## SBML Level 3 Package Specification

# Multistate, Multicomponent and Multicompartment Species Package for SBML Level 3

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This is a release candidate specification for the SBML Level 3 package called "*Multi*". Please send feedback to the package mailing list at sbml-multi@lists.sourceforge.net.

The latest release, past releases, and other materials related to this specification are available at <a href="http://sbml.org/Documents/Specifications/SBML\_Level\_3/Packages/Multistate\_and\_Multicomponent\_Species\_(multi)">http://sbml.org/Documents/Specifications/SBML\_Level\_3/Packages/Multistate\_and\_Multicomponent\_Species\_(multi)</a>

 $This \, {\it release} \, of \, the \, specification \, is \, available \, at \, $$ https://sourceforge.net/p/sbml/code/HEAD/tree/trunk/specifications/sbml-level-3/version-1/multi/spec/sbml-multi_spec_1.1.rc2.pdf$ 



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## 1 Introduction

This Multistate, Multicomponent and Multicompartment Species (Multi) package provides an extension of SBML Level 3 [Hucka et al. (2016)] that supports encoding models with molecular complexes that have multiple components and can exist in multiple states and in multiple compartments. One of its goals also is to provide a platform for sharing models based on the specifications of bi-molecular interactions and the rules governing such interactions [Angermann et al. (2012); Feret et al. (2009); Hlavacek et al. (2006); Zhang et al. (2013)]. This specification covers the goals and features described in the previous Multi proposal [Le Novère and Oellrich (2010)] for extending SBML to carry the information for *multistate multicomponent* species with revised data structure. In addition, this specification includes the feature for *multicompartment* species as described in the releases of the Multi proposal [Zhang and Meier-Schellersheim (2013a), Zhang et al. (2012)].

# 1.1 Proposal and specifications

The proposal corresponding to this package specification is available at:

http://sbml.org/Community/Wiki/SBML\_Level\_3\_Proposals/Multistate\_and\_Multicomponent\_Species\_ Proposal

The specifications (v1.0.1 to current) are located at:

https://sourceforge.net/p/sbml/code/HEAD/tree/trunk/specifications/sbml-level-3/version-1/multi/spec/

# 1.2 Package dependencies

The Multi package has no dependencies on other SBML Level 3 packages.

#### 1.3 Document conventions

UML 1.0 notation is used in this document to define the constructs provided by this package. Colors in the diagrams carry the following additional information for the benefit of those viewing the document on media that can display color:

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- Black Items colored black are components taken unchanged from their definitions in the SBML Level 3 Core specification document.
- **Green** Items colored green are components that exist in SBML Level 3 Core, but are extended by this package. Class boxes are also drawn with with dashed lines to further distinguish them.
- Blue Items colored blue are new components introduced in this package specification. They have no equivalent in the SBML Level 3 Core specification.

For other matters involving the use of UML, XML and typographical conventions, this document follows the conventions used in the SBML Level 3 Core specification document [Hucka et al. (2016)].

For simplicity, "..." in all example code refers to some unspecified code content, that is not important for the purpose of illustrating the issue at hand.

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# 2 Background and context

Rule-base, domain-detailed modeling has been extremely valuable in systems biology related studies [Manes et al. (2015) and Miskov-Zivanov1 et al. (2013)]. Rule-based, domain-detailed modeling approaches (*BioNetGen* [Faeder et al. (2009)], *Kappa* [Danos and Laneve (2004)], and *Simmune* [Angermann et al. (2012); Meier-Schellersheim et al. (2006)]) define rules for interactions between pairs of molecule domains, specifying how the interactions depend on particular states of the molecules (pattern) and their locations in specific compartments. In order to generate networks of biochemical reactions these rules are applied to the molecular components of the systems to be modeled, either at the beginning of the modeling (simulation) process or "on the fly" (as molecule complexes emerge from the interaction rules). Expressing such rule-based, domain-detailed reaction networks using the concepts of **Species** and **Compartment** in SBML (L3 core and L2) can be difficult for rules and molecule sets that lead to large numbers of resulting molecular complexes. It would therefore be desirable to have an SBML standard for encoding rule-based, domain-detailed models using their "native" concepts for describing reactions instead of having to apply the rules and unfold the networks prior to encoding in an SBML format.

We proposed a revised proposal of the Multi package: "Multistate, Multicomponent and Multicompartment Species Package for SBML Level 3" (abbreviated as Multi) [Zhang et al. (2012) and Zhang and Meier-Schellersheim (2013a)] which takes the scopes and some data structures developed in the previous Multi proposal [Le Novère and Oellrich (2010)] and addresses main issues arising from a rule-based, domain-detailed modeling point of view with the data structures consistent with that used in the available rule-based, domain-detailed modeling tools.

Note:

This specification was developed with the main goal of taking into account bi-molecular interactions mediated through specific binding domains (or sites). Models without such detailed description of the molecular interactions can be encoded as well if the other features in this specification such as **SpeciesFeatureType**, **SpeciesFeature**, and extended **Compartment** satisfy the model requirements.

# 2.1 Past work on this problem or similar topics

- Nicolas Le Novère and Anika Oellrich proposed the previous version of the Multi proposal [Le Novère and Oellrich (2010)]. However, it was realized that a more detailed treatment of molecular binding sites and their state-dependent interactions would be desirable.
- In August 2012, Fengkai Zhang from the *Simmune* group presented "Draft for discussion SBML Proposals for Revised Multi, Simple Spatial and Multi-Spatial Extensions" at COMBINE 2012 [Zhang et al. (2012)]. The three proposals cover the goals and scope of the previous Multi proposal (2010), revise it and add some new features that improve usage of the proposal for rule-based approaches.
- Based on the discussions and suggestions received during COMBINE 2012 as well as on feedback from the SBML discussion forum, the new Multi proposal [Rev 221, Zhang and Meier-Schellersheim (2012)] was released to the SBML-Multi community, which integrates and covers most of the features in the three previous proposals of August 2012.
- In May 2013, a new reversion (rev 280) of the Multi proposal [Zhang and Meier-Schellersheim (2013a)] was released before the meeting of HARMONY 2013. The extended Compartment class and its related classes have been reorganized. All optional boolean attributes have been removed/replaced. A new optional Multi attribute, "whichValue", was added to the ci elements in KineticLaw to identify the sources of species. (Lucian Smith gave many comments/suggestions about this proposal and William Hlavacek gave thoughtful feedback about the BioNetGen example in this proposal). This revision (rev 280) was presented at HARMONY 2013 [Zhang and Meier-Schellersheim (2013b)] with new features to configure multiple occurrences of SpeciesFeature-Type. Several new or revised features were discussed during and after HARMONY 2013, including multiple occurrences of SpeciesFeatureType, multiple copies of SpeciesTypeInstance, the numericValue attribute for PossibleSpeciesFeatureValue and concentration summation of pattern species. These features are covered

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or updated in the specifications from v1.0.1.

# 2.2 Revision history

The versioning convention used in this document:

x.y.z (status)

x: version of SBML Level 3 core.

y: version of the Multi package.

z: release of the Multi package at its version y.

status: "draft", "release candidate", or "release".

For example, the current version is "1.1.rc2 (release candidate)"

x = "1"

y = "1"

z = "rc2"

status = "release candidate"

The followings are the revision history of the Multi package:

### 2.2.1 Version: 1.1.rc3 (release candidate), this version

Modify the numbers of several rules to be consistent with the general SBML validation rule conventions.

#### 2.2.2 Version: 1.1.rc2 (release candidate), January 2017

Add a new validation rule 20306 (Section A on page 78) to prevent circular referencing among the extended **Compartment** objects.

Revise the specification text with minor changes towards a version of the official release candidate.

#### 2.2.3 Version: 1.1.rc1 (release candidate), November 2016

Revise the specification text with minor changes towards a version of the official release candidate.

## 2.2.4 Version: 1.0.7 (draft), August 2016

Remove the SpeciesFeatureChange and ListOfSpeciesFeatureChanges classes under SpeciesTypeComponentMap-InProduct. The relations expressed in SpeciesFeatureChange can be inferred from the speciesTypeComponent-MapInProduct and the species of the mapped reactant and product.

Add a new validation rule 21306, "an **outwardBindingSite** cannot be a binding site in a bond of the species" (see Section 3.16.3 on page 28 and Section A on page 75)

#### 2.2.5 Version: 1.0.6 (draft), March 2016

Remove recursively referencing relationship in the **ListOfSpeciesFeatures** class and add a **SubListOfSpeciesFeatures** class. See the details in **Species**.

Version 1.0.6.1 with minor document update is released in April 2016.

## 2.2.6 Version 1.0.5 (draft), November 2015

This version has been developed from the previous release v1.0.4 with the following modifications based on the discussion during and after COMBINE 2015 [Zhang (2015)]:

■ Drop the occur attribute in the class of **SpeciesTypeInstance**.

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- Drop the occur attribute in the class of **SpeciesTypeComponentIndex**.
- Drop the class of **DenotedSpeciesTypeComponentIndex**.
- Revise the scope of **PossibleSpeciesFeatureValue** ids to be global.

Version 1.0.5.1 with minor document update is released in Dec 2015.

## 2.2.7 Version 1.0.4 (draft), June 2015

This version has been developed from the previous release v1.0.3 with minor document update and complete validation rules.

## 2.2.8 Version 1.0.3 (draft), April 2015

This version has been developed from the previous release v1.0.2 mainly based on the discussion in COMBINE 2014 with focus on how to facilitate tools to export and import models encoded in the Multi format [Zhang and Meier-Schellersheim (2014)]

## 2.2.9 Version 1.0.2 (draft), November 2014

This version has been developed from the previous release v1.0.1 with the following modifications:

- A new BindingSiteSpeciesType sub-class inheriting the SpeciesType class for binding sites. Accordingly, the isBindingSite attribute has been dropped from SpeciesType.
- Restriction on **binding** sites which have to be atomic.
- Restriction on SpeciesType that a speciesType cannot have a listOfSpeciesFeatureTypes if it has a list-OfInSpeciesTypeBonds.
- A new IntraSpeciesReaction sub-class inheriting the Reaction class for the reactions happening within a Species object. Accordingly, the isIntraSpeciesReaction attribute has been dropped from Reaction.
- Validation rules.

#### 2.2.10 Version 1.0.1 (draft), September 2013

This was released and presented in COMBINE 2013 [Zhang and Meier-Schellersheim (2013c)], mainly addressing the scenario of multiple occurrences of identical components and/or identical features.

## 2.2.11 Revision history before draft version 1.0.1

See the past work (Section 2.1 on page 6).

# 3 Package syntax and semantics

This section contains a definition of the syntax and semantics of the Multi package for SBML Level 3 Core.

# 3.1 Namespace URI and other declarations necessary for using this package

The following is the namespace URI for this version of the Multi package for SBML Level 3 Core:

```
"http://www.sbml.org/sbml/level3/version1/multi/version1"
```

In addition, SBML documents using a given package must indicate whether the package can be used to change the mathematical interpretation of a model. This is done using the attribute **required** on the **<sbml>** element in the SBML document. For the Multi package, the value of this attribute must be "true".

The following fragment illustrates the beginning of a typical SBML model using SBML Level 3 Core and this version of the Multi package:

```
<?xml version="1.0" encoding="UTF-8"?>
<sbml xmlns="http://www.sbml.org/sbml/level3/version1/core" level="3" version="1"
    xmlns:multi="http://www.sbml.org/sbml/level3/version1/multi/version1" multi:required="true">
```

# 3.2 Primitive data types

The Multi package uses a number of the primitive data types described in Section 3.1 of the SBML Level 3 Core [Hucka et al. (2016)] specification such as SId, SIdRef, string, boolean, int and positiveInteger, and adds three additional primitive types described below.

#### 3.2.1 Type BindingStatus

The BindingStatus primitive data type is used in the definition of the **OutwardBindingSite** class. BindingStatus is derived from type string and its values are restricted to be one of the following possibilities: "bound", "unbound", and "either". Attributes of type BindingStatus cannot take on any other values. The meaning of these three values is discussed in the context of the **OutwardBindingSite** class in Section 3.16 on page 28.

#### 3.2.2 Type Relation

The Relation primitive data type is used in the definition of the **SubListOfSpeciesFeatures** class. Relation is derived from type string and its values are restricted to be one of the following possibilities: "and", "or", and "not". Attributes of type Relation cannot take on any other values. The meaning of these three values is discussed in the context of the **SubListOfSpeciesFeatures** class in Section 3.17 on page 29.

#### 3.2.3 Type RepresentationType

The RepresentationType primitive data type is used in the extension of the ci element.

RepresentationType is derived from type string and its values are restricted to be one of the following possibilities: "sum" or "numericValue". If present, attributes of type RepresentationType cannot take on any other values. The meaning of these values is discussed in the context of the ci element in Section 3.26 on page 40.

# 3.3 The new and extended classes in the Multi Package

The Multi package defines or extends the following object classes, Model, ListOfSpeciesTypes, Compartment, ListOfCompartmentReferences, CompartmentReference, SpeciesType, ListOfSpeciesTypeInstances, ListOfSpeciesFeatureTypes, ListOfInSpeciesTypeBonds, ListOfSpeciesTypeComponentIndexes, SpeciesFeatureType, ListOfPossibleSpeciesFeatureValues, PossibleSpeciesFeatureValue, SpeciesTypeInstance, InSpeciesTypeBond,

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SpeciesTypeComponentIndex, Species, ListOfOutwardBindingSites, ListOfSpeciesFeatures, SubListOfSpeciesFeatures, OutwardBindingSite, SpeciesFeature, ListOfSpeciesFeatureValues, SpeciesFeatureValue, Reaction,

SimpleSpeciesReference, SpeciesReference, ListOfSpeciesTypeComponentMapsInProduct, and

SpeciesTypeComponentMapInProduct.

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All the classes in the Multi package are directly or indirectly derived from **SBase**, and **SBase** provides the ability to attach SBO terms as well as MIRIAM annotations. The semantics of a given class in the Multi package can be made more precise by referencing to external controlled vocabularies and ontologies.

Like the classes in SBML Level 3 Core, most new Multi classes have the attribute id (typically mandatory but not all, and of type SId), which serves as an identifier to provide a way to identify the class object. The identifier of a class object reference may or may not carry mathematical interpretation or be used in mathematical formulas, depending on its class and the class object referencing it. The scope of ids is described in the section of "Namespace scoping rules for identifiers" (Section 3.27 on page 42).

3.4 Model

The Multi package extends the **Model** class of SBML Level 3 Core and adds an optional **ListOfSpeciesTypes** child to **Model**. Figure 1 provides the UML diagram for the extended **Model** class.

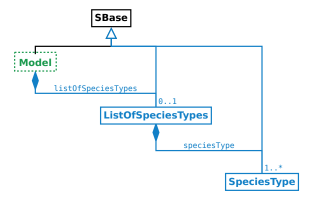


Figure 1: The extension of the Model class.

## 3.4.1 ListOfSpeciesTypes

**ListOfSpeciesTypes** is defined in Figure 1. If present, a **ListOfSpeciesTypes** object must contain at least one **Species-Type** object. Since **ListOfSpeciesTypes** is derived from **SBase**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

# 3.5 Extended Compartment

A **Compartment** object in SBML Level 3 Core represents a bounded space in which *species* are located. In the Multi package, **Compartment** is extended. A Multi **compartment** can be a **type** that multiple referencing **compartments** can map to. A Multi **compartment** can also be a composite **compartment** or a container that includes other **compartments**.

The extension of **Compartment** is defined in Figure 2. The extended **Compartment** class has a new required attribute isType, a new optional attribute compartmentType and an optional **ListOfCompartmentReferences** child. The example at Section 4.1 on page 43 illustrates the use of the extended **Compartment** class.

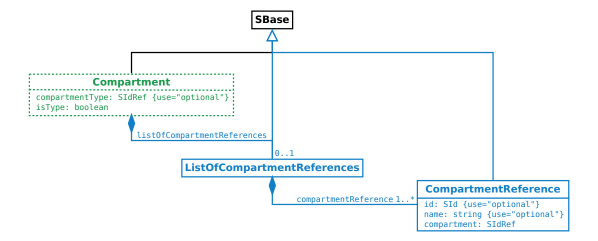


Figure 2: The definitions of Compartment, ListOfCompartmentReferences and CompartmentReference

## 3.5.1 The isType attribute

The required attribute isType, of type boolean, on the **Compartment** class serves to provide a way to indicate whether the **Compartment** object is a compartment type.

A **Compartment** object is a compartment type if the value of its isType attribute is "true". A compartment type is a template (in the sense of prototype) for all **Compartment** objects referencing it (via compartmentType attributes). A **Species** object directly referencing a compartment type is not a *fully defined* species (see Section 3.19 on page 33).

If the value of the isType attribute is "false", the Compartment object is a "not-a-type" compartment, and it is similar to a SBML core compartment except it can reference a compartment type and can have a ListOfCompartment-References child.

#### 3.5.2 The compartmentType attribute

The optional attribute compartmentType, of type SIdRef, is used for a "not-a-type" compartment to reference a compartment type. A compartment with the "true" value of its isType attribute cannot have the compartmentType attribute defined.

## 3.5.3 ListOfCompartmentReferences

**ListOfCompartmentReferences** is defined in Figure 2, and is extended from the **ListOf** class. A **listOfCompartmentReference** children. Since **ListOfCompartmentReferences** is derived from **SBase** through **ListOf**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

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# 3.6 CompartmentReference

**CompartmentReference** is defined in Figure 2 on the previous page. It has two optional attributes **id** and **name**, and a required attribute **compartment**. Since **CompartmentReference** is derived from **SBase**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

#### 3.6.1 The id and name attributes

The optional id attribute, of type SId, serves to provide a way to identify a compartmentReference. Compartment-Reference also has an optional name attribute of type string.

If some or all compartmentReferences within a ListOfCompartmentReferences object reference the same compartment, those compartmentReferences are required to have their id attributes defined to distinguish different compartmentReferences.

## 3.6.2 The compartment attribute

The required compartment attribute, of type SIdRef, serves to provide a way to reference a Compartment object.

Note:

A compartmentReference cannot reference a compartment that directly or indirectly contains the compartment-Reference. In other words, circular references are not allowed when constructing compartments and compartment-References.

# 3.7 The relationship of Compartment, CompartmentReference and ListOfCompartmentReferences

In a **ListOfCompartmentReferences** object, every children **compartmentReferences** must exclusively reference, directly or indirectly, "**not-a-type**" **compartment** which can be of the same compartment type. See the extended **Compartment** objects in the example in Section 4.1 on page 43.

All compartments referenced by a listOfCompartmentReferences must have the values of their isType attributes the same as that in the parent compartment of the listOfCompartmentReferences. For example, a compartment "A" with isType="true" has a listOfCompartmentReferences referencing two compartments "A1" and "A2". Then, "A1" and "A2" must have isType="true".

## 3.8 SpeciesType

SpeciesType is defined in Figure 3 and serves to provide backbone structures for species. SpeciesType has one required attribute, id, two optional attributes, name and compartment and four optional ListOf\_ objects of ListOfSpeciesFeatureTypes, ListOfSpeciesTypeInstances, ListOfInSpeciesTypeBonds and ListOfSpeciesTypeComponentIndexes respectively. Since SpeciesType is derived from SBase, it inherits the sboTerm and metaid attributes, as well as the optional children Notes and Annotation objects.

The **ListOfSpeciesTypeInstances** subobject provides a way to define multicomponents which are instances of other **SpeciesType** objects.

The **ListOfSpeciesFeatureTypes** subobject and its **SpeciesFeatureType** children set up a framework for the referencing **species** or the instances of **speciesTypes** to be able to have multistates. The **ListOfSpeciesTypeComponent-Indexes** subobject provides a flexible way to reference any **component** in a **speciesType**.

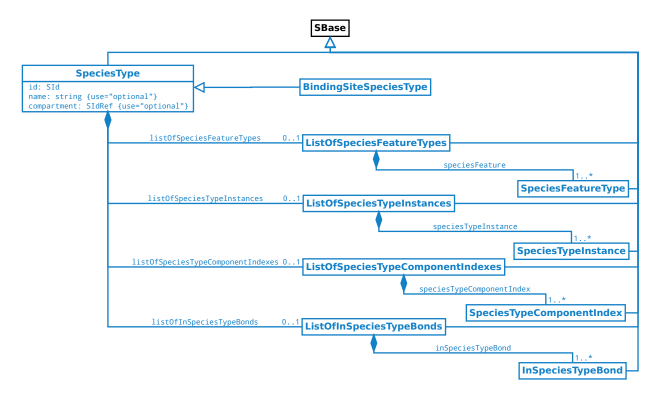


Figure 3: The definition of the SpeciesType class.

#### 3.8.1 The id and name attributes

The required id attribute, of type SId, serves to provide a way to identify a speciesType. SpeciesType also has an optional name attribute of type string.

#### 3.8.2 The compartment attribute

**SpeciesType** has an optional attribute compartment, of type SIdRef, to be used to identify the compartment where the speciesType is located. The attribute value must be the identifier of an existing compartment in the model. If present, it must be consistent with the compartment attributes of the referencing species (see Section 3.15 on page 26) and the compartmentReference attributes of its instances (see Section 3.11.3 on page 18). The example in Section 4.1 on page 43 illustrates how to keep the consistency of this attribute.

#### 3.8.3 ListOfSpeciesFeatureTypes

**ListOfSpeciesFeatureTypes** is defined in Figure 3 on the previous page, and is extended from the **ListOf** class. If present, a **listOfSpeciesFeatureTypes** must have one or more **SpeciesFeatureType** children. Since **ListOf-SpeciesFeatureTypes** is derived from **SBase** through **ListOf**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

#### 3.8.4 ListOfSpeciesTypeInstances

**ListOfSpeciesTypeInstances** is defined in Figure 3 on the preceding page, and is extended from the **ListOf** class. If present, a listOfSpeciesTypeInstances must have one or more **SpeciesTypeInstance** children. Since **ListOfSpeciesTypeInstances** is derived from **SBase** through **ListOf**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

#### 3.8.5 ListOfInSpeciesTypeBonds

**ListOfInSpeciesTypeBonds** class is defined in Figure 3 on the previous page, and is extended from the **ListOf** class. If present, a **listOfInSpeciesTypeBonds** must have one or more **InSpeciesTypeBond** children. Since **ListOfInSpeciesTypeBonds** is derived from **SBase** through **ListOf**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

## 3.8.6 ListOfSpeciesTypeComponentIndexes

**ListOfSpeciesTypeComponentIndexes** is defined in Figure 3 on the preceding page, and is extended from the **ListOf** class. If present, a **listOfSpeciesTypeComponentIndexes** must have one or more **SpeciesTypeComponentIndex** children. Since **ListOfSpeciesTypeComponentIndexes** is derived from **SBase** through **ListOf**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

#### 3.8.7 BindingSiteSpeciesType

BindingSiteSpeciesType inherits the SpeciesType class and is defined in Figure 3 on the previous page. A BindingSiteSpeciesType object is a binding site, and therefore its instance can further define the bindingStatus attribute and can participate a binding internally and explicitly in an InSpeciesTypeBond object, or externally and implicitly defined by an OutwardBindingSite object. A binding site must be an atomic component which means that a BindingSiteSpeciesType object cannot contain a ListOfSpeciesTypeInstances subobject.

