SBML Model Report

Model name: "KaDe output:"



August 8, 2017

1 General Overview

This is a document in SBML Level 2 Version 4 format. Table 1 gives an overview of the quantities of all components of this model.

Element	Quantity	Element	Quantity
compartment types	0	compartments	1
species types	0	species	2
events	0	constraints	0
reactions	3	function definitions	0
global parameters	4	unit definitions	2
rules	0	initial assignments	0

Table 1: The SBML components in this model.All components are described in more detail in the following sections.

2 Unit Definitions

This is an overview of five unit definitions. The units area, length, and time are predefined by SBML and not mentioned in the model.

2.1 Unit substance

Name substance

Definition mol

2.2 Unit volume

Name volume

Definition 1

2.3 Unit area

Name Predefined unit area

Notes

 $\text{Definition} \ m^2$

2.4 Unit length

Name Predefined unit length

Notes

 $\textbf{Definition} \ m$

2.5 Unit time

Name Predefined unit time

Notes

Definition s

3 Compartment

This model contains one compartment.

Id	Name	SBO	Spatial Dimensions	Size	Unit	Constant	Outside
default			3	1	litre		

Table 2: Properties of all compartments

3.1 Compartment default

This is a three-dimensional compartment with a constant size of one litre.

4 Species

This model contains two species. Section 7 provides further details and the derived rates of change of each species.

Id	Name	Compartment	Derived Unit	Constant	Boundary Condi- tion
s1	A(x, y)	default	mol		
s2	A(x, y 1), A(x, y 1)	default	mol		

Table 2. Dre nortion of each analis

5 Parameters

This model contains four global parameters.

	Table 4. I toperties of each parameter.					
Id	Name	SBO	Value	Unit	Constant	
tinit			0.00	S		
tend			1.00	S		
period			0.01	S		
k			1.00			

Table 4: Properties of each parameter.

6 Reactions

This model contains three reactions. All reactions are listed in the following table and are subsequently described in detail. If a reaction is affected by one or more modifiers, the identifiers of the modifier species are written above the reaction arrow.

Table 5: Overview of all reactions				
Nº Id	Name	Reaction Equation	SBO	
1 re1	1 $A(x,y), A(x,y) \rightarrow A(x,y!1), A(x,y!1)$	$2s1 \longrightarrow 2s2$		
2 re2	2 $A(x,y), A(x,y) \rightarrow A(x!1,y), A(x,y!1)$	$2s1 \longrightarrow 2s2$		
3 re3	$A(x,y), A(x,y) \to A(x!1,y), A(x!1,y)$	$2s1 \longrightarrow 2s2$		

Table 5: Overview of all reaction

6.1 Reaction re1

This is an irreversible reaction of one reactant forming one product.

Name $A(x,y), A(x,y) \rightarrow A(x,y!1), A(x,y!1)$

Reaction equation

$$2s1 \longrightarrow 2s2$$
 (1)

Reactant

Table 6: Properties of each reactant					
	Id	Name	SBO		
	s1	A(x, y)			

Product

Table 7: Properties of each product.					
Id	Name	SBO			
s2	A(x, y!1), A(x, y!1)				

Kinetic Law

Derived unit contains undeclared units

$$v_1 = \frac{\mathbf{k}}{2} \cdot [\mathbf{s1}] \cdot [\mathbf{s1}] \tag{2}$$

6.2 Reaction re2

This is an irreversible reaction of one reactant forming one product.

Name $A(x,y), A(x,y) \rightarrow A(x!1,y), A(x,y!1)$

Reaction equation

$$2s1 \longrightarrow 2s2$$
 (3)

Reactant

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Table 8: Properties of each reactant.

Id	Name	SBO
s1	A(x, y)	

Product

Table 9: Properties of each product.					
Id	Name	SBO			
s2	A(x, y!1), A(x, y!1)				

Kinetic Law

Derived unit contains undeclared units

$$v_2 = \mathbf{k} \cdot [\mathbf{s}\mathbf{1}] \cdot [\mathbf{s}\mathbf{1}] \tag{4}$$

6.3 Reaction re3

This is an irreversible reaction of one reactant forming one product.

Name $A(x,y), A(x,y) \rightarrow A(x!1,y), A(x!1,y)$

Reaction equation

$$2s1 \longrightarrow 2s2$$
 (5)

Reactant

Table 1	0: Pr	operties of	f each re	eactant.
	Id	Name	SBO	
	s1	A(x, y)		

Product

1	Table	11: Properties of each	product.
	Id	Name	SBO
	s2	A(x, y!1), A(x, y!1)	

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Kinetic Law

Derived unit contains undeclared units

$$v_3 = \frac{\mathbf{k}}{2} \cdot [\mathbf{s1}] \cdot [\mathbf{s1}] \tag{6}$$

7 Derived Rate Equations

When interpreted as an ordinary differential equation framework, this model implies the following set of equations for the rates of change of each species.

Identifiers for kinetic laws highlighted in gray cannot be verified to evaluate to units of SBML substance per time. As a result, some SBML interpreters may not be able to verify the consistency of the units on quantities in the model. Please check if

- parameters without a unit definition are involved or
- volume correction is necessary because the hasOnlySubstanceUnits flag may be set to false and spacialDimensions > 0 for certain species.

7.1 Species s1

Name A(x, y)

Initial amount 100 mol

This species takes part in three reactions (as a reactant in re1, re2, re3).

$$\frac{\mathrm{d}}{\mathrm{d}t}\mathbf{s}\mathbf{1} = -2 v_1 - 2 v_2 - 2 v_3 \tag{7}$$

7.2 Species s2

Name A(x, y!1), A(x, y!1)

Initial amount 0 mol

This species takes part in three reactions (as a product in re1, re2, re3).

$$\frac{\mathrm{d}}{\mathrm{d}t}\mathbf{s}^2 = 2 v_1 + 2 v_2 + 2 v_3 \tag{8}$$

References

Dräger, A., Planatscher, H., Wouamba, D. M., Schröder, A., Hucka, M., Endler, L., Golebiewski, M., Müller, W., and Zell, A. (2009). SBML2LATEX: Conversion of SBML files into humanreadable reports. Bioinformatics, 25(11), 1455–1456. 10.1093/bioinformatics/btp170.

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