AKL4	P*S*k

Example 2

This example shows the current list of user-defined kinetic laws by accessing the root object.

```
rootObj = sbioroot;
get(rootObj, 'UserDefinedKineticLaws')
```

Abstract Kinetic Law Object Array

Index:	Library:	Name:	Expression:
1	UserDefined	AKL1	S+P-S*P
2	UserDefined	AKL2	P+S*k
3	UserDefined	AKL3	P-S*k
4	UserDefined	AKL4	P*S*k

Example 3

This example shows you how to add a user-defined kinetic law and how it is displayed in UserDefinedKineticLaws.

- 1 Create an abstract kinetic law.

```
abstkineticlawObj = sbioabstractkineticlaw('mylaw1', '(k1*s)/(k2+k1*s)');
```

- 2 Assign the parameter and species variables to the expression.

```
set (abstkineticlawObj, 'SpeciesVariables', {'s'});
set (abstkineticlawObj, 'ParameterVariables', {'k1', 'k2'});
```

- 3 Add the new abstract kinetic law to the user-defined library.

```
sbioaddtolibrary(abstkineticlawObj);
```

SimBiology adds the abstract kinetic law to the user-defined library. You can verify this using sbiowhos.

```
sbiowhos -kineticlaw -userdefined
```

```
Abstract Kinetic Law Object Array
```

Index:	Library:	Name:	Expression:
1	UserDefined	mylaw1	(k1*s)/(k2+k1+s)

Alternatively,

```
rootObj = sbioroot;
get(rootObj, 'UserDefinedKineticlaws')
```

```
Abstract Kinetic Law Object Array
```

Index:	Library:	Name:	Expression:
1	UserDefined	mylaw1	(k1*s)/(k2+k1+s)

See Also

[BuiltInKineticLaws](#), [BuiltInUnits](#), [BuiltInUnitPrefixes](#)

MATLAB functions `get` and `set`

UserDefinedUnitPrefixes

Purpose

Property containing user-defined unit prefixes

Description

UserDefinedUnitPrefixes is a SimBiology root object property showing all user-defined unit prefixes. You can specify units with prefixes for species amounts and parameter values, because, SimBiology enables you to do dimensional analysis and unit conversion during simulation. The valid units and unit prefixes are either built-in or user-defined. Use the command `sbiowhos -userdefined -unit` to see the list of user-defined units.

You can add, modify, or delete UserDefinedUnitPrefixes. You can define a unit prefix with the command `sbioregisterunitprefix`, which enables you to create the unit and add it to the user-defined unit prefixes library, and also add it to the UserDefinedUnitPrefixes property of the root object.

Characteristics

Applies to	Object: root
Data type	char string
Data values	Valid unit prefixes
Access	Read/Write

Example

This example shows how to create a user-defined unit prefix and access it through the UserDefinedUnitPrefixes property.

- 1 Create a unit prefix with a multiplier of 10^{-5} (`sbioregisterunitprefix` requires you to specify the exponent).

```
sbioregisterunitprefix('peta', 15);
```

- 2 Display the unit prefix, using the command `sbiowhos`.

```
sbiowhos -userdefined -unitprefix
```

```
SimBiology UserDefined Unit Prefixes
```

Index:	Name:	Multiplier:
1	peta	1.000000e+015

Alternatively, to display only names, use the following commands:

```
r = sbioroot  
r.UserDefinedUnitPrefixes  
  
ans =  
  
'peta'
```

See Also

[BuiltInUnitPrefixes](#), [BuiltInUnits](#), [UserDefinedUnits](#),
[UserDefinedKineticLaws](#)

UserDefinedUnits

Purpose	Property containing user-defined units
Description	<p>UserDefinedUnits is a SimBiology root object property showing all user-defined units. You can specify units for species amounts and parameter values, because, SimBiology enables you to do dimensional analysis and unit conversion during simulation. The valid units are either built-in or user-defined. Use the command <code>sbiowhos -userdefined -unit</code> to see the list of user-defined units.</p> <p>You can add, modify, or delete UserDefinedUnits. You can define a unit with the command <code>sbioregisterunit</code>, which enables you to create the unit and add it to the user-defined units library, and also add it to the UserDefinedUnits property of the root object.</p>

Characteristics

Applies to	Object: root
Data type	char string
Data values	Valid units
Access	Read/Write

Example

This example shows how to create a user-defined unit and access it through the UserDefinedUnits property.

- 1 Create units for the rate constants of a first order and a second order reaction.

```
sbioregisterunit('firstorderconstant', '1/second', 1);
sbioregisterunit('secondorderconstant', '1/molecule*second', 1);
```

- 2 Display the unit, using the command `sbiowhos`.

```
sbiowhos -userdefined -unit
```

SimBiology UserDefined Units

Index:	Name:	Composition:	Multiplier:	Offset:
1	[1x18 char]	1/second	1.000000	0.000000
2	[1x19 char]	1/molecule*second	1.000000	0.000000

Alternatively, to display only names, use the following commands:

```
r = sbioroot  
r.UserDefinedUnits
```

```
ans =  
  
'secondorderconstant'  
'firstorderconstant'
```

See Also

[BuiltInUnitPrefixes](#), [BuiltInUnits](#), [UserDefinedUnitPrefixes](#),
[UserDefinedKineticLaws](#)

Value

Purpose Property to assign value to parameter object

Description The property `Value` is the value of the parameter object. The parameter object defines an assignment that can be used by the model object and/or the kinetic law object. Create parameters and assign `Value` using the method `addparameter`.

Characteristics

Applies to	Object: parameter
Data type	<code>double</code>
Data values	Any <code>double</code> . Default value is 1.0.
Access	Read/Write

Example Assign a parameter with value to the model object

- 1 Create a model object, then add a reaction object

```
modelObj = sbiomodel ('my_model');
```

- 2 Add a parameter to the model object (`modelObj`) with Value 0.5.

```
parameterObj1 = addparameter (modelObj, 'K1', 0.5)
```

MATLAB returns

Parameter Object Array

Index:	Name:	Value:	ValueUnits:
1	K2	0.5	

See Also `addparameter`, `sbioparameter`

Purpose	Property with parameter value units
Description	Indicates the unit definition of the parameter object Value property. ValueUnits can be one of the builtin units. To get a list of the builtin units use the <code>sbioshowunits</code> function. If ValueUnits changes from one unit definition to another, the Value does not automatically convert to the new units. The <code>sbioconvertunits</code> function does this conversion.
You can add a parameter object to a model object or a kinetic law object.	
Characteristics	
Applies to	Object: parameter
Data type	char string
Data values	Unit from units library, default is empty ''
Access	Read/Write
Example	Assign a parameter with value to the model object.
1 Create a model object, then add a reaction object.	
<pre>modelObj = sbiomodel('my_model');</pre>	
2 Add a parameter with Value 0.5 , assign it to the model object (modelObj).	
<pre>parameterObj1 = addparameter(modelObj, 'K1', 0.5, 'ValueUnits', 1/second)</pre>	
MATLAB returns	
<pre>Parameter Object Array</pre>	
<pre>Index: Name: Value: ValueUnits: 1 K4 0.5 1/second</pre>	
See Also	<code>sbioparameter</code> , <code>addparameter</code> , <code>sbioshowunits</code> , <code>sbioconvertunits</code>

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