Note:

In the Multi package, a binding site can only participate in one binding at a time. That means a binding site cannot bind two partners at the same time. The binding relationship is one-to-one.

# 3.9 SpeciesFeatureType

SpeciesFeature objects. SpeciesFeatureType has two required attributes id and occur, an optional attribute name, and a required child listOfPossibleSpeciesFeatureValues. The multiple possibleSpeciesFeatureValues of the ListOfPossibleSpeciesFeatureValues object permit constructing multistate species via its speciesFeatures under the ListOfSpeciesFeatureValues or SubListOfSpeciesFeatureValues object. Since SpeciesFeatureType is derived from SBase, it inherits the sboTerm and metaid attributes, as well as the optional children Notes and Annotation objects.

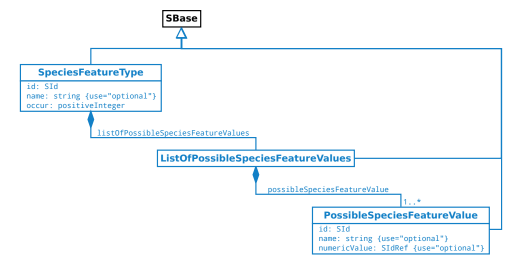


Figure 4: The definitions of SpeciesFeatureType, ListOfPossibleSpeciesFeatureValues and PossibleSpeciesFeatureValue and PossibleSpeciesFeatureValue classes.

#### 3.9.1 The id and name attributes

The required id attribute, of type SId, serves to provide a way to identify a speciesFeatureType. Its value must be unique within its direct parent speciesType. When a speciesFeatureType is referenced by a speciesFeature, a SpeciesTypeComponentIndex object indexing the containing component can be used to avoid ambiguity.

**SpeciesFeatureType** also has an optional name attribute of type string.

#### 3.9.2 The occur attribute

**SpeciesFeatureType** has a required attribute occur, of type positiveInteger, used to indicate the number of instances of the **speciesFeatureType**. This attribute can be used to infer the number of the instances in *don't care* state with the use of the occur attribute in a referencing **speciesFeature** (also see Section 3.18.3 on page 31).

## 3.9.3 ListOfPossibleSpeciesFeatureValues

**ListOfPossibleSpeciesFeatureValues** is defined in Figure 4, and is extended from the **ListOf** class. A **listOfPossible-SpeciesFeatureValues** must have one or more **PossibleSpeciesFeatureValue** children. Since **ListOfPossible-SpeciesFeatureValues** is derived from **SBase** through **ListOf**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

# 3.10 PossibleSpeciesFeatureValue

PossibleSpeciesFeatureValue is defined in Figure 4 on the previous page, and is used to define the possible values a speciesFeature can take. It has a required attribute id and two optional attributes name and numericValue. Since PossibleSpeciesFeatureValue is derived from SBase, it inherits the sboTerm and metaid attributes, as well as the optional children Notes and Annotation objects.

## 3.10.1 The id and name attributes

The required id attribute, of type SId, serves to provide a way to identify a possibleSpeciesFeatureValue.

If the id of a possibleSpeciesFeatureValue is the content of a ci element in a MathML expression, it can either represent the numericValue ( when the ci has representationType="numericValue") or the count of the feature instances (default) which have this value.

PossibleSpeciesFeatureValue also has an optional name attribute of type string.

## 3.10.2 The numeric Value attribute

**PossibleSpeciesFeatureValue** has an optional attribute numericValue to be used to provide a reference to a numeric value that the **PossibleSpeciesFeatureValue** object can have. This attribute has type of SIdRef, and the value must be the identifier of a **Parameter** object in the model. The numeric value along with the unit can be defined in the **Parameter** object.

The modeler can either use the identifier of the parameter, or the identifier of the possibleSpeciesFeatureValue (with ci's representationType and speciesReference attribute) as the content of a ci element to represent its value in MathML expressions in SBML.

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# 3.11 SpeciesTypeInstance

SpeciesTypeInstance serves to provide a way to construct speciesTypes and species with multiple components. A speciesType can contain a list of instances of other speciesTypes which can also have their own speciesTypeInstances, so the complete construct of a speciesType has a tree structure. A speciesType cannot contain an instance of any other speciesType that already contains the instance of it. In other words, circular references are not allowed when constructing speciesTypes. For example, if a speciesType "A" contains the instance of another speciesType "B", "B" must not contain the instance of "A" anywhere in the complete structure of "B".

**SpeciesTypeInstance** is defined in Figure 5. It has two required attributes, id, and speciesType, and two optional attributes name and compartmentReference. Since **SpeciesTypeInstance** is derived from **SBase**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

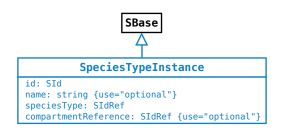


Figure 5: The definition of the SpeciesTypeInstance class

#### 3.11.1 The id and name attributes

The required attribute id, of type SId, serves to provide a way to identify a speciesTypeInstance. Its value must be unique within its direct parent speciesType.

**SpeciesTypeInstance** also has an optional name attribute of type string.

## 3.11.2 The speciesType attribute

The required attribute speciesType, of type SIdRef, is used to reference a speciesType.

#### 3.11.3 The compartmentReference attribute

The optional attribute compartmentReference, of type SIdRef, can be used to indicate which sub-compartment in a composite compartment the speciesTypeInstance is located in.

For example, a compartment "cA" has two sub-compartments "cB1" (referenced by compartmentReference "crB1") and "cB2" (referenced by compartmentReference "crB2") of the same compartment type "cB". A speciesType "stA" has two speciesTypeInstances "stiB1" and "stiB2" of the same speciesType "stB". The speciesType "stA" references the compartment "cA" and the speciesType "stB" references the compartment "cB". The speciesTypeInstance "stiB1" is located in "cB1" via the compartmentReference "crB1" and the speciesTypeInstance "stiB2" is located in "cB2" via the compartmentReference "crB2". The SBML code can be as follows:

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# 3.12 SpeciesTypeComponentIndex

SpeciesTypeComponentIndex provides a way to identify or index a component within a speciesType. A SpeciesTypeComponentIndex object can be referenced by other class objects, such as InSpeciesTypeBond, Outward-BindingSite, SpeciesFeature or SpeciesTypeComponentMapInProduct objects, which needs to identify a component in a particular speciesType.

**SpeciesTypeComponentIndex** is defined in Figure 6. It has two required attributes, id, and component, and two optional attributes name and identifyingParent. Since **SpeciesTypeComponentIndex** is derived from **SBase**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

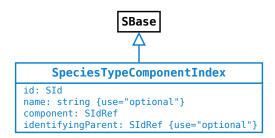


Figure 6: The definition of the SpeciesTypeComponentIndex class

See Section 3.16.3 on page 28 about how to use SpeciesTypeComponentIndex in an outwardBindingSite.

Note:

A speciesTypeComponentIndex should be unambiguous. For example, a speciesTypeComponentIndex should not reference to a speciesType which is referenced by two speciesTypeInstances contained in the same SpeciesType object.

#### 3.12.1 The id and name attributes

The id attribute, of type SId, provides a way to identify a speciesTypeComponentIndex. The value must be unique within the direct parent speciesType. SpeciesTypeComponentIndex also has an optional name attribute of type string.

#### 3.12.2 The component attribute

The component attribute, of type of SIdRef, references a speciesTypeInstance in the speciesType, or the speciesType itself. The value of this attribute can be the id of a speciesTypeInstance or a speciesType-ComponentIndex that is defined in the speciesType of a speciesTypeInstance.

## 3.12.3 The identifyingParent attribute

The component attribute itself may not be sufficient to uniquely reference a component in a speciesType. The identifyingParent attribute provides assistance for the identification of a component. It references a parent of the component and the value can be the id of an object of SpeciesTypeInstance, SpeciesTypeComponentIndex or SpeciesType.

This example illustrates the use of the identifyingParent attribute. There are three speciesTypes "stA", "stB" and "stC". The speciesType "stB" contains two speciesTypeInstances "C1" and "C2" of the same speciesType "stC". The speciesType "stA" contains two speciesTypeInstances "B1" and "B2" of the same speciesType "stB". The speciesType "A" may be required to index every "C1" and "C2" by its ListOfInSpeciesTypeBonds child or referencing species. The following SBML code demonstrates how to do the indexing with assistance from the identifyingParent attribute.

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```
<multi:listOfSpeciesTypes>
   <multi:speciesType multi:id="stC" ... />
<multi:speciesType multi:id="stB" ... >
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="C1" multi:speciesType="stC" />
         <multi:speciesTypeInstance multi:id="C2" multi:speciesType="stC" />
      </multi:listOfSpeciesTypeInstances>
   </multi:speciesType>
   <multi:speciesType multi:id="stA" ... >
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="B1" multi:speciesType="stB" />
         <multi:speciesTypeInstance multi:id="B2" multi:speciesType="stB" />
      </multi:listOfSpeciesTypeInstances>
      <multi:listOfSpeciesTypeComponentIndexes>
         <multi:speciesTypeComponentIndex multi:id="B1C1"</pre>
            multi:component="C1" multi:identifyingParent="B1" />
         <multi:speciesTypeComponentIndex multi:id="B1C2"</pre>
            multi:component="C2" multi:identifyingParent="B1" />
         <multi:speciesTypeComponentIndex multi:id="B2C1"</pre>
            multi:component="C1" multi:identifyingParent="B2" />
         <multi:speciesTypeComponentIndex multi:id="B2C2"</pre>
            multi:component="C2" multi:identifyingParent="B2" />
      </multi:listOfSpeciesTypeComponentIndexes>
   </multi:speciesType>
</multi:listOfSpeciesTypes>
```

In the speciesType "stA", "B1C1" identifies the "C1" in "B1" and "B2C1" identifies the "C1" in "B2". Similarly, "B1C2" identifies the "C2" in "B1" and "B2C2" identifies "C2" in "B2".

#### 3.12.4 Reference a component in a speciesType or a species

In the Multi package, component(s) of a speciesType (or a species via its speciesType attribute) can be referenced by objects of multiple classes such as **OutwardBindingSite** and **SpeciesFeature**. A component of a speciesType can be a speciesTypeInstance or the speciesType itself. For example:

```
<multi:listOfSpeciesTypes>
   <multi:bindingSiteSpeciesType multi:id="stA" ... />
   <multi:speciesType multi:id="stB" ...>
   </multi:speciesType>
   <multi:speciesType multi:id="stABB" ...>
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="stiA" multi:speciesType="stA" .../>
         <multi:speciesTypeInstance multi:id="stiB1" multi:speciesType="stB" .../>
         <multi:speciesTypeInstance multi:id="stiB2" multi:speciesType="stB" .../>
      </multi:listOfSpeciesTypeInstances>
   </multi:speciesType>
</multi:listOfSpeciesTypes>
<listOfSpecies>
   <species id="spA" multi:speciesType="stA" ...>
      <multi:listOfOutwardBindingSites>
         <multi:outWardBindingSite multi:component="stA" multi:bindingStatus="bound" />
      </multi:listOfOutwardBindingSites>
  </species>
   <species id="spABB" multi:speciesType="stABB" ...>
      <multi:listOfOutwardBindingSites>
         <multi:outWardBindingSite multi:component="stiA" multi:bindingStatus="bound" />
      </multi:listOfOutwardBindingSites>
    </species>
</listOfSpecies>
```

In this example, the component of the outwardBindingSite in species "spABB" is a speciesTypeInstance ("spABB"), and the component of the outwardBindingSite in species "spA" is a speciesType ("stA") which is directly referenced by the speciesType attribute of "spA".

In many cases, to reference a component, the id of the component will be sufficient and it is not necessary to create an index (speciesTypeComponentIndex). The example in Section 3.12.3 on page 20 illustrates two equivalent ways to reference a component, for example, the "B1" component in the "stA" speciesType. The creation of a speciesTypeComponentIndex cannot be avoided when a speciesType (indirectly) has two speciesTypeInstances that have the same id.

# 3.13 InSpeciesTypeBond

An **InSpeciesTypeBond** object defines a bond existing within a **speciesType**. The bond therefore exists in every **species** that references the **speciesType**.

**InSpeciesTypeBond** is defined in Figure 7. It has two optional attributes, **id** and **name**, and two required attributes, **bindingSite1** and **bindingSite2**. Since **InSpeciesTypeBond** is derived from **SBase**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

The binding relationship in an inSpeciesTypeBond is one-to-one (see Section 3.8.7 on page 15). The uniqueness of an inSpeciesTypeBond is ensured by the pair of referenced bindingSites. A speciesType cannot have two inSpeciesTypeBonds containing the same pair of bindingSites.

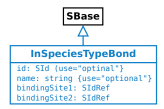


Figure 7: The definition of the InSpeciesTypeBond class

#### 3.13.1 The id and name attributes

The optional **id** attribute, of type SId, provides a way to identify an **inSpeciesTypeBond**. If present, the value of the **id** attribute must be unique within its direct parent **speciesType**.

**InSpeciesTypeBond** also has an optional **name** attribute of type string.

## 3.13.2 The bindingSite1 and bindingSite2 attributes

InSpeciesTypeBond has two required attributes, bindingSite1 and bindingSite2, both of type SIdRef, used to reference a pair of binding sites of the InSpeciesTypeBond object in a speciesType. The referenced identifiers of the binding sites can be the ids of the speciesTypeInstances (binding sites), or the ids of the speciesTypeComponent-Indexes indexing the binding sites and the ultimately referenced components must be the BindingSiteSpeciesType objects. Obviously, bindingSite1 and bindingSite2 must not reference the same BindingSiteSpeciesType object.

## 3.14 Uniqueness of SpeciesType definitions

In some special cases, it may be possible to define a speciesType in multiple equivalent ways.

Figure 8 shows an example of a speciesType constructed in two different formats. The two "st\_x" speciesTypes in the diagram can be the results of different reaction paths, but they are equivalent and define the same speciesType.

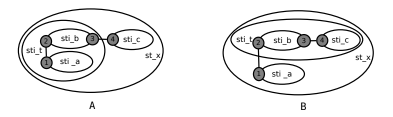


Figure 8: Different formats of the same speciesType

Construct 1: The definition of speciesType "st\_x" on the left (A) in Figure 8.

```
<multi:listOfSpeciesTypes>
   <multi:bindingSiteSpeciesType multi:id="st1" />
<multi:bindingSiteSpeciesType multi:id="st2" />
   <multi:bindingSiteSpeciesType multi:id="st3" />
<multi:bindingSiteSpeciesType multi:id="st3" />
<multi:bindingSiteSpeciesType multi:id="st3" />
   <multi:bindingSiteSpeciesType multi:id="st4" />
   <multi:speciesType multi:id="st_a">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="_1" multi:speciesType="st1" />
      </multi:listOfSpeciesTypeInstances>
   </multi:speciesType>
   <multi:speciesType multi:id="st_b">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="_2" multi:speciesType="st2" />
          <multi:speciesTypeInstance multi:id="_3" multi:speciesType="st3" />
      </multi:listOfSpeciesTypeInstances>
   </multi:speciesType>
   <multi:speciesType multi:id="st_c">
      <multi:listOfSpeciesTypeInstances>
          <multi:speciesTypeInstance multi:id="_4" multi:speciesType="st4" />
      </multi:listOfSpeciesTypeInstances>
   </multi:speciesType>
   <multi:speciesType multi:id="st_t">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="sti_a" multi:speciesType="st_a" />
         <multi:speciesTypeInstance multi:id="sti_b" multi:speciesType="st_b" />
      </multi:listOfSpeciesTypeInstances>
      <multi:listOfInSpeciesTypeBonds>
         <multi:inSpeciesTypeBond multi:bindingSite1="_1" multi:bindingSite2="_2" />
      </multi:listOfInSpeciesTypeBonds>
   </multi:speciesType>
   <multi:speciesType multi:id="st_x">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="sti_t" multi:speciesType="st_t" />
         <multi:speciesTypeInstance multi:id="sti_c" multi:speciesType="st_c" />
      </multi:listOfSpeciesTypeInstances>
      <multi:listOfInSpeciesTypeBonds>
         <multi:inSpeciesTypeBond multi:bindingSite1="_3" multi:bindingSite2="_4" />
      </multi:listOfInSpeciesTypeBonds>
   </multi:speciesType>
```

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#### </multi:listOfSpeciesTypes>

Construct 2: The definition of speciesType "st\_x" on the right (B) in Figure 8 on the preceding page.

```
<multi:listOfSpeciesTypes>
   <multi:bindingSiteSpeciesType multi:id="st1" />
   <multi:bindingSiteSpeciesType multi:id="st2"</pre>
   ~multi:bindingSiteSpeciesType multi:id="st2" />
<multi:bindingSiteSpeciesType multi:id="st3" />
<multi:bindingSiteSpeciesType multi:id="st3" />
   <multi:bindingSiteSpeciesType multi:id="st4" />
   <multi:speciesType multi:id="st_a">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="_1" multi:speciesType="st1" />
      </multi:listOfSpeciesTypeInstances>
   </multi:speciesType>
   <multi:speciesType multi:id="st_b">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="_2" multi:speciesType="st2" />
         <multi:speciesTypeInstance multi:id="_3" multi:speciesType="st3" />
      </multi:listOfSpeciesTypeInstances>
   </multi:speciesType>
   <multi:speciesType multi:id="st_c">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="_4" multi:speciesType="st4" />
      </multi:listOfSpeciesTypeInstances>
   </multi:speciesType>
   <multi:speciesType multi:id="st_t">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="sti_b" multi:speciesType="st_b" />
         <multi:speciesTypeInstance multi:id="sti_c" multi:speciesType="st_c" />
      </multi:listOfSpeciesTypeInstances>
      <multi:listOfInSpeciesTypeBonds>
         <multi:inSpeciesTypeBond multi:bindingSite1="_3" multi:bindingSite2="_4" />
      </multi:listOfInSpeciesTypeBonds>
   </multi:speciesType>
   <multi:speciesType multi:id="st_x">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="sti_a" multi:speciesType="st_a" />
         <multi:speciesTypeInstance multi:id="sti_t" multi:speciesType="st_t" />
      </multi:listOfSpeciesTypeInstances>
      <multi:listOfInSpeciesTypeBonds>
         <multi:inSpeciesTypeBond multi:bindingSite1="_1" multi:bindingSite2="_2" />
      </multi:listOfInSpeciesTypeBonds>
   </multi:speciesType>
</multi:listOfSpeciesTypes>
```

This kind of ambiguity cannot be avoided for speciesTypes involving more than two subcomponents connected by inSpeciesTypeBonds. A and B in Figure 8 on the previous page can be products of different association reactions. It is up to the modeler (parser) to identify whether the two speciesTypes such as those in the example above are identical.

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3.15 Species

A species in SBML Level 3 Core refers to a pool of entities. A species in the Multi package is extended from a pool to a template or pattern to which multiple pools may map. An extended species can reference a speciesType that provides the backbone for the species such as components (including binding sites) and speciesFeature-Types. When referencing a speciesType, a species can be further defined with regard to the binding statuses of its outwardBindingSites and the speciesFeatures. With the options to have variable values selected, such as "either" for the bindingStatus attribute and multiple possibleSpeciesFeatureValues for a speciesFeature, an extended species can work as a template or pattern how species participate in reactions.

The extension of the **Species** class is illustrated in Figure 9. The extended **Species** class has a new optional attribute speciesType, and two extra optional **ListOfOutwardBindingSites** and **ListOfSpeciesFeatures** children. A species may have a **listOfOutwardBindingSites** child and/or a **listOfSpeciesFeatures** child only when its speciesType attribute has been defined.

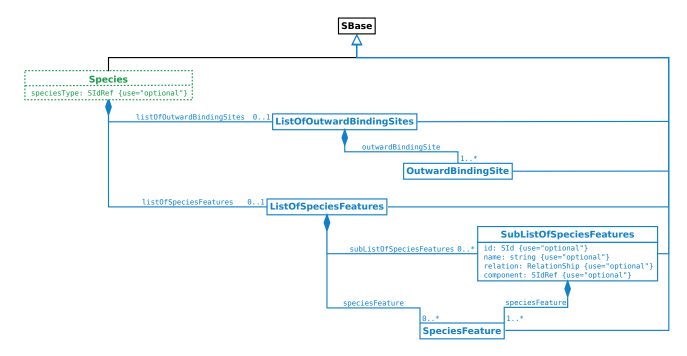


Figure 9: The extension of the Species class

#### 3.15.1 The speciesType attribute

The optional attribute speciesType, of type SIdRef, references a SpeciesType object.

#### 3.15.2 ListOfOutwardBindingSites

**ListOfOutwardBindingSites** is defined in Figure 9 and is extended from the **ListOf** class. A **listOfOutwardBinding-Sites** can only be defined when the **speciesType** attribute is defined. If present, it must have one or more **OutwardBindingSite** children. Since **ListOfOutwardBindingSites** is derived from **SBase** through **ListOf**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

Note:

The listOfOutwardBindingSites of a species is not necessary to list all the outwardBindingSites (the binding sites not involved in any inSpeciesTypeBond) defined by the referenced speciesType. If an outwardBindingSite

is not listed in the list0f0utwardBindingSites, the value of its bindingStatus is "either". In other words, the binding site is in a don't care state.

## 3.15.3 ListOfSpeciesFeatures

**ListOfSpeciesFeatures** is defined in Figure 9 on the preceding page and is extended from the **ListOf** class. A **list-OfSpeciesFeatureTypes** can only be defined when the **speciesType** attribute is defined. If present, it must have one or more children. A child can be a **SpeciesFeature**, or a **SubListOfSpeciesFeatures** object.

Note:

The listOfSpeciesFeatures of a species does not have to cover all the speciesFeatures corresponding to all speciesFeatureTypes (see Section 3.9 on page 16) of every component defined by the referenced speciesType. If a speciesFeatureType is defined and there is no speciesFeature explicitly referencing it, the species has an implicit speciesFeature having all the listOfPossibleSpeciesFeatureValues and "or" relationships between them. In other words, the implicit speciesFeature has a don't care state for the species.

Since **ListOfSpeciesFeatures** is derived from **SBase** through **ListOf**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

The example at Section 3.18.7 on page 31 illustrates the usage of the ListOfSpeciesFeatures class.

# 3.16 OutwardBindingSite

OutwardBindingSite is defined in Figure 10. It has two optional attributes, id and name, and two required attributes, bindingStatus and component. A binding site not involved in any InSpeciesTypeBond object in the species-Type referenced by a species is an outwardBindingSite. Since OutwardBindingSite is derived from SBase, it inherits the sboTerm and metaid attributes, as well as the optional children Notes and Annotation objects.

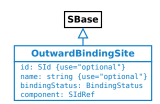


Figure 10: The definition of the OutwardBindingSite class

#### 3.16.1 The id and name attributes

The optional id attribute, of type SId, can serve to provide a way to identify an outwardBindingSite. If present, the value must be unique within the species. OutwardBindingSite also has an optional name attribute of type string.

## 3.16.2 The bindingStatus attribute

The bindingStatus attribute takes a value of type **BindingStatus**.

#### 3.16.3 The component attribute

The component attribute, of type SIdRef, references a component which ultimately reference a **BindingSiteSpecies- Type** object. The attribute value must be the identifier of a **SpeciesTypeInstance**, **SpeciesTypeComponentIndex** or **SpeciesType** object. An **outwardBindingSite** cannot be a binding site referenced by any **inSpeciesTypeBond** in the **species**.

There are three scenarios for the **component** attribute to have the value of an identifier of **SpeciesType**, **SpeciesTypeInstance**, or **SpeciesTypeComponentIndex** respectively.

- (1) When a species references a simple bindingSiteSpeciesType, the value of the component attribute of the outwardBindingSite of the species can only be the id of the referenced speciesType.
- (2) When a species references a speciesType with a speciesTypeInstance being a binding site (have an id of BindingSiteSpeciesType as its speciesType attribute) and the id of the speciesTypeInstance can identify the binding site within the speciesType (referenced by the species) unambiguously, and therefore, the value of the component attribute of an outwardBindingSite of the species can be the id of the species-TypeInstance.
- (3) When a species references a speciesType with a speciesTypeInstance being a binding site (directly or indirectly) and id of the speciesTypeInstance can NOT identify the binding site without ambiguity, an id of SpeciesTypeComponentIndex can be used as the value of the component attribute of an outwardBindingSite of the species.

#### 3.16.4 Example

Figure 11 on the following page illustrates the usage of the OutwardBindingSite class. Species "sp\_x" references speciesType "st\_x", which has three speciesTypeInstances "sti\_a", "sti\_b" and "sti\_c". SpeciesTypeInstance "sti\_a" has bindingSites "\_1" and "\_3", speciesTypeInstance "sti\_b" has bindingSites "\_2" and "\_4", and speciesTypeInstance "sti\_c" has bindingSite "\_5". The inSpeciesTypeBondin "st\_x" involves two bindingSites

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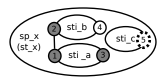


Figure 11: An example of OutwardBindingSite

"\_1" and "\_2". The other three bindingSites, "\_3", "\_4" and "\_5", in the species "sp\_x" are outwardBindingSites. The outwardBindingSite "\_3" is "bound" (filled circle with solid line in the diagram), the outwardBindingSite "\_4" is "unbound" (empty circle with solid line) and the outwardBindingSite "\_5" has binding status "either" (empty circle with dotted line). The corresponding SBML code would be as follows:

```
<multi:listOfSpeciesTypes>
   <multi:bindingSiteSpeciesType multi:id="st_1" />
   <multi:bindingSiteSpeciesType multi:id="st_2" />
   <multi:bindingSiteSpeciesType multi:id="st_3" />
   <multi:bindingSiteSpeciesType multi:id="st_4" />
   <multi:bindingSiteSpeciesType multi:id="st_5" />
   <multi:speciesType multi:id="st_a">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="_1" multi:speciesType="st_1" />
         <multi:speciesTypeInstance multi:id="_3" multi:speciesType="st_3" />
      </multi:listOfSpeciesTypeInstances>
  </multi:speciesType>
   <multi:speciesType multi:id="st_b">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="_2" multi:speciesType="st_2" />
         <multi:speciesTypeInstance multi:id="_4" multi:speciesType="st_4" />
      </multi:listOfSpeciesTypeInstances>
  </multi:speciesType>
   <multi:speciesType multi:id="st_c">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="_5" multi:speciesType="st_5" />
      </multi:listOfSpeciesTypeInstances>
  </multi:speciesType>
   <multi:speciesType multi:id="st_x">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="sti_a" multi:speciesType="st_a" />
         <multi:speciesTypeInstance multi:id="sti_b" multi:speciesType="st_b" />
         <multi:speciesTypeInstance multi:id="sti_c" multi:speciesType="st_c" />
      </multi:listOfSpeciesTypeInstances>
      <multi:listOfInSpeciesTypeBonds>
         <multi:inSpeciesTypeBond multi:bindingSite1="_1" multi:bindingSite2="_2" />
      </multi:listOfInSpeciesTypeBonds>
  </multi:speciesType>
</multi:listOfSpeciesTypes>
tofSpecies>
   <species id="sp_x" multi:speciesType="st_x">
      <multi:listOfOutwardBindingSites>
         <multi:outwardBindingSite multi:component="_3" multi:bindingStatus="bound" />
         <multi:outwardBindingSite multi:component="_4" multi:bindingStatus="unbound" />
         <multi:outwardBindingSite multi:component="_5" multi:bindingStatus="either" />
      </multi:listOfOutwardBindingSites>
   </species>
</listOfSpecies>
```

# 3.17 SubListOfSpeciesFeatures

**SubListOfSpeciesFeatures** is defined in Figure 9 on page 26, and is extended from the **ListOf** class. If present, a **subListOfSpeciesFeatures** must have one or more **SpeciesFeature** children. Since **SubListOfSpeciesFeatures** is

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derived from **SBase** through **ListOf**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

#### 3.17.1 The id and name attributes

The optional id attribute, of type SId, can serve to provide a way to identify a subListOfSpeciesFeatures. If present, the value must be unique within the species. SubListOfSpeciesFeatures also has an optional name attribute of type string.

#### 3.17.2 The relation attribute

**SubListOfSpeciesFeatures** has an optional attribute relation, of type **Relation**, to define the logic relationship among its children. The relation attribute cannot be defined if a **subListOfSpeciesFeatures** has only one child.

## 3.17.3 The component attribute

The optional component attribute, of type SIdRef, can be used to indicate which **component** of a species the subListOfSpeciesFeatures belongs to. It is required when the component of any speciesFeature contained in this subListOfSpeciesFeatures cannot be identified only based on its speciesFeatureType attribute.

## 3.18 SpeciesFeature

SpeciesFeature is defined in Figure 12. It has three optional attributes, id, name and component, and two required attributes, speciesFeatureType and occur, and a required child listOfSpeciesvFeatureValues. Since SpeciesFeature is derived from SBase, it inherits the sboTerm and metaid attributes, as well as the optional children Notes and Annotation objects. SpeciesFeature serves to define the state of a component in a species by selecting values from the listOfPossibleSpeciesFeatureValues of the referenced speciesFeatureType.

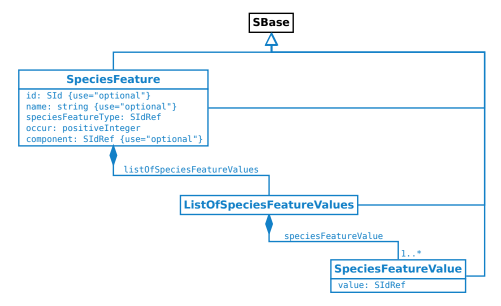


Figure 12: The definitions of the SpeciesFeature class and the SpeciesFeatureValue class

#### 3.18.1 The id and name attributes

The optional **id** attribute, of type SId, can serve to provide a way to identify a **speciesFeature**. If present, the value must be unique within the **species. SpeciesFeature** also has an optional **name** attribute of type string.

#### 3.18.2 The speciesFeatureType attribute

**SpeciesFeature** has a required attribute **speciesFeatureType**, of type SIdRef, used to reference a **speciesFeature**-Type.

#### 3.18.3 The occur attribute

**SpeciesFeature** has a required attribute **occur**, of type of **positiveInteger**, used to define the number of instances of the referenced **speciesFeatureType**.

The value of the occur attribute cannot be larger than the occur of the referenced speciesFeatureType. When a speciesFeatureType has multiple instances (speciesFeatureType's occur > "1"), the speciesFeature's occur attribute provides a way for a species to define the instances of the speciesFeatureType differently.

For example, in a speciesType, speciesFeatureType "ftA" has occur="2" and two possibleSpeciesFeature-Values "fva1" and "fva2". A species referencing the speciesType can be defined to have two speciesFeatures "sfA1" and "sfA2" both referencing "ftA". The speciesFeature "sfA1" has occur="1" and its value is "fva1". The speciesFeature "sfA2" has occur="1" and its value is "fva2".

If the occur of a speciesFeature is less than the occur of the referenced speciesFeatureType, the rest of the unspecified instances of the speciesFeatureType are in *don't care* state which means that the value of an unspecified instance can be any from the listOfPossibleSpeciesFeatureValues.

For example, in a speciesType, a speciesFeatureType "phosphorylation" has two possibleSpeciesFeature-Values "phosphorylated" and "unphosphorylated" and the occur is "5". A species referencing the speciesType can be defined to have a speciesFeature of the "phosphorylation" with the value of "phosphorylated" and the occur of "1". Then, the species is a pattern species with at least one "phosphorylated" site (the other four "phosphorylation" sites are in *don't care* state). This pattern species can be mapped by anyone of the *fully defined* species (see Section 3.19 on page 33) of the same type and with any of "1" to "5" phosphorylated sites.

## 3.18.4 The component attribute

The optional component attribute, of type SIdRef, can be used to indicate which component of a species the speciesFeature belongs to. It is required when the component cannot be identified only based on the speciesFeatureType attribute.

## 3.18.5 ListOfSpeciesFeatureValues

**ListOfSpeciesFeatureValues** is defined in Figure 12 on the preceding page, and is extended from the **ListOf** class. A listOfSpeciesvFeatureValues must have one or more **SpeciesFeatureValue** children. If a listOfSpecies-Features has multiple speciesFeatureValues, the interpretation of the relationship between them is "or". Since **ListOfSpeciesFeatureValues** is derived from **SBase** through **ListOf**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

#### 3.18.6 SpeciesFeatureValue

SpeciesFeatureValue is defined in Figure 12 on the previous page. A speciesFeatureValue serves to specify a value for a speciesFeature to select from the listOfPossibleSpeciesFeatureValues defined in the referenced speciesFeatureType. The SpeciesFeatureValue class has only one attribute value, of type SIdRef, used to reference a PossibleSpeciesFeatureValue object. Since SpeciesFeatureValue is derived from SBase, it inherits the sboTerm and metaid attributes, as well as the optional children Notes and Annotation objects.

#### 3.18.7 Example

Figure 13 on the following page is an example speciesType to illustrate the usage of the ListOfSpeciesFeatures, Sub-ListOfSpeciesFeatures and SpeciesFeature classes. SpeciesType "st\_X" has speciesTypeInstance "sti\_A" with speciesFeatureType "fA", and speciesTypeInstance "sti\_B" with speciesFeatureTypes "fB1" and "fB2". The

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speciesFeatureType "fA" has two possibleSpeciesFeatureValues "v1" and "v2". The speciesFeatureType "fB1" has "v3" and "v4", and "fB2" has "v5" and "v6". Here are several ways to construct the listOfSpeciesFeaturesof a species referencing the speciesType "st\_A":

```
listOfSpeciesFeatures("fA"="v1", "fB1"="v3", "fB2"="v5") is a state:
    "[fA=v1] AND [fB1=v3] AND [fB2=v5]"

listOfSpeciesFeatures("fA"="v1", "fB1"="v3") is a state:
    "[fA=v1] AND [fB1=v3] AND ([fB2=v5] OR fB2=v6)"
    "fB2" has a value of don't care

listOfSpeciesFeatures(
    "fA=v1",
    subListOfSpeciesFeatures ("fB1=v3", "fB2=v5", relation="not")
) is a state:
    "[fA=v1] and [fB1=v4] and [fB2=V5]" or
    "[fA=v1] and [fB1=v4] and [fB2=V6]" or
    "[fA=v1] and [fB1=v3] and [fB2=V6]"
```

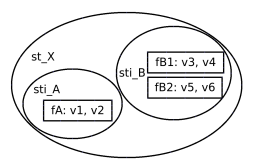


Figure 13: An example speciesFeatureType to illustrate the usage of ListOfSpeciesFeatures, SubListOfSpeciesFeatures and the SpeciesFeature

The SBML code can be as follows and the species "sp\_A1", "sp\_A2" and "sp\_A3" contain the tree listOfSpecies-Features above respectively.

# 3.19 Fully defined species and mapping to pattern species

An extended **Species** object functions as a template or a pattern which allows multiple pools of entities to map to it. A **species** is *fully defined* if there is only one pool mapping to it. A *fully defined* **species** can be considered the same as an SBML core **species**, and can be initialized with the **initialAmount** attribute, or the **initial-Concentration** attribute, or via an **InitialAssignment** object. In the Multi package, a **species** is *fully defined* if the following conditions are fulfilled:

- All outwardBindingSites must be free (bindingStatus="unbound"), since "bound" sites imply that there is a non-specified binding partner.
- Each speciesFeature occurrence can only have one speciesFeatureValue, and every occurrence of every speciesFeatureType of every component of the referenced speciesType must be referenced by exactly one speciesFeature occurrence.
- If applicable, only "and" values are allowed for the relation attributes of the SubListOfSpeciesFeatures objects.
- Only one single **SpeciesFeatureValue** object is allowed for any **speciesFeature**.
- The referenced compartment cannot be a compartment type, which means the value of the isType attribute of the referenced compartment can only be "false".

The mapping from a *fully defined* species to a *pattern* species is implicit and can be inferred from the structure of the species. For example, a speciesType "stA" has one speciesFeatureType with two possibleSpeciesFeatureValues "v1" and "v2". The species "spA1" references "stA" and has the speciesFeature with the value of "v1". Another species "spA" also references "stA" and has no speciesFeature explicitly defined. Thus, the species "spA1" is a fully defined species and can map to the *pattern* species "spA" because species "spA" has an implicit speciesFeature which can take either value "v1" or value "v2" (see the note in Section 3.15.3 on page 27).

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3.20 Reaction

Reaction itself in the Multi package is not extended, but it may use the Multi Species objects to construct reactions. The Reaction class in the Multi package cannot only define the relations among pools (SBML core species), but also the relations among patterns (Multi extended species). Several related classes including SimpleSpecies-Reference and SpeciesReference are extended to handle some issues specific to the Multi package. A new class, IntraSpeciesReaction, is derived from Reaction to explicitly define those reactions within the same Species object.

The changes under the **Reaction** class in the Multi package are illustrated in Figure 14.

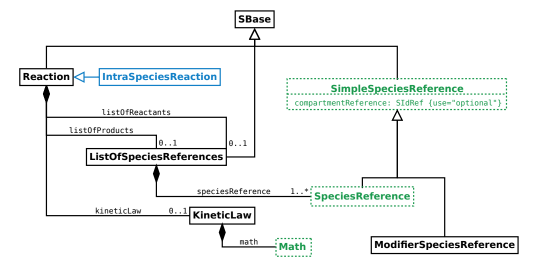


Figure 14: The changes under the Reaction class including IntraSpeciesReaction, SimpleSpeciesReference, Species-Reference and Math

## 3.21 IntraSpeciesReaction

**IntraSpeciesReaction** is derived from **Reaction** for the reactions happening within a species (see the example "Extended Reaction class" at page 23 of the slides at HARMONY 2013 [Zhang and Meier-Schellersheim (2013b)]).

A particular reaction may happen within a species as an intraSpeciesReaction if the following conditions are fulfilled:

- The reaction is either an association reaction or a dissociation reaction.
- If it is an association reaction, each of the two reactant species has at least one outwardBindingSite free ("unbound").
- If it is a dissociation reaction, each of the two product species has at least one outwardBindingSite free ("unbound").

#### Note:

Technically, transformations are also reactions happening with one species, but they do not have the ambiguity of association and dissociation reactions. Therefore, transformation reactions do not have to be defined as intra-SpeciesReactions.

# 3.22 Extended SimpleSpeciesReference

The **SimpleSpeciesReference** class is extended with a new optional attribute **compartmentReference**, of type SIdRef, to reference a **compartmentReference**. The **compartmentReference** attribute can serve to indicate in which sub-compartment an object of a class (**SpeciesReference** or **ModifierSpeciesReference**) inheriting **Simple-SpeciesReference** is located.

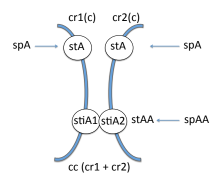


Figure 15: Reaction: spA(cr1) + spA(cr2) -> spAA

This example illustrates the use of the compartmentReference attribute in simpleSpeciesReferences. The situation described here could correspond to interactions among species located on two adjacent membranes. A model has a composite compartment "cc" with two compartmentReferences "cr1" and "cr2", and both reference "c" subcomponents. Species "spA" references compartment "c", and species "spAA" references the composite compartment "cc". A reaction happens between two "spA" species from the two "c" compartments and results in a cross-compartment product. The SBML code can be as follows:

```
<listOfCompartments>
   <compartment id="c" constant="true" multi:isType="true" />
   <compartment id="cc" constant="true" multi:isType="true">
      <multi:listOfCompartmentReferences>
         <multi:compartmentReference multi:id="cr1" multi:compartment="c" />
         <multi:compartmentReference multi:id="cr2" multi:compartment="c" />
      </multi:listOfCompartmentReferences>
   </compartment>
</listOfCompartments>
<multi:listOfSpeciesTypes>
   <multi:bindingSiteSpeciesType multi:id="stA" multi:compartment="c" />
   <multi:speciesType multi:id="stAA" multi:compartment="cc">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="stiA1" multi:speciesType="stA"
            multi:compartmentReference="cr1" />
         <multi:speciesTypeInstance multi:id="stiA2" multi:speciesType="stA"</pre>
            multi:compartmentReference="cr2" />
      </multi:listOfSpeciesTypeInstances>
      <multi:listOfInSpeciesTypeBonds>
         <multi:inSpeciesTypeBond multi:bindingSite1="stiA1" multi:bindingSite2="stiA2" />
      </multi:listOfInSpeciesTypeBonds>
  </multi:speciesType>
</multi:listOfSpeciesTypes>
st0fSpecies>
   <species id="spA" multi:speciesType="stA" compartment="c" ... />
   <species id="spAA" multi:speciesType="stAA" compartment="cc" ... />
</listOfSpecies>
<listOfReactions>
```

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## 3.23 Extended SpeciesReference

The **SpeciesReference** class is extended from SBML Level 3 Core and can establish **component** mappings between the reactant **species** and the product **species** when the mappings cannot be inferred from the **id**s of the **Species-TypeInstance** objects. The **SpeciesReference** class has an optional **ListOfSpeciesTypeComponentMapsInProduct** child, as defined in Figure 16. Only a **reaction product** can contain the **ListOfSpeciesTypeComponentMapsIn-Product** child and it is not necessary to store the mappings again in the **reactants**.

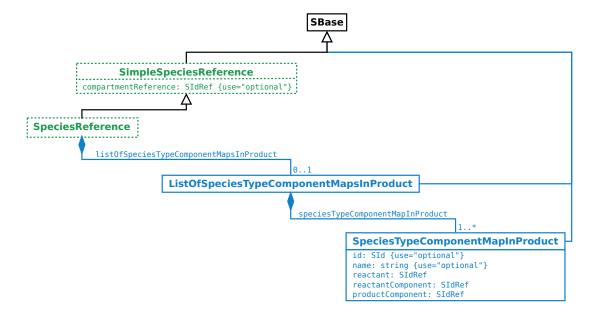


Figure 16: The extension of the SpeciesReference class

#### 3.23.1 ListOfSpeciesTypeComponentMapsInProduct

**ListOfSpeciesTypeComponentMapsInProduct** is defined in Figure 16, and is extended from the **ListOf** class. If present, a listOfSpeciesTypeComponentMapsInProduct must have one or more **SpeciesTypeComponentMapsInProduct** children. Since **ListOfSpeciesTypeComponentMapsInProduct** is derived from **SBase** through **ListOf**, it inherits the **sboTerm** and **metaid** attributes, as well as the optional children **Notes** and **Annotation** objects.

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### 3.24 SpeciesTypeComponentMapInProduct

SpeciesTypeComponentMapInProduct is defined in Figure 16 on the previous page. It has two optional attributes, id and name, and three required attributes, reactant, reactantComponent and productComponent. Since SpeciesTypeComponentMapInProduct is derived from SBase, it inherits the sboTerm and metaid attributes, as well as the optional children Notes and Annotation objects.

A speciesTypeComponentMapInProduct defines the mapping between a component in a reactant and a component in a product. The identifications of a **component** and the speciesReference should be sufficient to identify the component in the context of a reaction. The attributes reactant and reactantComponent can identify the component in a reactant, and the productComponent attribute and the product storing the mapping information can identify the component in a product.

#### 3.24.1 The id and name attributes

The optional id attribute, of type SId, can serve to provide a way to identify an speciesTypeComponentMapIn-Product. If present, the value must be unique within the reaction. SpeciesTypeComponentMapInProduct also has an optional name attribute of type string.

#### 3.24.2 The reactant attribute

**SpeciesTypeComponentMapInProduct** has a required **reactant** attribute, of type SIdRef, to reference the **id** of a reactant **speciesReference** in a **reaction**.

#### 3.24.3 The reactantComponent attribute

**SpeciesTypeComponentMapInProduct** has a required **reactantComponent** attribute, of type SIdRef, to reference a **component** in a reactant **species**.

#### 3.24.4 The productComponent attribute

**SpeciesTypeComponentMapInProduct** has a required **productComponent** attribute, of type SIdRef, to reference a **component** in a product **species**.

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# 3.25 OutwardBindingSites and speciesFeatures in *don't care* state in reaction products

An outwardBindingSite is in *don't care* state if its bindingStatus is "either" or is not specified (also see Section 3.15.2 on page 26). A speciesFeature is in *don't care* state if it is not specified in the referencing species (also see Section 3.18 on page 30).

For a species as a product in a reaction, if it has don't care outwardBindingSites or don't care speciesFeatures, the interpretation of the don't care is don't change. In a product, a don't care outwardBindingSite has the same bindingStatus as the mapped outwardBindingSite in the reactant, and a don't care speciesFeature has the same value as the mapped speciesFeature in the reactant.

#### 3.26 Extended ci elements in Math objects

The Multi package extends the ci element in **Math** in **Reaction** with optional attributes **speciesReference** and **representationType**.

#### 3.26.1 The speciesReference attribute

The optional speciesReference attribute, of type SIdRef, can only be used when the content of the ci element is a species id, or when the content of the ci element is a speciesFeature id. The speciesReference attribute can identify which species is referenced in a reaction, and the speciesReference attribute must have a value of a speciesReference id within the same reaction.

If the ci content references a species' id, the id represents the concentration or amount of the species.

If the ci content references a speciesFeature's id, the id represents the count of the speciesFeature instances with the speciesFeatureValue (also see Section 3.18.1 on page 30).

The example in Section 3.22 on page 35 can be further extended with a block of kineticLaw in the reaction to illustrate the use of the speciesReference attribute with a species' id.

Two "spA" species are distinguished by the "r1" and "r2" speciesReferences respectively.

#### 3.26.2 The representationType attribute

The optional representationType attribute, of type RepresentationType, can only be used when the content of the ci element is a species' id or a possibleSpeciesFeatureValue's id. The representationType and speciesReference attributes can both be used for the same ci element at the same time.

The representationType attribute can only have the value of "sum" when the content of the ci is the id of a species. The interpretation of such a ci element is the total concentration or amount of all *fully defined* species (see Section 3.19 on page 33) mapping to the referenced pattern species.

The representationType attribute can have the value of numericValue when the content of the ci is the id of a possibleSpeciesFeatureValue and the speciesReference attribute must be defined. The interpretation of such a ci is the same as a ci element having a parameter which the possibleSpeciesFeatureValue links via its numericValue attribute.

The following example demonstrates the use of this attribute for "sum" of species concentrations.

```
k1*Si/(k2+SUM(Si))
```

In this example, the reactant "Si" is a pattern species which may have multiple *fully defined* species mapping to it, for example species "S1", "S2", ..., "Sn". "SUM(Si)" is a function to calculate the total concentration of all *fully* 

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*defined* **species** mapping to "Si". The **product** can be another pattern **species** "Pi". The SBML code for the math expression can be as follows:

```
<reaction id="r">
  <listOfReactants>
     <speciesReference species="Si" />
   </list0fReactants>
   tofProducts>
      <speciesReference species="Pi" />
   </list0fProducts>
   <kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
         <apply>
            <divide>
               <apply>
                  <times />
                  <ci>Si</ci>
                  <ci>k1</ci>
               </apply>
               <apply>
                  <plu> <plu> />
                  <ci>k2</ci>
                  <ci multi:representationType="sum">Si</ci>
               </apply>
            </divide>
         </apply>
      <listOfLocalParameters>
         <localParameter id="k1" ... />
         <localParameter id="k2" ... />
      </listOfLocalParameters>
   </kineticLaw>
</reaction>
```

The math expressions for the individual species in the example can be:

```
For species S1: k1*S1/(k2 + (S1 + S2 + ... + Sn))

For species S2: k1*S2/(k2 + (S1 + S2 + ... + Sn))

...

For species Sn: k1*Sn/(k2 + (S1 + S2 + ... + Sn))
```

### 3.27 Namespace scoping rules for identifiers

In the Multi package, as in SBML Level 3 Version 1 Core, the **Model** object contains the main components of an SBML model, such as **species**, **compartments** and **reactions**. The package defines new classes within a model and the scope of the identifiers of those new classes should be defined to prevent identifier collisions. In this section, we describe the scoping rules for all of the types and classes defined in Section 3.4 to Section 3.26 on pages 11–40.

- 1. The namespace for **SId** identifiers defined within a **Model** object used in the Multi package follows the same rules as those defined in SBML Level 3 Version 1 Core for plain **Model** objects. The scope of the identifiers is limited to the enclosing **Model** object.
- 2. The identifier of every **SpeciesType** and **PossibleSpeciesFeatureValue** object defined in the Multi package must be unique across the set of all identifiers in the **Model** object in which it is located.
- 3. The identifier of every SpeciesTypeInstance, SpeciesTypeComponentIndex, InSpeciesTypeBond and SpeciesFeatureType object defined in the Multi package must be unique across the set of all identifiers of the same class under the direct parent SpeciesType object in which it is located.
- 4. The identifier of every **SpeciesFeature** and **SubListOfSpeciesFeatures** object defined in the Multi package must be unique across the set of all identifiers in the **Species** object in which it is located.
- 5. The identifier, if defined, of every **CompartmentReference** object defined in the Multi package must be unique across the set of all identifiers in the **Compartment** object in which it is located.

# 4 Examples

This section contains examples employing the Multi package for SBML Level 3.

### 4.1 Example: Compartment, Species Type, and Species

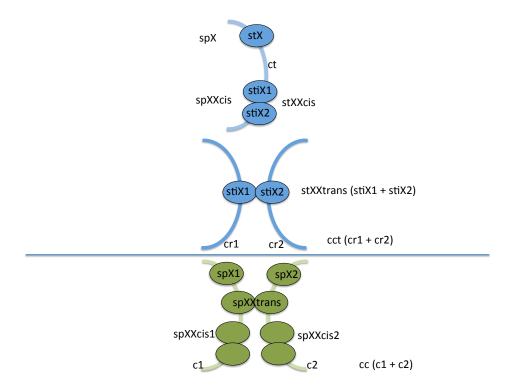


Figure 17: Diagram for an example of Compartment, Species Type and Species

Figure 17 shows an example illustrating the usages of and relations among the **Compartment**, **SpeciesType** and **Species** classes.

"ct" is a compartment type. "cct" is a composite compartment type with two compartmentReferences "cr1" and "cr2" both referencing "ct". "c1" is a "not-a-type" compartment and references "ct". Similarly, "c2" is also a "not-a-type" compartment and references "ct". "cc" is a composite "not-a-type" compartment composed of "c1" and "c2".

"stX" is a speciesType on the "ct" compartment. "stXXcis" is a speciesType on the "ct" compartment, and has two speciesTypeInstances "stiX1" and "stX2" both of that reference the "stX" speciesType. "stXXtrans" is a speciesType on the "cct" compartment with two speciesTypeInstances "stiX1" and "stiX2" sitting in different sub-compartments.

"spX" is a species referencing speciesType "stX". "spXXcis" is a species referencing "stXXcis". "spX1" is a species referencing "stX" and sitting in the "c1" compartment. "spX2" is a species also referencing "stX", but sitting in "c2". "spXXtrans" is a species referencing "stXXtrans". "spXXcis1" is a species referencing "stXXtrans" and sitting in "c1". "spXXcis1" is a species referencing "stXXtrans" and sitting in "c2".

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"spX1", "spX2", "spXXtrans", "spXXcis1" and "spXXcis2" are fully defined species (see Section 3.19 on page 33).

The SBML code can be as follows:

```
<listOfCompartments>
   <compartment id="ct" multi:isType="true" />
   <compartment id="cct" multi:isType="true">
      <multi:listOfCompartmentReferences>
         <multi:compartmentReference multi:id="cr1" multi:compartment="ct" />
         <multi:compartmentReference multi:id="cr2" multi:compartment="ct" />
      </multi:listOfCompartmentReferences>
   </compartment>
   <compartment id="c1" multi:isType="false" multi:compartmentType="ct" />
   <compartment id="c2" multi:isType="false" multi:compartmentType="ct" />
   <compartment id="cc" multi:isType="false" multi:compartmentType="cct">
      <multi:listOfCompartmentReferences>
         <multi:compartmentReference multi:compartment="c1" />
         <multi:compartmentReference multi:compartment="c2" />
      </multi:listOfCompartmentReferences>
   </compartment>
</listOfCompartments>
<multi:listOfSpeciesTypes>
   <multi:bindingSiteSpeciesType multi:id="stX" multi:compartment="ct" />
   <multi:speciesType multi:id="stXXcis" multi:compartment="ct">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="stiX1" multi:speciesType="stX" />
         <multi:speciesTypeInstance multi:id="stiX2" multi:speciesType="stX" />
      </multi:listOfSpeciesTypeInstances>
      <multi:listOfInSpeciesTypeBonds>
         <multi:inSpeciesTypeBond multi:bindingSite1="stiX1" multi:bindingSite2="stiX2" />
      </multi:listOfInSpeciesTypeBonds>
   </multi:speciesType>
   <multi:speciesType multi:id="stXXtrans" multi:compartment="cct">
      <multi:listOfSpeciesTypeInstances>
         <multi:speciesTypeInstance multi:id="stiX1" multi:speciesType="stX"
            multi:compartmentReference="cr1" />
         <multi:speciesTypeInstance multi:id="stiX2" multi:speciesType="stX"</pre>
            multi:compartmentReference="cr2" />
      </multi:listOfSpeciesTypeInstances>
      <multi:listOfInSpeciesTypeBonds>
         <multi:inSpeciesTypeBond multi:bindingSite1="stiX1" multi:bindingSite2="stiX2" />
      </multi:listOfInSpeciesTypeBonds>
   </multi:speciesType>
</multi:listOfSpeciesTypes>
st0fSpecies>
   <species id="spX" multi:speciesType="stX" compartment="ct" />
   <species id="spXXcis" multi:speciesType="stXXcis" compartment="ct" />
   <species id="spX1" multi:speciesType="stX" compartment="c1" /> <!-- Fully defined -->
   <species id="spX2" multi:speciesType="stX" compartment="c2" /> <!-- Fully defined -->
  <species id="spXXtrans" multi:speciesType="stXXtrans" compartment="cc" /> <!-- Fully defined -->
  <species id="spXXcis1" multi:speciesType="stXXcis" compartment="c1" /> <!-- Fully defined -->
<species id="spXXcis2" multi:speciesType="stXXcis" compartment="c2" /> <!-- Fully defined -->
</listOfSpecies>
```

# 4.2 *Simmune* example: the Ecad model

The Simmune toolset (http://go.usa.gov/QeH) has some example models including the published Ecad model [Angermann et al. (2012)]. The Ecad model describes the interactions between E-cadherin receptors that can associate either with other E-cadherin receptors within the same membrane (in "cis") or with E-cadherin receptors on adjacent membranes (in "trans"). This model is transformed into the SBML Level 3 format with use of the Multi package.

```
<?xml version="1.0" encoding="UTF-8"?>
```

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```
<sbml xmlns="http://www.sbml.org/sbml/level3/version1/core" level="3" version="1"</pre>
   xmlns:multi="http://www.sbml.org/sbml/level3/version1/multi/version1" multi:required="true">
   <model name="E-cadherin_mediated_adhesion">
      <!-- Definitions -->
      <listOfUnitDefinitions>
         <unitDefinition id="litre_per_mole_per_sec">
            tofUnits>
               <unit kind="litre" exponent="1" scale="0" multiplier="1" />
               <unit kind="mole" exponent="-1" scale="0" multiplier="1" />
               <unit kind="second" exponent="-1" scale="0" multiplier="1" />
            </listOfIInits>
         </unitDefinition>
         <unitDefinition id="micron_square_per_sec">
            listOfUnits>
               <unit kind="metre" exponent="2" scale="-6" multiplier="1" />
               <unit kind="second" exponent="-1" scale="0" multiplier="1" />
            </listOfUnits>
         </unitDefinition>
         <unitDefinition id="micrometre_per_sec">
            listOfUnits>
               <unit kind="metre" exponent="1" scale="-6" multiplier="1" />
               <unit kind="second" exponent="-1" scale="0" multiplier="1" />
            </listOfUnits>
         </unitDefinition>
         <unitDefinition id="per_sec">
            <listOfUnits>
               <unit kind="second" exponent="-1" scale="0" multiplier="1" />
            </listOfUnits>
         </unitDefinition>
      </listOfUnitDefinitions>
      <!-- Compartments -->
      <listOfCompartments>
         <compartment id="membrane" constant="true" multi:isType="true" />
         <compartment id="inter_membrane" constant="true" multi:isType="true">
            <multi:listOfCompartmentReferences>
               <multi:compartmentReference multi:id="m1" multi:compartment="membrane" />
               <multi:compartmentReference multi:id="m2" multi:compartment="membrane" />
            </multi:listOfCompartmentReferences>
         </compartment>
      </list0fCompartments>
      <!-- SpeciesTypes -->
      <multi:listOfSpeciesTypes>
         <!-- Ecad with cis-binding site and trans-binding site: -->
         <multi:bindingSiteSpeciesType multi:id="st_Cis_Interface" />
         <multi:bindingSiteSpeciesType multi:id="st_Trans_Interface" />
         <multi:speciesType multi:id="st_Ecad" multi:compartment="membrane">
            <multi:listOfSpeciesTypeInstances>
               <multi:speciesTypeInstance multi:id="cis" multi:speciesType="st_Cis_Interface" />
               <multi:speciesTypeInstance multi:id="trans" multi:speciesType="st_Trans_Interface" />
            </multi:listOfSpeciesTypeInstances>
         </multi:speciesType>
         <!-- cis dimer: -->
         <multi:speciesType multi:id="st_Ecad_cis_dimer" multi:compartment="membrane">
            <multi:listOfSpeciesTypeInstances>
               <multi:speciesTypeInstance multi:id="Ecad1" multi:speciesType="st_Ecad" />
               <multi:speciesTypeInstance multi:id="Ecad_2" multi:speciesType="st_Ecad" />
            </multi:listOfSpeciesTypeInstances>
            <multi:listOfSpeciesTypeComponentIndexes>
               <multi:speciesTypeComponentIndex multi:id="Ecad1cis"
                  multi:component="cis" multi:identifyingParent="Ecad1" />
               <multi:speciesTypeComponentIndex multi:id="Ecad2cis"</pre>
```

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```
multi:component="cis" multi:identifyingParent="Ecad2" />
      <multi:speciesTypeComponentIndex multi:id="Ecad1trans"
         multi:component="trans" multi:identifyingParent="Ecad1" />
      <multi:speciesTypeComponentIndex multi:id="Ecad2trans"</pre>
         multi:component="trans" multi:identifyingParent="Ecad2" />
   </multi:listOfSpeciesTypeComponentIndexes>
   <multi:listOfInSpeciesTypeBonds>
      <multi:inSpeciesTypeBond multi:bindingSite1="Ecad1cis"
         multi:bindingSite2="Ecad2cis" />
   </multi:listOfInSpeciesTypeBonds>
</multi:speciesType>
<!-- trans dimer: -->
<multi:speciesType multi:id="st_Ecad_trans_dimer" multi:compartment="inter_membrane">
   <multi:listOfSpeciesTypeInstances>
      <multi:speciesTypeInstance multi:id="Ecad1" multi:speciesType="st_Ecad"</pre>
         multi:compartmentReference="m1" />
      <multi:speciesTypeInstance multi:id="Ecad2" multi:speciesType="st_Ecad"</pre>
         multi:compartmentReference="m2" />
   </multi:listOfSpeciesTypeInstances>
   <multi:listOfSpeciesTypeComponentIndexes>
      <multi:speciesTypeComponentIndex multi:id="Ecad1trans"</pre>
         multi:component="trans" multi:identifyingParent="Ecad1" />
      <multi:speciesTypeComponentIndex multi:id="Ecad2trans"
         multi:component="trans" multi:identifyingParent="Ecad2" />
      <multi:speciesTypeComponentIndex multi:id="Ecad1cis"</pre>
         multi:component="cis" multi:identifyingParent="Ecad1" />
      <multi:speciesTypeComponentIndex multi:id="Ecad2cis"
         multi:component="cis" multi:identifyingParent="Ecad2" />
   </multi:listOfSpeciesTypeComponentIndexes>
   <multi:listOfInSpeciesTypeBonds>
      <multi:inSpeciesTypeBond multi:bindingSite1="Ecad1trans"</pre>
         multi:bindingSite2="Ecad2trans" />
   </multi:listOfInSpeciesTypeBonds>
</multi:speciesType>
<!-- trimer: -->
<multi:speciesType multi:id="st_Ecad_trimer" multi:compartment="inter_membrane">
   <multi:listOfSpeciesTypeInstances>
      <multi:speciesTypeInstance multi:id="Ecad1" multi:speciesType="st_Ecad"</pre>
         multi:compartmentReference="m1" />
      <multi:speciesTypeInstance multi:id="Ecad2" multi:speciesType="st_Ecad"</pre>
         multi:compartmentReference="m1" />
      <multi:speciesTypeInstance multi:id="Ecad3" multi:speciesType="st_Ecad"</pre>
         multi:compartmentReference="m2" />
   </multi:listOfSpeciesTypeInstances>
   <multi:listOfSpeciesTypeComponentIndexes>
      <multi:speciesTypeComponentIndex multi:id="Ecad1cis"
         multi:component="cis" multi:identifyingParent="Ecad1" />
      <multi:speciesTypeComponentIndex multi:id="Ecad1trans"</pre>
         multi:component="trans" multi:identifyingParent="Ecad1" />
      <multi:speciesTypeComponentIndex multi:id="Ecad2cis"
         multi:component="cis" multi:identifyingParent="Ecad2" />
      <multi:speciesTypeComponentIndex multi:id="Ecad2trans"
         multi:component="trans" multi:identifyingParent="Ecad2" />
      <multi:speciesTypeComponentIndex multi:id="Ecad3cis"
         multi:component="cis" multi:identifyingParent="Ecad3" />
      <multi:speciesTypeComponentIndex multi:id="Ecad3trans"
         multi:component="trans" multi:identifyingParent="Ecad3" />
   </multi:listOfSpeciesTypeComponentIndexes>
   <multi:listOfInSpeciesTypeBonds>
      <multi:inSpeciesTypeBond multi:bindingSite1="Ecad1cis"</pre>
         multi:bindingSite2="Ecad2cis" />
      <multi:inSpeciesTypeBond multi:bindingSite1="Ecad1trans"</pre>
         multi:bindingSite2="Ecad3trans" />
   </multi:listOfInSpeciesTypeBonds>
```

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```
</multi:speciesType>
</multi:listOfSpeciesTypes>
<!-- Species -->
Species>
   <!-- free Ecad -->
   <species id="sp_Ecad_unbound" name="Ecad_unbound" compartment="membrane"</pre>
      hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false"
      multi:speciesType="st_Ecad">
      <multi:listOfOutwardBindingSites>
         <multi:outwardBindingSite multi:component="cis"
            multi:bindingStatus="unbound" />
         <multi:outwardBindingSite multi:component="trans"</pre>
            multi:bindingStatus="unbound" />
      </multi:listOfOutwardBindingSites>
   </species>
   <!-- Pattern species: Ecad trans unbnd -->
   <species id="sp_Ecad_trans_unbnd" name="Ecad_trans_unbnd" compartment="membrane"</pre>
      hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false"
      multi:speciesType="st_Ecad">
      <multi:listOfOutwardBindingSites>
         <multi:outwardBindingSite multi:component="trans"
            multi:bindingStatus="unbound" />
      </multi:listOfOutwardBindingSites>
   </species>
   <!-- Pattern species: Ecad trans bnd -->
   <species id="sp_Ecad_trans_bnd" name="Ecad_trans_bnd" compartment="membrane"
hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false"</pre>
      multi:speciesType="st_Ecad">
      <multi:listOfOutwardBindingSites>
         <multi:outwardBindingSite multi:component="trans"</pre>
            multi:bindingStatus="bound" />
      </multi:listOfOutwardBindingSites>
   </species>
   <!-- Pattern species: Ecad all -->
   <species id="sp_Ecad_all" name="Ecad_all" compartment="membrane"</pre>
      hasOnlySubstanceUnits="false"
      boundaryCondition="false" constant="false" multi:speciesType="st_Ecad" />
   <!-- Pattern species: Ecad cis unbnd -->
   <species id="sp_Ecad_cis_unbnd" name="Ecad_cis_unbnd" compartment="membrane"</pre>
      hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false"
      multi:speciesType="st_Ecad">
      <multi:listOfOutwardBindingSites>
         <multi:outwardBindingSite multi:component="cis"</pre>
            multi:bindingStatus="unbound" />
      </multi:listOfOutwardBindingSites>
   </species>
   <!-- Pattern species: Ecad cis unbnd, trans bnd -->
   <species id="sp_Ecad_6" name="Ecad_6" compartment="membrane" hasOnlySubstanceUnits="false"</pre>
      boundaryCondition="false" constant="false" multi:speciesType="st_Ecad">
      <multi:listOfOutwardBindingSites>
         <multi:outwardBindingSite multi:component="cis"
            multi:bindingStatus="unbound" />
         <multi:outwardBindingSite multi:component="trans"</pre>
            multi:bindingStatus="bound" />
      </multi:listOfOutwardBindingSites>
   </species>
   <!-- Pattern species: Ecad cis bnd, trans unbnd -->
   <species id="sp_Ecad_7" name="Ecad_7" compartment="membrane" hasOnlySubstanceUnits="false"</pre>
```

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```
boundaryCondition="false" constant="false" multi:speciesType="st_Ecad">
      <multi:listOfOutwardBindingSites>
         <multi:outwardBindingSite multi:component="cis"
            multi:bindingStatus="bound" />
         <multi:outwardBindingSite multi:component="trans"</pre>
            multi:bindingStatus="unbound" />
      </multi:listOfOutwardBindingSites>
   </species>
   <!-- Pattern species: Ecad cis dimer -->
   <species id="sp_Ecad_cis_dimer" name="Ecad_cis_dimer" compartment="membrane"</pre>
      hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false"
      multi:speciesType="st_Ecad_cis_dimer" />
   <!-- Pattern species: Ecad cis dimer: all trans bnd -->
   <species id="sp_EcadEcad_2" name="Ecad.Ecad_2" compartment="membrane"</pre>
      hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false"
      multi:speciesType="st_Ecad_cis_dimer">
      <multi:listOfOutwardBindingSites>
         <multi:outwardBindingSite multi:component="Ecad1trans"</pre>
            multi:bindingStatus="bound" />
         <multi:outwardBindingSite multi:component="Ecad2trans"</pre>
            multi:bindingStatus="bound" />
      </multi:listOfOutwardBindingSites>
   </species>
   <!-- Pattern species: Ecad trans dimer -->
   <species id="sp_EcadEcad_1" name="Ecad.Ecad_1" compartment="inter_membrane"</pre>
      hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false"
      multi:speciesType="st_Ecad_trans_dimer" />
   <!-- Pattern species: Ecad trans dimer: all cis bnd -->
   <species id="sp_Ecad_trans_dimer_2" name="Ecad_trans_dimer_2" compartment="inter_membrane"</pre>
      hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false"
      multi:speciesType="st_Ecad_trans_dimer">
      <multi:listOfOutwardBindingSites>
         <multi:outwardBindingSite multi:component="Ecad1cis"
            multi:bindingStatus="bound" />
         <multi:outwardBindingSite multi:component="Ecad2cis"</pre>
            multi:bindingStatus="bound" />
      </multi:listOfOutwardBindingSites>
   </species>
   <!-- Pattern species: Ecad True Trimer -->
   <species id="sp_Ecad_True_Trimer" compartment="inter_membrane" hasOnlySubstanceUnits="false"</pre>
      boundaryCondition="false" constant="false" multi:speciesType="st_Ecad_trimer">
      <multi:listOfOutwardBindingSites>
         <multi:outwardBindingSite multi:component="Ecad2trans"
            multi:bindingStatus="unbound" />
         <multi:outwardBindingSite multi:component="Ecad3cis"</pre>
            multi:bindingStatus="unbound" />
      </multi:listOfOutwardBindingSites>
   </species>
   <!-- Pattern species: Ecad All Trimer -->
   <species id="sp_Ecad_All_Trimer" compartment="inter_membrane" hasOnlySubstanceUnits="false"</pre>
      boundaryCondition="false" constant="false" multi:speciesType="st_Ecad_trimer" />
</listOfSpecies>
<!-- Reactions -->
1istOfReactions>
   <!-- cis association: -->
   <reaction id="rc_Cis_Association" name="Cis_Association" reversible="false" fast="false"</pre>
      compartment="membrane">
```

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```
<speciesReference id="Cis_Association_r1" species="sp_Ecad_6"</pre>
         stoichiometry="1" constant="false" />
      <speciesReference id="Cis_Association_r2" species="sp_Ecad_6"</pre>
         stoichiometry="1" constant="false" />
   </list0fReactants>
   <listOfProducts>
      <speciesReference species="sp_EcadEcad_2" constant="false" />
   </listOfProducts>
   <kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
         <apply>
            <times />
            <ci> kon </ci>
            <ci multi:speciesReference="Cis_Association_r1"> sp_Ecad_6 </ci>
            <ci multi:speciesReference="Cis_Association_r2"> sp_Ecad_6 </ci>
         </apply>
      <listOfLocalParameters>
         <localParameter id="kon" value="9000" units="litre_per_mole_per_sec" />
      </listOfLocalParameters>
   </kineticLaw>
</reaction>
<!-- In species cis association: Here the model requires that the two interacting molecules
 are part of one connected complex already prior to the association. Since the necessary
 connectivity can only be mediated by the trans binding sites here, these sites must be
 bound to the subcomplex (not shown) linking the two interacting molecules.
<multi:intraSpeciesReaction id="rc_Intra_Complex_Cis_Association"</pre>
name="Intra-Complex_Cis_Association"
  reversible="false" fast="false" compartment="membrane">
   t0fReactants>
      <speciesReference id="Intra_Complex_Cis_Association_r1" species="sp_Ecad_6"</pre>
         stoichiometry="1" constant="false" />
      <speciesReference id="Intra_Complex_Cis_Association_r2" species="sp_Ecad_6"</pre>
         stoichiometry="1" constant="false" />
   </listOfReactants>
   tofProducts>
      <speciesReference species="sp_EcadEcad_2" constant="false" />
   </listOfProducts>
   <kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
         <apply>
            <times />
            <ci multi:speciesReference="Intra_Complex_Cis_Association_r1"> sp_Ecad_6 </ci>
         </apply>
      <listOfLocalParameters>
         <localParameter id="kon" value="100" units="per_sec" />
      </listOfLocalParameters>
   </kineticLaw>
</multi:intraSpeciesReaction>
<!-- trans association: -->
<reaction id="rc_Trans_Association" name="Trans_Association" reversible="false" fast="false"</pre>
   compartment="inter_membrane">
   <listOfReactants>
      <speciesReference id="Trans_Association_r1" species="sp_Ecad_trans_unbnd"</pre>
         multi:compartmentReference="m1" constant="false" />
      <speciesReference id="Trans_Association_r2" species="sp_Ecad_trans_unbnd"</pre>
         multi:compartmentReference="m2" constant="false" />
   </listOfReactants>
   tofProducts>
      <speciesReference species="sp_EcadEcad_1" constant="false" />
   </listOfProducts>
```

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```
<kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
         <apply>
            <times />
            <ci> kon </ci>
            <ci multi:speciesReference="Trans_Association_r1"> sp_Ecad_trans_unbnd </ci>
            <ci multi:speciesReference="Trans_Association_r2"> sp_Ecad_trans_unbnd </ci>
         </apply>
      <listOfLocalParameters>
         <localParameter id="kon" value="90000" units="litre_per_mole_per_sec" />
      </listOfLocalParameters>
   </kineticLaw>
</reaction>
<!-- In complex trans association: Here the model requires that the two interacting molecules
 are part of one connected complex already prior to the association. Since the necessary
 connectivity can only be mediated by the cis binding sites here, these sites must be bound
 to the subcomplex (not shown) linking the two interacting molecules.
<multi:intraSpeciesReaction id="rc_Intra_Complex_Trans_Association"</pre>
name="Intra-Complex_Trans_Association"
   reversible="false" fast="false" compartment="inter_membrane" >
   <listOfReactants>
      <speciesReference id="Intra_Complex_Trans_Association_r1" species="sp_Ecad_7"</pre>
         multi:compartmentReference="m1" constant="false" />
      <speciesReference id="Intra_Complex_Trans_Association_r2" species="sp_Ecad_7"</pre>
         multi:compartmentReference="m2" constant="false" />
   </listOfReactants>
   t0fProducts>
      <speciesReference species="sp_Ecad_trans_dimer_2" constant="false" />
   </listOfProducts>
   <kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
         <apply>
            <ci> kon </ci>
            <ci multi:speciesReference="Intra_Complex_Trans_Association_r1"> sp_Ecad_7 </ci>
         </apply>
      <listOfLocalParameters>
         <localParameter id="kon" value="100" units="per_sec" />
      </listOfLocalParameters>
   </kineticLaw>
</multi:intraSpeciesReaction>
<!-- cis dissociation: -->
<reaction id="rc_Cis_dissociation" name="Cis_dissociation" reversible="false" fast="false"</pre>
   compartment="membrane">
   <listOfReactants>
      <speciesReference species="sp_Ecad_cis_dimer" constant="false" />
   </listOfReactants>
   <listOfProducts>
      <speciesReference id="Cis_dissociation_p1" species="sp_Ecad_cis_unbnd"</pre>
         stoichiometry="1" constant="false" />
      <speciesReference id="Cis_dissociation_p2" species="sp_Ecad_cis_unbnd"</pre>
         stoichiometry="1" constant="false" />
   </listOfProducts>
   <kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
         <apply>
            <times />
            <ci> koff </ci>
            <ci> sp_Ecad_cis_unbnd </ci>
         </apply>
```

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```
<listOfLocalParameters>
         <localParameter id="koff" value="1" units="per_sec" />
      </listOfLocalParameters>
   </kineticLaw>
</reaction>
<!-- In-species cis dissociation: By specifying that this reaction breaks only an inner bond,
 the model limits the application of this reaction to dissociations that result in only one
 reaction product. The complex is still connected through a subcomplex that is not shown
 here but that links the two molecules involved in the reaction at their trans binding
 sites. Note that the modeler application has to ensure the correct application of this
 rule and its consistent definition. For instance, specifying the one or both of the trans
 binding sites to be unbound would lead to a rule that could never be applied because the
 trans bindings are required for the connectivity of the result complex.
 -->
<multi:intraSpeciesReaction id="rc_Intra_Complex_Cis_dissociation"</pre>
name="Intra-Complex_Cis_dissociation"
   reversible="false" fast="false" compartment="membrane" >
   <listOfReactants>
      <speciesReference species="sp_EcadEcad_2" constant="false" />
   </list0fReactants>
   <listOfProducts>
      <speciesReference id="Intra_Complex_Cis_dissociation_p1" species="sp_Ecad_6"</pre>
         stoichiometry="2" constant="false" />
      <speciesReference id="Intra_Complex_Cis_dissociation_p2" species="sp_Ecad_6"</pre>
         stoichiometry="2" constant="false" />
   </list0fProducts>
   <kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
         <apply>
            <times />
            <ci> koff </ci>
            <ci> sp_Ecad_6 </ci>
         </apply>
      <listOfLocalParameters>
         <localParameter id="koff" value="0.01" units="per_sec" />
      </listOfLocalParameters>
   </kineticLaw>
</multi:intraSpeciesReaction>
<!-- trans dissociation: -->
<reaction id="rc_Trans_dissociation" name="Trans_dissociation" reversible="false"</pre>
   fast="false" compartment="inter_membrane">
   listOfReactants>
      <speciesReference species="sp_EcadEcad_1" constant="false" />
   </listOfReactants>
   <speciesReference id="Trans_dissociation_p1" species="sp_Ecad_trans_unbnd"</pre>
         multi:compartmentReference="m1" constant="false" />
      <speciesReference id="Trans_dissociation_p2" species="sp_Ecad_trans_unbnd"</pre>
         multi:compartmentReference="m2" constant="false" />
   </listOfProducts>
   <kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
         <apply>
            <times />
            <ci> koff </ci>
            <ci> sp_Ecad_trans_unbnd </ci>
         </apply>
      <listOfLocalParameters>
         <localParameter id="koff" value="1" units="per_sec" />
      </listOfLocalParameters>
   </kineticLaw>
</reaction>
```

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```
<!-- In species trans dissociation: By specifying that this reaction breaks only an inner
           bond, the model limits the application of this reaction to dissociations that result in
           only one reaction product. The complex is still connected through a subcomplex that is
           not shown here but that links the two molecules involved in the reaction at their cis
           binding sites. Note that the modeler application has to ensure the correct application
           of this rule and its consistent definition. For instance, specifying the one or both of
           the cis binding sites to be unbound would lead to a rule that could never be applied
           because the cis bindings are required for the connectivity of the result complex.
         <multi:intraSpeciesReaction id="rc_Intra_Complex_Trans_dissociation"</pre>
          name="Intra-Complex_Trans_dissociation"
            reversible="false" fast="false" compartment="inter_membrane" >
            <listOfReactants>
                <speciesReference species="sp_Ecad_trans_dimer_2" constant="false" />
            </listOfReactants>
            tofProducts>
                <speciesReference id="Intra_Complex_Trans_dissociation_p1" species="sp_Ecad_7"</pre>
                   multi:compartmentReference="m1" constant="false" />
               <speciesReference id="Intra_Complex_Trans_dissociation_p2" species="sp_Ecad_7"</pre>
                   multi:compartmentReference="m2" constant="false" />
            </list0fProducts>
            <kineticLaw>
               <math xmlns="http://www.w3.org/1998/Math/MathML">
                   <apply>
                      <times />
                      <ci> koff </ci>
                      <ci> sp_Ecad_7 </ci>
                   </apply>
               <listOfLocalParameters>
                   <localParameter id="koff" value="0.01" units="per_sec" />
                </listOfLocalParameters>
            </kineticLaw>
         </multi:intraSpeciesReaction>
      </listOfReactions>
   </model>
</sbml>
```

## 4.3 A *BioNetGen* example from its user manual

egfr\_simple.bngl (http://bionetgen.org/index.php/BNGManual:Listing\_1)

```
begin parameters
 NA 6.02e23
                           # Avogadro's number (molecules/mol)
 f 1
                           # Fraction of the cell to simulate
 Vo f*1.0e-10
                           # Extracellular volume=1/cell_density (L)
 V f*3.0e-12
                           # Cytoplasmic volume (L)
 EGF_init 20*1e-9*NA*Vo
                           # Initial amount of ligand (20 nM)
                           # converted to copies per cell
 # Initial amounts of cellular components (copies per cell)
 EGFR_init f*1.8e5
 Grb2_init f*1.5e5
 Sos1_init f*6.2e4
 # Rate constants
 # Divide by NA*V to convert bimolecular rate constants
 # from /M/sec to /(molecule/cell)/sec
 kp1 9.0e7/(NA*Vo) # ligand-monomer binding
 km1 0.06
                     # ligand-monomer dissociation
 kp2 1.0e7/(NA*V)
                     # aggregation of bound monomers
 km2 0.1
                    # dissociation of bound monomers
```

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```
# dimer transphosphorylation
 kp3 0.5
                     # dimer dephosphorylation
  km3 4.505
 kp4 1.5e6/(NA*V) # binding of Grb2 to receptor
                     # dissociation of Grb2 from receptor
 km4 0.05
 kp5 1.0e7/(NA*V) # binding of Grb2 to Sos1
                      # dissociation of Grb2 from Sos1
 km5 0.06
 deg 0.01
                     # degradation of receptor dimers
end parameters
begin molecule types
 EGF(R)
 EGFR(L,CR1,Y1068~U~P)
 Grb2(SH2,SH3)
                                                                                                              13
 Sos1(PxxP)
 Trash()
end molecule types
begin seed species
                                                                                                              18
 EGF(R)
                                                                                                              19
 EGFR(L,CR1,Y1068~U) EGFR_init
 Grb2(SH2,SH3)
                     Grb2 init
                                                                                                              21
 Sos1(PxxP)
                      Sos1_init
                                                                                                              22
end seed species
                                                                                                              24
begin observables
 1 Molecules EGFR_tot EGFR()
 2 Molecules Lig_free EGF(R)
                                                                                                              27
 3 Species Dim EGFR(CR1!+)
 4 Molecules RP
                         EGFR(Y1068~P!?)
 5 Molecules Grb2Sos1 Grb2(SH2,SH3!1).Sos1(PxxP!1)
6 Molecules Sos1_act EGFR(Y1068!1).Grb2(SH2!1,SH3!2).Sos1(PxxP!2)
                                                                                                              30
end observables
                                                                                                              32
                                                                                                              33
begin reaction rules
 # Ligand-receptor binding
                                                                                                              35
 1 EGFR(L,CR1) + EGF(R) \leftarrow EGFR(L!1,CR1).EGF(R!1) kp1, km1
                                                                                                              36
 # Receptor-aggregation
                                                                                                              38
 2 EGFR(L!+,CR1) + EGFR(L!+,CR1) <-> EGFR(L!+,CR1!1).EGFR(L!+,CR1!1) kp2,km2
                                                                                                              39
 # Transphosphorylation of EGFR by RTK
                                                                                                              41
 3 EGFR(CR1!+,Y1068~U) -> EGFR(CR1!+,Y1068~P) kp3
                                                                                                              42
 # Dephosphorylation
                                                                                                              44
 4 EGFR(Y1068~P) -> EGFR(Y1068~U) km3
 # Grb2 binding to pY1068
                                                                                                              47
 5 EGFR(Y1068~P) + Grb2(SH2) <-> EGFR(Y1068~P!1).Grb2(SH2!1) kp4,km4
 # Grb2 binding to Sos1
                                                                                                              50
 6 Grb2(SH3) + Sos1(PxxP) <-> Grb2(SH3!1).Sos1(PxxP!1) kp5,km5
                                                                                                              51
                                                                                                              52
 # Receptor dimer internalization/degradation
                                                                                                              53
  7 EGF(R!1).EGF(R!2).EGFR(L!1,CR1!3).EGFR(L!2,CR1!3) -> Trash()
end reaction rules
                                                                                                              55
                                                                                                              56
#actions
generate_network({overwrite=>1});
                                                                                                              58
                                                                                                              59
# Equilibration
simulate_ode({suffix=>equil,t_end=>100000,n_steps=>10,sparse=>1,steady_state=>1});
setConcentration("EGF(R)","EGF_init");
                                                                                                              62
saveConcentrations(); # Saves concentrations for future reset
                                                                                                              64
# Kinetics
                                                                                                              65
writeSBML({});
```

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```
simulate_ode({t_end=>120,n_steps=>120});
resetConcentrations(); # reverts to saved Concentrations
simulate_ssa({suffix=>ssa,t_end=>120,n_steps=>120});
```

The SBML code can be as follows. Please note, the SBML code does not cover the content other than the model in the bngl file, such as the "actions", "Equilibration" and "Kinetics" sections.

```
<?xml version="1.0" encoding="UTF-8"?>
<sbml xmlns="http://www.sbml.org/sbml/level3/version1/core" level="3" version="1"</pre>
   xmlns:multi="http://www.sbml.org/sbml/level3/version1/multi/version1" multi:required="true">
   <model name="bionetgen_example_egfr_simple">
      <listOfUnitDefinitions>
         <unitDefinition id="molecules_per_mol">
            listOfUnits>
               <unit kind="mole" scale="0" mulitplier="1" exponent="-1" />
            </listOfUnits>
         </unitDefinition>
      </listOfUnitDefinitions>
      <!-- compartments -->
      <listOfCompartments>
         <compartment id="Vo" constant="true" spatialDimensions="3" units="liter"</pre>
            multi:isType="false" />
         <compartment id="V" constant="true" spatialDimensions="3" units="liter"</pre>
            multi:isType="false" />
      </listOfCompartments>
      <!-- speciesType -->
      <multi:listOfSpeciesTypes>
         <!-- EGF(R) -->
         <multi:bindingSiteSpeciesType multi:id="st_EGF_bs_R" />
         <multi:speciesType multi:id="st_EGF">
            <multi:listOfSpeciesTypeInstances>
               <multi:component multi:id="R" multi:speciesType="st_EGF_bs_R" />
            </multi:listOfSpeciesTypeInstances>
         </multi:speciesType>
         <!-- EGFR(L, CR1, Y1068~U~P) -->
         <multi:bindingSiteSpeciesType multi:id="st_EGFR_bs_L" />
         <multi:bindingSiteSpeciesType multi:id="st_EGFR_bs_CR1" />
         <multi:bindingSiteSpeciesType multi:id="st_EGFR_bs_Y1068">
            <multi:listOfSpeciesFeatureTypes>
               <multi:speciesFeatureType multi:id="sft_Y1068">
                  <multi:listOfPossibleSpeciesFeatureValues>
                     <multi:possibleSpeciesFeatureValue multi:id="U" />
                     <multi:possibleSpeciesFeatureValue multi:id="P" />
                  </multi:listOfPossibleSpeciesFeatureValues>
               </multi:speciesFeatureType>
            </multi:listOfSpeciesFeatureTypes>
         </multi:bindingSiteSpeciesType>
         <multi:speciesType multi:id="st_EGFR">
            <multi:listOfSpeciesTypeInstances>
               <multi:component multi:id="L" multi:speciesType="st_EGFR_bs_L" />
               <multi:component multi:id="CR1" multi:speciesType="st_EGFR_bs_CR1" />
               <multi:component multi:id="Y1068" multi:speciesType="st_EGFR_bs_Y1068" />
            </multi:listOfSpeciesTypeInstances>
         </multi:speciesType>
         <!-- EGFR dimer: [EGFR(CR1!1).EGFR(CR1!1)] -->
         <multi:speciesType multi:id="st_EGFR_dimer">
            <multi:listOfSpeciesTypeInstances>
               <multi:component multi:id="EGFR1" multi:speciesType="st_EGFR" />
```

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```
<multi:component multi:id="EGFR2" multi:speciesType="st_EGFR" />
   </multi:listOfSpeciesTypeInstances>
   <multi:listOfSpeciesTypeComponentIndexes>
      <multi:speciesTypeComponentIndex multi:id="EGFR1CR1"</pre>
         multi:component="CR1" identifyingParent="EGFR1" />
      <multi:speciesTypeComponentIndex multi:id="EGFR2CR1"</pre>
         multi:component="CR1" identifyingParent="EGFR2" />
   </multi:listOfSpeciesTypeComponentIndexes>
   <multi:listOfInSpeciesTypeBonds>
      <multi:inSpeciesTypeBond multi:bindingSite1="EGFR1CR1"</pre>
         multi:bindingSite2="EGFR2CR1" />
   </multi:listOfInSpeciesTypeBonds>
</multi:speciesType>
<!-- EGFR-EGF dimer: [EGF(R!1).EGF(R!2).EGFR(L!1,CR1!3).EGFR(L!2,CR1!3)] -->
<multi:speciesType multi:id="st_EGFR_EGF_dimer">
   <multi:listOfSpeciesTypeInstances>
      <multi:component multi:id="EGF1" multi:speciesType="st_EGF" />
      <multi:component multi:id="EG2" multi:speciesType="st_EGF" />
      <multi:component multi:id="EGFR1" multi:speciesType="st_EGFR" />
      <multi:component multi:id="EGFR2" multi:speciesType="st_EGFR" />
   </multi:listOfSpeciesTypeInstances>
   <multi:listOfSpeciesTypeComponentIndexes>
      <multi:speciesTypeComponentIndex multi:id="EGF1R"</pre>
         multi:component="R" identifyingParent="EGF1" />
      <multi:speciesTypeComponentIndex multi:id="EGF2R"</pre>
         multi:component="R" identifyingParent="EGF2" />
      <multi:speciesTypeComponentIndex multi:id="EGFR1L"</pre>
         multi:component="L" identifyingParent="EGFR1" />
      <multi:speciesTypeComponentIndex multi:id="EGFR2L"
         multi:component="L" identifyingParent="EGFR2" />
      <multi:speciesTypeComponentIndex multi:id="EGFR1CR1"</pre>
         multi:component="CR1" identifyingParent="EGFR1" />
      <multi:speciesTypeComponentIndex multi:id="EGFR2CR1"</pre>
         multi:component="CR1" identifyingParent="EGFR2" />
   </multi:listOfSpeciesTypeComponentIndexes>
   <multi:listOfInSpeciesTypeBonds>
      <multi:inSpeciesTypeBond multi:bindingSite1="EGFR1CR1" multi:bindingSite2="EGFR2CR1" />
      <multi:inSpeciesTypeBond multi:bindingSite1="EGF1R" multi:bindingSite2="EGFR1L" />
      <multi:inSpeciesTypeBond multi:bindingSite1="EGF2R" multi:bindingSite2="EGFR2L" />
   </multi:listOfInSpeciesTypeBonds>
</multi:speciesType>
<!-- Grb2(SH2, SH3) -->
<multi:bindingSiteSpeciesType multi:id="st_Grb2_bs_SH2" />
<multi:bindingSiteSpeciesType multi:id="st_Grb2_bs_SH3" />
<multi:speciesType multi:id="st_Grb2">
   <multi:listOfSpeciesTypeInstances>
      <multi:component multi:id="SH2" multi:speciesType="st_Grb2_bs_SH2" />
      <multi:component multi:id="SH3" multi:speciesType="st_Grb2_bs_SH3" />
   </multi:listOfSpeciesTypeInstances>
</multi:speciesType>
<multi:bindingSiteSpeciesType multi:id="st_Sos1_bs_PxxP" />
<multi:speciesType multi:id="st_Sos1">
   <multi:listOfSpeciesTypeInstances>
      <multi:component multi:id="PxxP" multi:speciesType="st_Sos1_bs_PxxP" />
   </multi:listOfSpeciesTypeInstances>
</multi:speciesType>
<!-- Trash -->
<multi:speciesType multi:id="trash" />
<!-- Grb2-Sos1 -->
<multi:speciesType multi:id="st_Grb2_Sos1">
```

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```
<multi:listOfSpeciesTypeInstances>
          <multi:component multi:id="Grb2" multi:speciesType="st_Grb2" />
<multi:component multi:id="Sos1" multi:speciesType="st_Sos1" />
      </multi:listOfSpeciesTypeInstances>
      <multi:listOfInSpeciesTypeBonds>
          <multi:inSpeciesTypeBond multi:bindingSite1="SH3" multi:bindingSite2="PxxP" />
      </multi:listOfInSpeciesTypeBonds>
   </multi:speciesType>
   <!-- EGFR(Y1068!1).Grb1(SH2!1,SH3!2).Sos1(PxxP!2) -->
   <multi:speciesType multi:id="st_EGFR_Grb2_Sos1">
      <multi:listOfSpeciesTypeInstances>
         <multi:component multi:id="EGFR" multi:speciesType="st_EGFR" />
<multi:component multi:id="Grb2" multi:speciesType="st_Grb2" />
<multi:component multi:id="Sos1" multi:speciesType="st_Sos1" />
                                                                                                               13
      </multi:listOfSpeciesTypeInstances>
      <multi:listOfInSpeciesTypeBonds>
          <multi:inSpeciesTypeBond multi:bindingSite1="Y1068" multi:bindingSite2="SH2" />
                                                                                                               18
          <multi:inSpeciesTypeBond multi:bindingSite1="SH3" multi:bindingSite2="PxxP" />
                                                                                                               19
      </multi:listOfInSpeciesTypeBonds>
                                                                                                               20
   </multi:speciesType>
                                                                                                               21
   <!-- EGFR(L!1).EGF(R!1) -->
   <multi:speciesType multi:id="st_EGFR_EGF">
                                                                                                               24
      <multi:listOfSpeciesTypeInstances>
          <multi:component multi:id="EGFR" multi:speciesType="st_EGFR" />
          <multi:component multi:id="EGF" multi:speciesType="st_EGF" />
                                                                                                               27
      </multi:listOfSpeciesTypeInstances>
      <multi:listOfInSpeciesTypeBonds>
                                                                                                               29
         <multi:inSpeciesTypeBond multi:bindingSite1="L" multi:bindingSite2="R" />
                                                                                                               30
      </multi:listOfInSpeciesTypeBonds>
   </multi:speciesType>
                                                                                                               32
                                                                                                               33
   <!-- EGFR(Y1068!1).Grb2(SH2!1) -->
   <multi:speciesType multi:id="st_EGFR_Grb2">
                                                                                                               35
      <multi:listOfSpeciesTypeInstances>
                                                                                                               36
          <multi:component multi:id="EGFR" multi:speciesType="st_EGFR" />
                                                                                                               37
          <multi:component multi:id="Grb2" multi:speciesType="st_Grb2" />
                                                                                                               38
      </multi:listOfSpeciesTypeInstances>
                                                                                                               39
      <multi:listOfInSpeciesTypeBonds>
          <multi:inSpeciesTypeBond multi:bindingSite1="Y1068" multi:bindingSite2="SH2" />
                                                                                                               41
      </multi:listOfInSpeciesTypeBonds>
                                                                                                               42
   </multi:speciesType>
                                                                                                               43
                                                                                                               44
</multi:listOfSpeciesTypes>
<!-- species -->
                                                                                                               47
Species>
   <species id="sp_EGF_free" name="EGF(R)" multi:speciesType="st_EGF"</pre>
                                                                                                               50
      hasOnlySubstanceUnits="false"
      boundaryCondition="false" constant="false">
      <multi:listOfOutwardBindingSites>
                                                                                                               53
          <multi:outwardBindingSite multi:component="R" multi:bindingStatus="unbound" />
      </multi:listOfOutwardBindingSites>
                                                                                                               55
   </species>
                                                                                                               56
   <species id="sp_EGFR_free_U" name="EGFR(L,CR1,Y1068~U)" multi:speciesType="st_EGFR"</pre>
                                                                                                               58
      hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
                                                                                                               59
      <multi:listOfOutwardBindingSites>
          <multi:outwardBindingSite multi:component="L" multi:bindingStatus="unbound" />
                                                                                                               61
         <multi:outwardBindingSite multi:component="CR1" multi:bindingStatus="unbound" />
                                                                                                               62
          <multi:outwardBindingSite multi:component="Y1068" multi:bindingStatus="unbound" />
      </multi:listOfOutwardBindingSites>
                                                                                                               64
      <multi:listOfSpeciesFeatures>
                                                                                                               65
         <multi:speciesFeature multi:speciesFeatureType="sft_Y1068">
```

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```
<multi:listOfSpeciesFeatureValues>
            <multi:speciesFeatureValue multi:value="U" />
         </multi:listOfSpeciesFeatureValues>
      </multi:speciesFeature>
   </multi:listOfSpeciesFeatures>
</species>
<species id="sp_Grb2_free" name="Grb2(SH2,SH3)" multi:speciesType="st_Grb2"
hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
   <multi:listOfOutwardBindingSites>
      <multi:outwardBindingSite multi:component="SH2" multi:bindingStatus="unbound" />
      <multi:outwardBindingSite multi:component="SH3" multi:bindingStatus="unbound" />
   </multi:listOfOutwardBindingSites>
</species>
<species id="sp_Grb2_SH2" name="Grb2(SH2)" multi:speciesType="st_Grb2"</pre>
   hasOnlySubstanceUnits="false"
   boundaryCondition="false" constant="false">
   <multi:listOfOutwardBindingSites>
      <multi:outwardBindingSite multi:component="SH2" multi:bindingStatus="unbound" />
   </multi:listOfOutwardBindingSites>
</species>
<species id="sp_Grb2_SH3" name="Grb2(SH3)" multi:speciesType="st_Grb2"</pre>
   hasOnlySubstanceUnits="false"
   boundaryCondition="false" constant="false">
   <multi:listOfOutwardBindingSites>
      <multi:outwardBindingSite multi:component="SH3" multi:bindingStatus="unbound" />
   </multi:listOfOutwardBindingSites>
</species>
<species id="sp_Sos1_free" name="Sos1(PxxP)" multi:speciesType="st_Sos1"</pre>
   hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
   <multi:listOfOutwardBindingSites>
      <multi:outwardBindingSite multi:component="PxxP" multi:bindingStatus="unbound" />
   </multi:listOfOutwardBindingSites>
</species>
<species id="sp_EGF_tot" name="EGF()" multi:speciesType="st_EGF"</pre>
   hasOnlySubstanceUnits="false"
   boundaryCondition="false" constant="false" />
<species id="sp_EGFR_dimerized" name="EGFR(CR1!+)" multi:speciesType="st_EGFR"</pre>
   hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
   <multi:listOfOutwardBindingSites>
      <multi:outwardBindingSite multi:component="CR1" multi:bindingStatus="bound" />
   </multi:listOfOutwardBindingSites>
</species>
<species id="sp_EGFR_U" name="EGFR(Y1068~P!?)" multi:speciesType="st_EGFR"</pre>
  hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
   <multi:listOfSpeciesFeatures>
      <multi:speciesFeature multi:speciesFeatureType="sft_Y1068">
         <multi:listOfSpeciesFeatureValues>
            <multi:speciesFeatureValue multi:value="P" />
         </multi:listOfSpeciesFeatureValues>
      </multi:speciesFeature>
   </multi:listOfSpeciesFeatures>
</species>
<species id="sp_EGFR_L_CR1" name="EGFR(L,CR1)" multi:speciesType="st_EGFR"</pre>
   hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
   <multi:listOfOutwardBindingSites>
      <multi:outwardBindingSite multi:component="L" multi:bindingStatus="unbound" />
      <multi:outwardBindingSite multi:component="CR1" multi:bindingStatus="unbound" />
   </multi:listOfOutwardBindingSites>
</species>
<species id="sp_EGFR_EGF_CR1" name="EGFR(L!1,CR1).EGF(R!1)" multi:speciesType="st_EGFR_EGF"</pre>
  hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
   <multi:listOfOutwardBindingSites>
      <multi:outwardBindingSite multi:component="CR1" multi:bindingStatus="unbound" />
   </multi:listOfOutwardBindingSites>
</species>
<species id="sp_EGFR_bL_CR1" name="EGFR(L!+,CR1)" multi:speciesType="st_EGFR"</pre>
```

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```
hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
   <multi:listOfOutwardBindingSites>
      <multi:outwardBindingSite multi:component="L" multi:bindingStatus="bound" />
      <multi:outwardBindingSite multi:component="CR1" multi:bindingStatus="unbound" />
   </multi:listOfOutwardBindingSites>
</species>
<species id="sp_EGFR_dimer_bL" name="EGFR(L!+,CR1!1).EGFR(L!+,CR1!1)"</pre>
   multi:speciesType="st_EGFR_dimer'
   hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
   <multi:listOfOutwardBindingSites>
      <multi:outwardBindingSite multi:component="EGFR1L"
         multi:bindingStatus="bound" />
      <multi:outwardBindingSite multi:component="EGFR2L"</pre>
         multi:bindingStatus="bound" />
   </multi:listOfOutwardBindingSites>
</species>
<species id="sp_EGFR_EGF_dimer" name="EGF(R!1).EGF(R!2).EGFR(L!1,CR1!3).EGFR(L!2,CR1!3)"</pre>
   multi:speciesType="st_EGFR_EGF_dimer" hasOnlySubstanceUnits="false"
   boundaryCondition="false"
   constant="false" />
<species id="sp_EGFR_bCR1_Y1068_U" name="EGFR(CR1!+,Y1068~U)" multi:speciesType="st_EGFR"</pre>
   hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
   <multi:listOfOutwardBindingSites>
      <multi:outwardBindingSite multi:component="CR1" multi:bindingStatus="bound" />
      <multi:outwardBindingSite multi:component="Y1068" multi:bindingStatus="unbound" />
   </multi:listOfOutwardBindingSites>
   <multi:listOfSpeciesFeatures>
      <multi:speciesFeature multi:speciesFeatureType="sft_Y1068">
         <multi:listOfSpeciesFeatureValues>
            <multi:speciesFeatureValue multi:value="U" />
         </multi:listOfSpeciesFeatureValues>
      </multi:speciesFeature>
   </multi:listOfSpeciesFeatures>
<species id="sp_EGFR_bCR1_Y1068_P" name="EGFR(CR1!+,Y1068~P)" multi:speciesType="st_EGFR"</pre>
   hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
   <multi:listOfOutwardBindingSites>
      <multi:outwardBindingSite multi:component="CR1" multi:bindingStatus="bound" />
      <multi:outwardBindingSite multi:component="Y1068" multi:bindingStatus="unbound" />
   </multi:listOfOutwardBindingSites>
   <multi:listOfSpeciesFeatures>
      <multi:speciesFeature multi:speciesFeatureType="sft_Y1068">
         <multi:listOfSpeciesFeatureValues>
            <multi:speciesFeatureValue multi:value="P" />
         </multi:listOfSpeciesFeatureValues>
      </multi:speciesFeature>
   </multi:listOfSpeciesFeatures>
</species>
<species id="sp_EGFR_Y1068_P" name="EGFR(Y1068~P)" multi:speciesType="st_EGFR"</pre>
   hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
   <multi:listOfOutwardBindingSites>
      <multi:outwardBindingSite multi:component="Y1068" multi:bindingStatus="unbound" />
   </multi:listOfOutwardBindingSites>
   <multi:listOfSpeciesFeatures>
      <multi:speciesFeature multi:speciesFeatureType="sft_Y1068">
         <multi:listOfSpeciesFeatureValues>
            <multi:speciesFeatureValue multi:value="P" />
         </multi:listOfSpeciesFeatureValues>
      </multi:speciesFeature>
   </multi:listOfSpeciesFeatures>
</species>
<species id="sp EGFR Y1068 U" name="EGFR(Y1068~U)" multi:speciesType="st EGFR"</pre>
   hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
   <multi:listOfOutwardBindingSites>
      <multi:outwardBindingSite multi:component="Y1068" multi:bindingStatus="unbound" />
   </multi:listOfOutwardBindingSites>
```

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```
<multi:listOfSpeciesFeatures>
         <multi:speciesFeature multi:speciesFeatureType="sft_Y1068">
            <multi:listOfSpeciesFeatureValues>
               <multi:speciesFeatureValue multi:value="U" />
            </multi:listOfSpeciesFeatureValues>
         </multi:speciesFeature>
      </multi:listOfSpeciesFeatures>
   </species>
   <species id="sp_EGFR_Grb2_P" name="EGFR(Y1068~P!1).Grb2(SH2!1)"</pre>
      multi:speciesType="st_EGFR_Grb2"
      hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
      <multi:listOfSpeciesFeatures>
         <multi:speciesFeature multi:speciesFeatureType="sft_Y1068">
            <multi:listOfSpeciesFeatureValues>
               <multi:speciesFeatureValue multi:value="P" />
            </multi:listOfSpeciesFeatureValues>
         </multi:speciesFeature>
      </multi:listOfSpeciesFeatures>
   </species>
   <species id="sp_Grb2_Sos1" name="Grb2(SH3!1).Sos1(PxxP!1)" multi:speciesType="st_Grb2_Sos1"</pre>
     hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false" />
   <species id="sp_Trash" name="Trash()" multi:speciesType="st_Trash"</pre>
      hasOnlySubstanceUnits="false"
      boundaryCondition="false" constant="false" />
</listOfSpecies>
<!-- parameters -->
<listOfParameters>
   <parameter id="NA" value="6.02e23" constant="true" units="molecules_per_mol" />
   <parameter id="f" value="1" constant="true" />
   <parameter id="kp1" constant="true" />
   <parameter id="km1" value="0.06" constant="true" />
   <parameter id="kp2" constant="true" />
   <parameter id="km2" value="0.1" constant="true" />
  <parameter id="kp3" value="0.5" constant="true" />
<parameter id="km3" value="4.505" constant="true" />
<parameter id="kp4" constant="true" />
   <parameter id="km4" value="0.05" constant="true" />
   <parameter id="kp5" constant="true" />
  <parameter id="km5" value="0.06" constant="true" />
   <parameter id="deg" value="0.01" constant="true" />
</list0fParameters>
<!-- intiialAssignments -->
<listOfIntialAssignments>
   <initialAssignment symbol="Vo">
      <math xmls="http://www.w3.org/1998/Math/MathML">
         <apply>
            <times />
            <ci>f</ci>
            <cn> 1e-10 </cn>
         </apply>
      </initialAssignment>
   <initialAssignment symbol="V">
      <math xmls="http://www.w3.org/1998/Math/MathML">
         <applv>
            <times />
            <ci>f</ci>
            <cn> 3e-12 </cn>
         </apply>
      </initialAssignment>
   <initialAssignment symbol="kp1">
```

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<math xmls="http://www.w3.org/1998/Math/MathML">
      <apply>
        <divide />
        <cn>9.02e7</cn>
        <apply>
           <times />
           <ci>NA</ci>
           <ci>Vo</ci>
        </apply>
     </apply>
  </initialAssignment>
<initialAssignment symbol="kp2">
   <math xmls="http://www.w3.org/1998/Math/MathML">
     <apply>
        <divide />
        <cn>1.0e7</cn>
        <apply>
           <times />
           <ci>NA</ci>
           <ci>V</ci>
        </apply>
     </apply>
  </initialAssignment>
<initialAssignment symbol="kp4">
  <math xmls="http://www.w3.org/1998/Math/MathML">
     <apply>
        <divide />
        <cn>1.5e6</cn>
        <apply>
           <times />
           <ci>NA</ci>
           <ci>V</ci>
        </apply>
     </apply>
  </initialAssignment>
<initialAssignment symbol="kp5">
  <math xmls="http://www.w3.org/1998/Math/MathML">
     <apply>
        <divide />
        <cn>1.0e7</cn>
        <apply>
           <times />
           <ci>NA</ci>
           <ci>V</ci>
        </apply>
     </apply>
  </initialAssignment>
<initialAssignment symbol="sp_EGF_free">
  <math xmls="http://www.w3.org/1998/Math/MathML">
     <apply>
        <times />
        <cn>20</cn>
        <cn>1e-9</cn>
        <ci>NA</ci>
        <ci>Vo</ci>
     </apply>
  </initialAssignment>
<initialAssignment symbol="sp_EGFR_free_U">
  <math xmls="http://www.w3.org/1998/Math/MathML">
     <apply>
        <times />
```

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```
<ci>f</ci>
            <cn>1.8e5</cn>
         </apply>
      </initialAssignment>
   <initialAssignment symbol="sp_Grb2_free">
      <math xmls="http://www.w3.org/1998/Math/MathML">
         <apply>
           <times />
            <ci>f</ci>
            <cn>1.5e5</cn>
         </apply>
      </initialAssignment>
   <initialAssignment symbol="sp_Sos1_free">
      <math xmls="http://www.w3.org/1998/Math/MathML">
         <apply>
            <times />
            <ci>f</ci>
            <cn>6.2e4</cn>
         </apply>
      </initialAssignment>
</listOfIntialAssignments>
<!-- reactions -->
<listOfReactions>
  <!-- # Ligand-receptor binding -->
   <!-- 1 EGFR(L,CR1) + EGF(R) <-> EGFR(L!1,CR1).EGF(R!1) kp1, km1 -->
   <reaction id="rc_Ligand_receptor_binding" reversible="true" fast="false">
      <listOfReactants>
         <speciesReference species="sp_EGFR_L_CR1" constant="false" />
         <speciesReference species="sp_EGF_free" constant="false" />
      </list0fReactants>
      <speciesReference species="sp_EGFR_EGF_CR1" constant="false" />
      </list0fProducts>
      <kineticLaw>
         <math xmlns="http://www.w3.org/1998/Math/MathML">
            <apply>
               <minus />
               <apply>
                  <times />
                  <ci> kp1 </ci>
                  <ci> sp_EGFR_L_CR1 </ci>
                  <ci> sp_EGF_free </ci>
               </apply>
               <apply>
                  <times />
                  <ci> km1 </ci>
                  <ci> sp_EGFR_EGF_CR1 </ci>
               </apply>
            </apply>
         </kineticLaw>
   </reaction>
  <!-- # Receptor-aggregation -->
   <!-- 2 EGFR(L!+, CR1) + EGFR(L!+, CR1) <-> EGFR(L!+, CR1!1) .EGFR(L!+, CR1!1) kp2, km2 -->
   <reaction id="rc_Receptor_aggregation" reversible="true" fast="false">
      <listOfReactants>
         <speciesReference species="sp_EGFR_bL_CR1" constant="false" stoichiometry="2" />
      </list0fReactants>
      <listOfProducts>
         <speciesReference species="sp_EGFR_dimer_bL" constant="false" />
```

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</listOfProducts>
   <kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
         <apply>
            <minus />
            <apply>
              <times />
              <ci> kp2 </ci>
               <ci> sp_EGFR_bL_CR1 </ci>
               <ci> sp_EGFR_bL_CR1 </ci>
            </apply>
            <apply>
              <times />
              <ci> km3 </ci>
              <ci> sp_EGFR_dimer_bL </ci>
            </apply>
         </apply>
      </kineticLaw>
</reaction>
<!-- # Transphosphorylation of EGFR by RTK -->
<!-- 3 EGFR(CR1!+,Y1068~U) -> EGFR(CR1!+,Y1068~P) kp3 -->
<reaction id="rc_Transphosphorylation" reversible="false" fast="false">
   <listOfReactants>
      <speciesReference species="sp_EGFR_bCR1_Y1068_U" constant="false" />
   </listOfReactants>
   <speciesReference species="sp_EGFR_bCR1_Y1068_P" constant="false" />
   </listOfProducts>
   <kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
         <apply>
            <times />
            <ci>kp3</ci>
            <ci>sp_EGFR_bCR1_Y1068_U</ci>
         </apply>
      </kineticLaw>
</reaction>
<!-- # Dephosphorylation -->
<!-- 4 EGFR(Y1068~P) -> EGFR(Y1068~U) km3 -->
<reaction id="rc_Dephosphorylation" reversible="false" fast="false">
   <listOfReactants>
      <speciesReference species="sp_EGFR_Y1068_P" constant="false" />
   </listOfReactants>
   <speciesReference species="sp_EGFR_Y1068_U" constant="false" />
   </list0fProducts>
   <kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
         <apply>
            <times />
            <ci>km3</ci>
            <ci>sp_EGFR_Y1068_P</ci>
         </apply>
      </kineticLaw>
</reaction>
<!-- # Grb2 binding to pY1068 -->
<!-- 5 EGFR(Y1068~P) + Grb2(SH2) <-> EGFR(Y1068~P!1).Grb2(SH2!1) kp4,km4 -->
<reaction id="rc_Grb2_binding_to_pY1068" reversible="true" fast="false">
   t0fReactants>
     <speciesReference species="sp_EGFR_Y1068_P" constant="false" />
```

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```
<speciesReference species="sp_Grb2_SH2" constant="false" />
   </list0fReactants>
   tofProducts>
      <speciesReference species="sp_EGFR_Grb2_P" constant="false" />
   </listOfProducts>
   <kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
         <apply>
            <minus />
            <apply>
               <times />
               <ci> kp4 </ci>
               <ci> sp_EGFR_Y1068_P </ci>
                <ci> sp_Grb2_SH2 </ci>
            </apply>
            <apply>
               <times />
               <ci> km4 </ci>
               <ci> sp_EGFR_Grb2_P </ci>
            </apply>
         </apply>
      </kineticLaw>
</reaction>
<!-- # Grb2 binding to Sos1 -->
<!-- 6 Grb2(SH3) + Sos1(PxxP) <-> Grb2(SH3!1).Sos1(PxxP!1) kp5,km5 -->
<reaction id="rc_Grb2_binding_to_Sos1" reversible="true" fast="false">
   <listOfReactants>
      <speciesReference species="sp_Grb2_SH3" constant="false" />
<speciesReference species="sp_Sos1_free" constant="false" />
   </list0fReactants>
   <listOfProducts>
      <speciesReference species="sp_Grb2_Sos1" constant="false" />
   </listOfProducts>
   <kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
         <apply>
            <minus />
            <apply>
               <times />
               <ci> kp5 </ci>
               <ci> sp_Grb2_SH3 </ci>
                <ci> sp_Sos1_free </ci>
            </apply>
            <apply>
               <times />
               <ci> km5 </ci>
                <ci> sp_Grb2_Sos1 </ci>
            </apply>
         </apply>
      </kineticLaw>
</reaction>
<!-- # Receptor dimer internalization/degradation -->
<!-- 7 EGF(R!1).EGF(R!2).EGFR(L!1,CR1!3).EGFR(L!2,CR1!3) -> Trash() -->
<reaction id="rc_EGFR_EGF_dimer_degration" reversible="false" fast="false">
   <listOfReactants>
      <speciesReference species="sp_EGFR_EGF_dimer" constant="false" />
   </list0fReactants>
   <listOfProducts>
      <speciesReference species="sp_Trash" constant="false" />
   </listOfProducts>
   <kineticLaw>
      <math xmlns="http://www.w3.org/1998/Math/MathML">
```

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### 4.4 Example from Kappa's documentation

Here is the example "An Introduction to Kappa Syntax" at *Kappa* website (http://www.kappalanguage.org/syntax.html).

Rule in English: "Unphosphorylated Site1 of A binds to Site1 of B"

Kappa Rule: A(Site1 u),B(Site1) -> A(Site1 u!1),B(Site1!1)

```
<?xml version="1.0" encoding="UTF-8"?>
<sbml xmlns="http://www.sbml.org/sbml/level3/version1/core" level="3" version="1"</pre>
   xmlns:multi="http://www.sbml.org/sbml/level3/version1/multi/version1" multi:required="true">
   <model name="An_Introduction_to_Kappa_Syntax">
      <!-- speciesType -->
      <multi:listOfSpeciesTypes>
         <!-- A:Site1 -->
         <multi:bindingSiteSpeciesType multi:id="st_A_Site1">
            <multi:listOfSpeciesFeatureTypes>
               <multi:speciesFeatureType multi:id="phosphorylation">
                  <multi:listOfPossibleSpeciesFeatureValues>
                     <multi:possibleSpeciesFeatureValue multi:id="U" />
                     <multi:possibleSpeciesFeatureValue multi:id="P" />
                  </multi:listOfPossibleSpeciesFeatureValues>
               </multi:speciesFeatureType>
            </multi:listOfSpeciesFeatureTypes>
         </multi:bindingSiteSpeciesType>
         <multi:speciesType multi:id="st_A">
            <multi:listOfSpeciesTypeInstances>
               <multi:speciesTypeInstance multi:id="Asite1" multi:speciesType="st_A_Site1" />
            </multi:listOfSpeciesTypeInstances>
         </multi:speciesType>
         <!-- B:Site1 -->
         <multi:bindingSiteSpeciesType multi:id="st_B_Site1" />
         <!-- B -->
         <multi:speciesType multi:id="st_B">
            <multi:listOfSpeciesTypeInstances>
               <multi:speciesTypeInstance multi:id="Bsite1" multi:speciesType="st_B_Site1" />
            </multi:listOfSpeciesTypeInstances>
         </multi:speciesType>
         <!-- A.B -->
         <multi:speciesType multi:id="st_AB">
            <multi:listOfSpeciesTypeInstances>
               <multi:speciesTypeInstance multi:id="A" multi:speciesType="st_A" />
               <multi:speciesTypeInstance multi:id="B" multi:speciesType="st_B" />
```

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```
</multi:listOfSpeciesTypeInstances>
      <multi:listOfInSpeciesTypeBonds>
         <multi:inSpeciesTypeBond multi:bindingSite1="Asite1"
            multi:bindingSite2="Bsite1" />
      </multi:listOfInSpeciesTypeBonds>
   </multi:speciesType>
</multi:listOfSpeciesTypes>
<!-- species -->
tofSpecies>
  <!-- species A with free unphosphorylated Site1 -->
   <species id="sp_A" name="A_with_Unphosphorylated_Site_1" multi:speciesType="st_A"</pre>
      hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false">
      <multi:listOfOutwardBindingSites>
         <multi:outwardBindingSite multi:component="Asite1"</pre>
            multi:bindingStatus="unbound" />
      </multi:listOfOutwardBindingSites>
      <multi:listOfSpeciesFeatures>
         <multi:speciesFeature multi:speciesFeatureType="phosphorylation">
            <multi:listOfSpeciesFeatureValues>
               <multi:speciesFeatureValue multi:value="U" />
            </multi:listOfSpeciesFeatureValues>
         </multi:speciesFeature>
      </multi:listOfSpeciesFeatures>
   </species>
   <!-- species B with free Site 1 -->
   <species id="sp_B" name="B" multi:speciesType="st_B" hasOnlySubstanceUnits="false"</pre>
      boundaryCondition="false" constant="false">
      <multi:listOfOutwardBindingSites>
         <multi:outwardBindingSite multi:component="sti_B_Site1"</pre>
            multi:bindingStatus="unbound" />
      </multi:listOfOutwardBindingSites>
   </species>
   <!-- species AB: unphosphorylated -->
   <species id="sp_AB" name="AB" multi:speciesType="st_AB" hasOnlySubstanceUnits="false"</pre>
      boundaryCondition="false" constant="false">
      <multi:listOfSpeciesFeatures>
         <multi:speciesFeature multi:speciesFeatureType="phosphorylation">
            <multi:listOfSpeciesFeatureValues>
               <multi:speciesFeatureValue multi:value="U" />
            </multi:listOfSpeciesFeatureValues>
         </multi:speciesFeature>
      </multi:listOfSpeciesFeatures>
   </species>
</listOfSpecies>
<!-- reactions -->
tofReactions>
   <!-- Unphosphorylated Site1 of A binds to Site1 of B -->
   <!-- Kappa Rule: A(Site1~u), B(Site1) -> A(Site1~u!1), B(Site1!1) -->
   <reaction id="rc_AB" reversible="false" fast="false">
      <speciesReference species="sp_A" constant="false" />
         <speciesReference species="sp_B" constant="false" />
      </listOfReactants>
      listOfProducts>
         <speciesReference species="sp_AB" constant="false" />
      </listOfProducts>
      <kineticLaw>
      </kineticLaw>
```

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# A Validation of SBML documents using Multi constructs

This section summarizes all the conditions that should be true of an SBML Level 3 Version 1 model that uses the Multi package. We use the same conventions that are used in the SBML Level 3 Version 1 Core specification document. In particular, there are different degrees of rule strictness. Formally, the differences are expressed in the statement of a rule: either a rule states that a condition *must* be true, or a rule states that it *should* be true. Rules of the former kind are strict SBML validation rules—a model encoded in SBML must conform to all of them in order to be considered valid. Rules of the latter kind are consistency rules. To help highlight these differences, we use the three symbols next to the rule numbers as described in section A of the SBML Level 3 Version 1 Core specification document:

- ✓ A checked box indicates a *requirement* for SBML conformance. If a model does not follow this rule, it does not conform to the Multi package specification. (Mnemonic intention behind the choice of symbol: "This must be checked.")
- A triangle indicates a *recommendation* for model consistency. If a model does not follow this rule, it is not considered strictly invalid as far as the Multi package specification is concerned; however, it indicates that the model contains a physical or conceptual inconsistency. (Mnemonic intention behind the choice of symbol: "This is a cause for warning.")
- ★ A star indicates a strong recommendation for good modeling practice. This rule is not strictly a matter of SBML encoding, but the recommendation comes from logical reasoning. As in the previous case, if a model does not follow this rule, it is not considered an invalid SBML encoding. (Mnemonic intention behind the choice of symbol: "You're a star if you heed this.")

The validation rules listed in the following subsections are all stated or implied in the rest of this specification document. They are enumerated here for convenience. Unless explicitly stated, all validation rules concern objects and attributes specifically defined in the Multi package.

For convenience and brevity, we use the shorthand "multi:x" to stand for an attribute or element name x in the namespace for the Multi package, using the namespace prefix multi. We use "multi:x" because it is shorter than to write a full explanation everywhere we refer to an attribute or element in the Multi package namespace.

## General rules about the Multi package

- multi-10101 ☑ To conform to Version 1 of the Multi package specification for SBML Level 3, an SBML document must declare the use of the following XML Namespace:
  - "http://www.sbml.org/sbml/level3/version1/multi/version1". (References: SBMLLevel 3 Package Specification for Multi Version 1, Section 3.1 on page 9.)
- multi-10102 
  ☑ Wherever they appear in an SBML document, elements and attributes from the Multi package must be declared either implicitly or explicitly to be in the XML namespace "http://www.sbml.org/sbml/level3/version1/multi/version1". (References: SBML Level 3 Package Specification for Multi Version 1, Section 3.1 on page 9.)

## General rules about MathML content in the Multi package

- multi-10201 A ci element in a Math object may have the optional attributes multi:speciesReference and multi:representationType. No other attributes from the Multi namespace are permitted on a ci element. (References: Section 3.26 on page 40).
- multi-10202 ☑ The value of the multi:speciesReference attribute on a given ci element must be the identifier of a SpeciesReference object within the same reaction. (References: Section 3.26.1 on page 40.)

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multi-10203 

✓ The value of the multi:representationType attribute on a given ci element must conform to the syntax of the Multi data type RepresentationType. (References: Section 3.26.2 on page 40.)

#### General rules about identifiers

- multi-10301 
  ✓ (Extends validation rule #10301 in the SBML Level 3 Version 1 Core specification.) Within a Model object, the values of the attributes id and multi:id on every instance of the following classes of objects must be unique across the set of all id and multi:id attribute values of all such objects in a model: the Model itself, plus all contained FunctionDefinition, Compartment, Species, Reaction, SpeciesReference, ModifierSpeciesReference, Event, and Parameter objects, plus the SpeciesType and PossibleSpeciesFeatureValue objects defined by the Multi package, and any objects defined by any other package with package:id attributes defined as falling in the 'SId' namespace. (References: Section 3.27 on page 42.)
- multi-10302 ☑ The value of a multi:id attribute must always conform to the syntax of the SBML data type SId. (References: SBML Level 3 Version 1 Core, Section 3.1.7.)
- multi-10303 ☑ The value of a multi:name attribute must always conform to the syntax of type string. (References: SBML Level 3 Version 1 Core, Section 3.1.1.)
- multi-10304 ☑ The value of a multi:id attribute on SpeciesTypeInstance objects must be unique across the set of all multi:id attribute values of all the SpeciesTypeInstance objects under the direct parent SpeciesType object in which it is located. (References: Section 3.11.1 on page 18 and Section 3.27 on page 42.)
- multi-10305 ☑ The value of a multi:id attribute on SpeciesTypeComponentIndex objects must be unique across the set of all multi:id attribute values of all the SpeciesTypeComponentIndex objects under the direct parent SpeciesType object in which it is located. (References: Section 3.12.1 on page 20 and Section 3.27 on page 42.)
- multi-10306 ☑ The value of a multi:id attribute on InSpeciesTypeBond objects must be unique across the set of all multi:id attribute values of all the InSpeciesTypeBond objects under the direct parent SpeciesType object in which it is located. (References: Section 3.13.1 on page 23 and Section 3.27 on page 42.)
- multi-10307 

  ✓ The value of a multi:id attribute on SpeciesFeatureType objects must be unique across the set of all multi:id attribute values of all the SpeciesFeatureType objects under the direct parent SpeciesType object in which it is located. (References: Section 3.9.1 on page 16 and Section 3.27 on page 42.)
- multi-10308 The value of a multi:id attribute on SubListOfSpeciesFeatures objects must be unique across the set of all id and multi:id attribute values of all objects in the Species object in which it is located. (References: Section 3.17.1 on page 30 and Section 3.27 on page 42.)
- multi-10309 

  ✓ The value of a multi:id attribute on SpeciesFeature objects must be unique across the set of all id and multi:id attribute values of all objects in the Species object in which it is located. (References: Section 3.18.1 on page 30 and Section 3.27 on page 42.)
- multi-10310 ☑ The value of a multi:id attribute on CompartmentReference objects must be unique across the set of all id and multi:id attribute values of all objects in the Compartment object in which it is located. (References: Section 3.6.1 on page 13 and Section 3.27 on page 42.)
- multi-10311 ☑ The value of a multi:compartment attribute on SpeciesType objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.8.2 on page 14.)

multi-10312 ☑	The value of a multi:numericValue attribute on PossibleSpeciesFeatureValue objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.10.2 on page 17.)	1
multi-10313 ☑	The value of a multi:speciesType attribute on SpeciesTypeInstance objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.11.2 on page 18.)	3
multi-10314 ☑	The value of a multi:compartmentReference attribute on SpeciesTypeInstance objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.11.3 on page 18.)	5
multi-10315 ☑	The value of a multi:component attribute on SpeciesTypeComponentIndex objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.12.2 on page 20.)	7
multi-10316 ☑	The value of a multi:identifyingParent attribute on SpeciesTypeComponentIndex objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.12.3 on page 20.)	9 10 11
multi-10317 ☑	The value of a multi:bindingSite1 attribute on InSpeciesTypeBond objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.13.2 on page 23.)	12
multi-10318 ☑	The value of a multi:bindingSite2 attribute on InSpeciesTypeBond objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.13.2 on page 23.)	14 15
multi-10319 ☑	The value of a multi:speciesType attribute on Species objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.15.1 on page 26.)	16 17
multi-10320 ☑	The value of a multi:component attribute on OutwardBindingSite objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.16.3 on page 28.)	18 19
multi-10321 ☑	The value of a multi:speciesFeatureType attribute on SpeciesFeature objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.18.2 on page 31.)	20 21
multi-10322 ☑	The value of a multi:component attribute on <b>SpeciesFeature</b> objects must conform to the syntax of the SBML data type <b>SIdRef</b> . (References: Section 3.18.4 on page 31.)	22 23
multi-10323 ☑	The value of a multi:value attribute on SpeciesFeatureValue objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.18.6 on page 31.)	24 25
multi-10324 ☑	The value of a multi:compartmentReference attribute on SimpleSpeciesReference objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.22 on page 35.)	26 27 28
multi-10325 ☑	The value of a multi:reactant attribute on SpeciesTypeComponentMapInProduct objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.24.2 on page 38.)	29 30 31
multi-10326 ☑	The value of a multi:reactantComponent attribute on SpeciesTypeComponentMapInProduct objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.24.3 on page 38.)	32 33 34
multi-10327 ☑	The value of a multi:productComponent attribute on SpeciesTypeComponentMapInProduct objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.24.4 on page 38.)	35 36 37
multi-10328 ☑	The value of a multi:compartmentType attribute on Compartment objects must conform to the syntax of the SBML data type SIdRef. (References: Section 3.5.2 on page 12.)	38
multi-10329 ☑	The value of a multi:compartment attribute on CompartmentReference objects must conform to the syntax of the SBML data type SIdRef (References: Section 3.6.2 on page 13.)	40

Rules for exte	nded SBML object	1
multi-20101 ☑	The multi:required attribute is required on the <sbml> element in the Multi package. (References: SBML Level 3 Package Specification for Multi Version 1, Section 3.1 on page 9.)</sbml>	2
multi-20102 ☑	The multi:required attribute on the <sbml> element must be Boolean. (References: SBML Level 3 Package Specification for Multi Version 1, Section 3.1 on page 9.)</sbml>	4
multi-20103 ☑	The value of the multi:required attribute on the <sbml> element must be "true". (References: SBML Level 3 Package Specification for Multi Version 1, Section 3.1 on page 9.)</sbml>	6 7
Rules for exte	nded Model objects	8
multi-20201    ✓	There may be at most one <b>ListOfSpeciesTypes</b> container object within a <b>Model</b> object. (References: Section 3.4 on page 11.)	9
multi-20202	A <b>ListOfSpeciesTypes</b> object within an extended <b>Model</b> object is optional, but if present, must not be empty. (References: Section 3.4 on page 11.)	11
multi-20203	A <b>ListOfSpeciesTypes</b> object may have the optional SBML core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace or the Multi namespace are permitted on a <b>ListOfSpeciesTypes</b> object. (References: Section 3.4.1 on page 11.)	13 14 15
multi-20204 ☑	Apart from the general <b>notes</b> and <b>annotation</b> subobjects permitted on all SBML objects, a <b>ListOfSpeciesTypes</b> container object may only contain <b>SpeciesType</b> objects. (References: Section 3.4.1 on page 11.)	16 17 18
Rules for exte	nded Compartment objects	19
multi-20301 ☑	An extended <b>Compartment</b> object must have the required attribute multi:isType, and may also have the optional attribute multi:comparetmentType. No other attributes from the Multi namespace are permitted on an extended <b>Compartment</b> object.(References: Section 3.5 on page 12.)	20 21 22 23
multi-20302 ☑	The value of a multi:isType attribute on an extended Compartment object must always confirm to the syntax of the SBML data type boolean. (References: Section 3.5.1 on page 12.)	24 25
multi-20303 ☑	The multi:isType attribute on an extended Compartment object is required. (References: Section 3.5.1 on page 12.)	26 27
multi-20304	The value of the multi:isType attribute of the Compartment object referenced by a Compartment-Reference object must be the same as that of the multi:isType attribute of the parent Compartment object of the ListOfCompartmentReferences object which contains the Compartment Reference object. (References: Section 3.7 on page 13.)	29
multi-20305 ☑	The multi:compartmentType attribute on a Compartment object must not be defined if the value of the multi:isType is "true". (References: Section 3.5.2 on page 12.)	32
multi-20306 ☑	There may be at most one <b>ListOfCompartmentReferences</b> container object within a <b>Compartment</b> object. (References: Section 3.5.3 on page 12.)	34 35
multi-20307 ☑	A <b>ListOfCompartmentReferences</b> object within a <b>Compartment</b> object is optional, but if present, must not be empty. (References: Section 3.5.3 on page 12.)	36 37
multi-20308 ☑	A <b>ListOfCompartmentReferences</b> object may have the optional SBML core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace or the Multi namespace are permitted on a <b>ListOfCompartmentReferences</b> object. (References: Section 3.5.3 on page 12.)	38 39 40 41

objects. (References: Section 3.5.3 on page 12.) Rules for SpeciesType objects A SpeciesType object may have the optional SBML Level 3 Core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace are permitted on a SpeciesType object. (References: Section 3.8 on page 14). A SpeciesType object may have the optional SBML Level 3 Core subobjects for notes and multi-20402 V annotation. No other elements from the SBML Level 3 Core namespace are permitted on a **SpeciesType** object. (References: Section 3.8 on page 14). multi-20403 V A Species Type object must have the required attribute multi:id, and may have the optional attributes multi:name and multi:compartment. No other attributes from the Multi namespace are permitted on a **SpeciesType** object. (References: Section 3.8 on page 14.) multi-20404 🗸 The value of the multi:compartment attribute, if set on a given SpeciesType object, must be the value of an id attribute on an existing Compartment object in the SId namespace of the parent **Model** object. (References: Section 3.8.2 on page 14.) multi-20405 V The various **ListOf** subobjects within a **SpeciesType** object are optional, but if present, these container objects must not be empty. Specifically, if any of the following classes of objects are present with a SpeciesType object, it must not be empty: ListOfSpeciesFeatureTypes, ListOf-SpeciesTypeInstances, ListOfSpeciesTypeComponentIndexes and ListOfInSpeciesTypeBonds. (References: Section 3.8 on page 14.) There may be at most one ListOfSpeciesFeatureTypes container object within a SpeciesType object. (References: Section 3.8 on page 14.) multi-20407 V Apart from the general notes and annotation subobjects permitted on all SBML objects, a ListOfSpeciesFeatureTypes container object may only contain SpeciesFeatureType objects. (References: Section 3.8.3 on page 15.) A ListOfSpeciesFeatureTypes object may have the optional SBML core attributes metaid and 27 sboTerm. No other attributes from the SBML Level 3 Core namespace or the Multi namespace are permitted on a ListOfSpeciesFeatureTypes object. (References: Section 3.8.3 on page 15.) multi-20409 V There may be at most one **ListOfSpeciesTypeInstances** container object within a **SpeciesType** object. (References: Section 3.8 on page 14.) 31 multi-20410 V Apart from the general notes and annotation subobjects permitted on all SBML objects, a ListOfSpeciesTypeInstances container object may only contain SpeciesTypeInstance objects. (References: Section 3.8.4 on page 15.) 34 multi-20411 🗸 A ListOfSpeciesTypeInstances object may have the optional SBML core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace or the Multi namespace are permitted on a **ListOfSpeciesTypeInstances**. (References: Section 3.8.4 on page 15.) There may be at most one ListOfSpeciesTypeComponentIndexes container object within a multi-20412 V **SpeciesType** object. (References: Section 3.8 on page 14.) multi-20413 🗹 Apart from the general notes and annotation subobjects permitted on all SBML objects, a ListOfSpeciesTypeComponentIndexes container object may only contain **SpeciesTypeComponentIndex** objects. (References: Section 3.8.6 on page 15.)

Apart from the general **notes** and **annotation** subobjects permitted on all SBML objects, a **ListOfCompartmentReferences** container object may only contain **CompartmentReference** 

multi-20414   ✓	A <b>ListOfSpeciesTypeComponentIndexes</b> object may have the optional SBML core attributes <b>metaid</b> and <b>sboTerm</b> . No other attributes from the SBML Level 3 Core namespace or the Multi namespace are permitted on a <b>ListOfSpeciesTypeComponentIndexes</b> object. (References: Section 3.8.6 on page 15.)	1 2 3 4
multi-20415   ✓	There may be at most one <b>ListOfInSpeciesTypeBonds</b> container object within a <b>SpeciesType</b> object. (References: Section 3.8 on page 14.)	5
multi-20416	Apart from the general notes and annotation subobjects permitted on all SBML objects, a <b>ListOfInSpeciesTypeBonds</b> container object may only contain <b>InSpeciesTypeBond</b> objects. (References: Section 3.8.5 on page 15.)	7 8 9
multi-20417   ✓	A <b>ListOfInSpeciesTypeBonds</b> object may have the optional SBML core attributes <b>metaid</b> and <b>sboTerm</b> . No other attributes from the SBML Level 3 Core namespace or the Multi namespace are permitted on a <b>ListOfInSpeciesTypeBonds</b> object. (References: Section 3.8.5 on page 15.)	10 11 12
Rules for Bind	dingSiteSpeciesType objects	13
multi-20501   ✓	A <b>BindingSiteSpeciesType</b> object is not permitted to have any <b>ListOfSpeciesTypeInstances</b> subobject. (References: Section 3.8.7 on page 15).	14 15
Rules for Spe	ciesFeatureType objects	16
multi-20601   ✓	A <b>SpeciesFeatureType</b> object may have the optional SBML Level 3 Core attributes <b>metaid</b> and <b>sboTerm</b> . No other attributes from the SBML Level 3 Core namespace are permitted on a <b>SpeciesFeatureType</b> object. (References: Section 3.9 on page 16).	17 18 19
multi-20602   ✓	A <b>SpeciesFeatureType</b> object may have the optional SBML Level 3 Core subobjects for <b>notes</b> and <b>annotation</b> . No other elements from the SBML Level 3 Core namespace are permitted on a <b>SpeciesFeatureType</b> object. (References: Section 3.9 on page 16).	20 21 22
multi-20603    ✓	A <b>SpeciesFeatureType</b> object must have the required attributes multi:id and multi:occur, and may have the optional attribute multi:name. No other attributes from the Multi namespace are permitted on a <b>SpeciesFeatureType</b> object. (References: Section 3.9 on page 16.)	23 24 25
multi-20604   ✓	The value of the multi:occur attribute on a given SpeciesFeatureType object must conform to the syntax of the SBML data type positiveInteger. (References: Section 3.9.2 on page 16.)	26 27
multi-20605   ✓	One <b>ListOfPossibleSpeciesFeatureValues</b> subobject in a <b>SpeciesFeatureType</b> object is required. (References: Section 3.9.3 on page 16.)	28 29
multi-20606 🗹	A <b>ListOfPossibleSpeciesFeatureValues</b> object may have the optional SBML core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace or the Multi namespace are permitted on a <b>ListOfPossibleSpeciesFeatureValues</b> object. (References: Section 3.9.3 on page 16.)	30 31 32 33
multi-20607    ✓	Apart from the general notes and annotation subobjects permitted on all SBML objects, a ListOfPossibleSpeciesFeatureValues container object may only contain PossibleSpeciesFeatureValue objects. (References: Section 3.9.3 on page 16.)	34 35 36
multi-20608	A <b>ListOfPossibleSpeciesFeatureValues</b> object must not be empty. (References: Section 3.9.3 on page 16.)	37 38

# Rules for PossibleSpeciesFeatureValue objects

multi-20701 ✓ A PossibleSpeciesFeatureValue object may have the optional SBML Level 3 Core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace are permit-

	ted on a <b>PossibleSpeciesFeatureValue</b> object. (References: Section 3.10 on page 17).
multi-20702 🗹	A <b>PossibleSpeciesFeatureValue</b> object may have the optional SBML Level 3 Core subobjects for <b>notes</b> and <b>annotation</b> . No other elements from the SBML Level 3 Core namespace are permitted on a <b>PossibleSpeciesFeatureValue</b> object. (References: Section 3.10 on page 17).
multi-20703	A <b>PossibleSpeciesFeatureValue</b> object must have the required attribute <b>multi:id</b> , and may have the optional attributes <b>multi:name</b> and <b>multi:numericValue</b> . No other attributes from the Multi namespace are permitted on a <b>PossibleSpeciesFeatureValue</b> object. (References: Section 3.10 on page 17.)
multi-20704   ✓	The value of the multi:numericValue attribute on a given PossibleSpeciesFeatureValue object must be the identifier of a Parameter object defined in the same Model object. (References: Section 3.10.2 on page 17.)
Rules for Spe	ciesTypelnstance objects
multi-20801   ✓	A <b>SpeciesTypeInstance</b> object may have the optional SBML Level 3 Core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace are permitted on a <b>SpeciesTypeInstance</b> object. (References: Section 3.11 on page 18).
multi-20802   ✓	A <b>SpeciesTypeInstance</b> object may have the optional SBML Level 3 Core subobjects for <b>notes</b> and <b>annotation</b> . No other elements from the SBML Level 3 Core namespace are permitted on a <b>SpeciesTypeInstance</b> object. (References: Section 3.11 on page 18).
multi-20803	A SpeciesTypeInstance object must have the required attributes multi:id and multi:species- Type, and may have the optional attributes multi:name and multi:compartmentReference. No other attributes from the Multi namespace are permitted on a SpeciesTypeInstance object. (References: Section 3.11 on page 18.)
multi-20805   ✓	The value of the multi:speciesType attribute on a given SpeciesTypeInstance object must be the identifier of a SpeciesType object defined in the same Model object. (References: Section 3.11.2 on page 18.)
multi-20806	The value of the multi:compartmentReference attribute, if present on a given SpeciesType-Instance object, must be the identifier of a CompartmentReference object defined in the same Model object. (References: Section 3.11.3 on page 18.)
Rules for Spe	ciesTypeComponentIndex objects
multi-20901 ☑	A <b>SpeciesTypeComponentIndex</b> object may have the optional SBML Level 3 Core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace are permitted on a <b>SpeciesTypeComponentIndex</b> object. (References: Section 3.12 on page 20).
multi-20902 🗹	A <b>SpeciesTypeComponentIndex</b> object may have the optional SBML Level 3 Core subobjects for <b>notes</b> and <b>annotation</b> . No other elements from the SBML Level 3 Core namespace are permitted on a <b>SpeciesTypeComponentIndex</b> object. (References: Section 3.12 on page 20).
multi-20903	A SpeciesTypeComponentIndex object must have the required attributes multi:id and multi:compartment, and may have the optional attributes multi:name and multi:identifying-Parent. No other attributes from the Multi namespace are permitted on a SpeciesType-ComponentIndex object. (References: Section 3.12 on page 20.)
multi-20904    ✓	The value of the multi:component attribute on a given SpeciesTypeComponentIndex object must be the identifier of a SpeciesTypeInstance object, or a SpeciesTypeComponentIndex object under the SpeciesType object that this SpeciesTypeComponentIndex object belongs to, or the SpeciesType object itself. (References: Section 3.12.2 on page 20.)

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The value of the multi:identifyingParent attribute on a given SpeciesTypeComponentIndex object must be the identifier of a component object under the SpeciesType object
that this SpeciesTypeComponentIndex object belongs to. A component object can be an
object of SpeciesTypeInstance, SpeciesTypeComponentIndex or SpeciesType. (References:
Section 3.12.3 on page 20.)

## Rules for InSpeciesTypeBond objects

- multi-21101 ✓ An InSpeciesTypeBond object may have the optional SBML Level 3 Core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace are permitted on an InSpeciesTypeBond object. (References: Section 3.13 on page 23).
- multi-21102 ✓ An InSpeciesTypeBond object may have the optional SBML Level 3 Core subobjects for notes and annotation. No other elements from the SBML Level 3 Core namespace are permitted on an InSpeciesTypeBond object. (References: Section 3.13 on page 23).
- multi-21103 ☑ An InSpeciesTypeBond object must have the required attributes, multi:bindingSite1 and multi:bindingSite2, and may have the optional attributes, multi:id and multi:name. No other attributes from the Multi namespace are permitted on an InSpeciesTypeBond object. (References: Section 3.13 on page 23.)
- multi-21104 ✓ The value of the multi:bindingSite1 attribute on a given InSpeciesTypeBond object must be the identifier of a SpeciesTypeInstance object or SpeciesTypeComponentIndex which ultimately reference a object of BindingSiteSpeciesType. (References: Section 3.13.2 on page 23.)
- multi-21105 

  ✓ The value of the multi:bindingSite2 attribute on a given InSpeciesTypeBond object must be the identifier of a SpeciesTypeInstance object or SpeciesTypeComponentIndex which ultimately reference a object of BindingSiteSpeciesType. (References: Section 3.13.2 on page 23.)
- multi-21106 

  ✓ The multi:bindingSite1 and multi:bindingSite2 attributes must not reference the same

  BindingSiteSpeciesType object. (References: Section 3.13.2 on page 23.)

### **Rules for extended Species objects**

- multi-21201 ✓ A Species object may have the optional attribute, multi:speciesType. No other attributes from the Multi namespace are permitted on a Species object. (References: Section 3.15 on page 26.)
- multi-21202 
  ✓ The value of a multi:speciesType attribute, if present on a Species object, must be the identifier of a SpeciesType object. (References: Section 3.15.1 on page 26.)
- multi-21203 ☑ Two ListOf \_\_ subobjects with a Species object are optional, but if present, these container object must not be empty. Specifically, if any of the following two classes of objects are present on the Species object, it must not be empty: ListOfOutwardBindingSites and ListOfSpecies-Features. (References: Section 3.15 on page 26.)
- multi-21204 ✓ A ListOfOutwardBindingSites object may have the optional SBML core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace or the Multi namespace are permitted on a ListOfOutwardBindingSites object. (References: Section 3.15.2 on page 26.)
- multi-21205 
  ✓ Apart from the general notes and annotation subobjects permitted on all SBML objects, a ListOfOutwardBindingSites container object may only contain OutwardBindingSite objects. (References: Section 3.15.2 on page 26.)
- multi-21206 ✓ A ListOfSpeciesFeatures object may have the optional SBML core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace or the Multi namespace are permitted on a ListOfSpeciesFeatures object. (References: Section 3.15.3 on page 27.)

multi-21207   ✓	A <b>SubListOfSpeciesFeatures</b> object may have the optional attributes multi:id, multi:name, multi:relation and multi:component. No other attributes from the Multi namespace are permitted on a <b>SubListOfSpeciesFeatures</b> object. (References: Section 3.17 on page 29.)	1 2 3
multi-21208   ✓	The value of the multi:relation attribute, if presented on a SubListOfSpeciesFeatures object, must conform to the syntax of the Multi data type Relation. (References: Section 3.17.2 on page 30.)	4 5
multi-21209   ✓	Apart from the general <b>notes</b> and <b>annotation</b> subobjects permitted on all SBML objects, a <b>ListOfSpeciesFeatures</b> container object may only contain <b>SpeciesFeature</b> and/or <b>SubListOfSpeciesFeatures</b> objects. (References: Section 3.15.3 on page 27.)	7 8 9
multi-21210   ✓	A <b>SubListOfSpeciesFeatures</b> object may have the optional SBML core attributes <b>metaid</b> and <b>sboTerm</b> . No other attributes from the SBML Level 3 Core namespace are permitted on a <b>SubListOfSpeciesFeatures</b> object. (References: Section 3.17 on page 29.)	1 1
multi-21211   ✓	Apart from the general <b>notes</b> and <b>annotation</b> subobjects permitted on all SBML objects, a <b>SubListOfSpeciesFeatures</b> container object may only contain <b>SpeciesFeature</b> objects. (References: Section 3.17 on page 29.)	1:
multi-21212 ☑	The value of the multi:component attribute on a given SubListOfSpeciesFeatures object must be the identifier of an object of SpeciesTypeInstance, SpeciesTypeComponentIndex or SpeciesType which contains the SpeciesFeature objects in this subListOfSpeciesFeatures. (References: Section 3.17.3 on page 30.)	1 1 1
Rules for Out	wardBindingSite objects	2
multi-21301   ✓	An <b>OutwardBindingSite</b> object may have the optional SBML Level 3 Core attributes <b>metaid</b> and <b>sboTerm</b> . No other attributes from the SBML Level 3 Core namespace are permitted on an <b>OutwardBindingSite</b> object. (References: Section 3.16 on page 28).	2 2
multi-21302   ✓	An <b>OutwardBindingSite</b> object may have the optional SBML Level 3 Core subobjects for <b>notes</b> and <b>annotation</b> . No other elements from the SBML Level 3 Core namespace are permitted on an <b>OutwardBindingSite</b> object. (References: Section 3.16 on page 28).	2 2
multi-21303   ✓	An OutwardBindingSite object must have the required attributes, multi:bindingStatus and multi:component, and may have the optional attributes multi:id and multi:name. No other attributes from the Multi namespace are permitted on an OutwardBindingSite object. (References: Section 3.16 on page 28.)	2 2 3
multi-21304   ✓	The value of the multi:bindingStatus attribute on a given OutwardBindingSite object must conform to the syntax of the Multi data type BindingStatus. (References: Section 3.16.2 on page 28.)	3
multi-21305   ✓	The value of the multi:component attribute on a given OutwardBindingSite object must be the identifier of an object of SpeciesTypeInstance, SpeciesTypeComponentIndex or SpeciesType which ultimately reference an object of BindingSiteSpeciesType. (References: Section 3.16.3 on page 28.)	3 3 3
multi-21306	An outwardBindingSite cannot be a binding site referenced by any inSpeciesTypeBond in the species. (References: Section 3.16.3 on page 28.)	3

# **Rules for SpeciesFeature objects**

multi-21401 ✓ A SpeciesFeature object may have the optional SBML Level 3 Core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace are permitted on a SpeciesFeature object. (References: Section 3.18 on page 30).

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A SpeciesFeature object may have the optional SBML Level 3 Core subobjects for notes and annotation. No other elements from the SBML Level 3 Core namespace are permitted on a **SpeciesFeature** object. (References: Section 3.18 on page 30). multi-21403 🗹 A SpeciesFeature object must have the required attributes, multi:speciesFeatureType and multi:occur, and may have the optional attributes, multi:id, multi:name, and multi:component. No other attributes from the Multi namespace are permitted on a SpeciesFeature object. (References: Section 3.18 on page 30.) The value of the multi:speciesFeatureType attribute on a given SpeciesFeature object multi-21404 🗹 must be the identifier of a SpeciesFeatureType object which is in the SpeciesType object referenced by the Species object containing this SpeciesFeature object. (References: Section 3.18.2 on page 31.) The value of the multi:occur attribute on a given SpeciesFeature object must conform to the syntax of the SBML data type positiveInteger. The value of the multi:occur attribute must not be larger than that of the multi:occur attribute of the SpeciesFeatureType object referenced by this **SpeciesFeature** object. (References: Section 3.18.3 on page 31.) The value of the multi:component attribute on a given SpeciesFeature object must be the identifier of an object of SpeciesTypeInstance, SpeciesTypeComponentIndex or SpeciesType which contains this **SpeciesFeature** object. (References: Section 3.18.4 on page 31.) One and only one ListOfSpeciesFeatureValues subobject within a SpeciesFeature object is 19 required. (References: Section 3.18.5 on page 31.) multi-21408 V A ListOfSpeciesFeatureValues object must not be empty. (References: Section 3.18.5 on page 31.) multi-21409 V A ListOfSpeciesFeatureValues object may have the optional SBML core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace or the Multi namespace are permitted on a ListOfSpeciesFeatureValues object. (References: Section 3.18.5 on page 31.) multi-21410 🗹 Apart from the general notes and annotation subobjects permitted on all SBML objects, a ListOfSpeciesFeatureValues container object may only contain SpeciesFeatureValue objects. 28 (References: Section 3.18.5 on page 31.) Rules for SpeciesFeatureValue objects multi-21501 🗹 A SpeciesFeatureValue object may have the optional SBML Level 3 Core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace are permitted on a 32 **SpeciesFeatureValue** object. (References: Section 3.18.6 on page 31). multi-21502 V A SpeciesFeatureValue object may have the optional SBML Level 3 Core subobjects for notes and annotation. No other elements from the SBML Level 3 Core namespace are permitted on a SpeciesFeatureValue object. (References: Section 3.18.6 on page 31). 36 multi-21503 V A SpeciesFeatureValue object must have the required attribute multi:value. No other attributes from the Multi namespace are permitted on a SpeciesFeatureValue object. (Refer-38 ences: Section 3.18.6 on page 31.) The value of the multi:value attribute on a given SpeciesFeatureValue object must be the identifier of a PossibleSpeciesFeatureValue object defined in the SpeciesFeatureType object referenced by the **SpeciesFeature** object containing this **SpeciesFeatureValue** object. (Refer-

ences: Section 3.18.6 on page 31.)

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## Rules for IntraSpeciesReaction objects

- multi-21601 ✓ An IntraSpeciesReaction object may have the optional SBML Level 3 Core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace and the Multi namespace are permitted on an IntraSpeciesReaction object. (References: Section 3.21 on page 34).
- multi-21602 ✓ An IntraSpeciesReaction object may have the optional SBML Level 3 Core subobjects for notes and annotation. No other elements from the SBML Level 3 Core namespace are permitted on an IntraSpeciesReaction object. (References: Section 3.21 on page 34).

#### Rules for extended SimpleSpeciesReference objects

- multi-21701 ✓ An extended SimpleSpeciesReference object may have the optional attribute, multi:compartmentReference. No other attributes from the Multi namespace are permitted on a SimpleSpeciesReference object. (References: Section 3.22 on page 35.)
- multi-21702 ✓ The value of a multi:compartmentReference attribute, if present on a SimpleSpeciesReference object, must be the identifier of a CompartmentReference object. (References: Section 3.22 on page 35.)

### Rules for extended SpeciesReference objects

- multi-21801 ✓ A ListOfSpeciesTypeComponentMapsInProduct object within an extended SpeciesReference object is optional, but if present, must not be empty. (References: Section 3.23.1 on page 37.)
- multi-21802 ✓ A ListOfSpeciesTypeComponentMapsInProduct object may have the optional SBML core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace or the Multi namespace are permitted on a ListOfSpeciesTypeComponentMapsInProduct object. (References: Section 3.23.1 on page 37.)
- multi-21803 ✓ Apart from the general notes and annotation subobjects permitted on all SBML objects, a ListOfSpeciesTypeComponentMapsInProduct container object may only contain Species-TypeComponentMapInProduct objects. (References: Section 3.23.1 on page 37.)

#### Rules for SpeciesTypeComponentMapInProduct objects

- multi-21901 ✓ A SpeciesTypeComponentMapInProduct object may have the optional SBML Level 3 Core attributes metaid and sboTerm. No other attributes from the SBML Level 3 Core namespace are permitted on a SpeciesTypeComponentMapInProduct object. (References: Section 3.24 on page 38).
- multi-21902 ✓ A SpeciesTypeComponentMapInProduct object may have the optional SBML Level 3 Core subobjects for notes and annotation. No other elements from the SBML Level 3 Core namespace are permitted on a SpeciesTypeComponentMapInProduct object. (References: Section 3.24 on page 38).
- Magnetical Magnetical
- multi-21904 ✓ The value of the multi:reactant attribute on a given SpeciesTypeComponentMapInProduct object must be the identifier of a reactant SpeciesReference object within a reaction. (References: Section 3.24.2 on page 38.)

multi-21905   ✓	The value of the multi:reactantComponent attribute on a given SpeciesTypeComponent-MapInProduct object must be the identifier of an object of SpeciesTypeInstance, SpeciesType-ComponentIndex or SpeciesType. (References: Section 2.24.2 on page 20.)	1 2
	ComponentIndex or SpeciesType. (References: Section 3.24.3 on page 38.)	3
multi-21906   ✓	The value of the multi:productComponent attribute on a given SpeciesTypeComponentMap-	4
	InProduct object must be the identifier of an object of SpeciesTypeInstance, SpeciesType-	5
	ComponentIndex or SpeciesType. (References: Section 3.24.4 on page 38.)	6
Rules for Con	mpartmentReference objects	7
multi-22001 ☑	A CompartmentReference object may have the optional SBML Level 3 Core attributes metaid	8
	and sboTerm. No other attributes from the SBML Level 3 Core namespace are permitted on a	9
	CompartmentReference object. (References: Section 3.6 on page 13).	10
multi-22002    ✓	A CompartmentReference object may have the optional SBML Level 3 Core subobjects for	11
	notes and annotation. No other elements from the SBML Level 3 Core namespace are	12
	permitted on a <b>CompartmentReference</b> object. (References: Section 3.6 on page 13).	13
multi-22003 ☑	A CompartmentReference object must have the required attribute multi:compartment, and	14
	may have the optional attributes multi:id and multi:name. No other attributes from the	15
	Multi namespace are permitted on a <b>CompartmentReference</b> object. (References: Section 3.6 on	16
	page 13.)	17
multi-22004	The value of the multi:compartment attribute must be the value of an id attribute on an exist-	18
	ing Compartment object in the SId namespace of the parent Model. (References: Section 3.6 on	19
	page 13.)	20
multi-22005	If some or all CompartmentReference objects within a ListOfCompartmentReferences object	21
	reference the same Compartment object, those compartmentReferences are required to have	22
	its multi:id attribute defined. (References: Section 3.6.1 on page 13.)	23
multi-22006	A compartmentReference cannot reference a compartment that directly or indirectly contains	24
	the compartmentReference. (References: Section 3.6.2 on page 13.)	25

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