Systems Biology Markup Language (SBML) Level 2: Structures and Facilities for Model Definitions

Andrew Finney, Michael Hucka
{afinney,mhucka}@cds.caltech.edu
Systems Biology Workbench Development Group
ERATO Kitano Symbiotic Systems Project
Control and Dynamical Systems, MC 107-81
California Institute of Technology, Pasadena, CA 91125, USA
http://www.cds.caltech.edu/erato

Principal Investigators: John Doyle and Hiroaki Kitano

SBML Level 2, Version 1 (Final Draft) June 18, 2003

Contents		5 Example Models Expressed in XML Using SBML	32
1 Introduction	2	5.1 A Simple Example Application of SBML	32
1.1 Scope and Limitations	2	5.2 Example Involving Units	33
1.2 Notational Conventions	2	5.3 Example Involving Assignment Rules	36
	2	5.4 Example Involving Algebraic Rules	37
2 Overview of SBML	3	5.5 Example Involving Combinations of	
3 Preliminary Definitions	4	boundaryCondition and constant Values on	
3.1 Type SBase and the SBML Type Inheritance		Species with RateRule structures	39
Hierarchy	4	5.6 Example of translation from a multi-	
3.2 Guidelines for the Use of the annotation Field		compartmental model containing reactions and	
in SBase	5	rate rules to ODEs	41
3.3 The id and name Fields on SBML Components	6	5.7 Example Involving Function Definitions	43
3.4 Type SId	7	5.8 Example Involving <i>delay</i> Functions	44
3.5 Component Identifiers and Namespaces in SBML	8	5.9 Example Involving Events	45
3.6 Mathematical Formulas in SBML Level 2	8	5.10 Example Involving Two-Dimensional Compart-	
4 SBML Components	11	ments	47
4.1 The SBML Container	11	6 Discussion	50
4.2 Models	11	6.1 Future Enhancements: SBML Level 3 and Beyond	51
4.3 Function Definitions	12	6.2 Relationships to Other Efforts	52
4.4 Unit Definitions	13	Acknowledgments	52
4.5 Compartments	16	Appendix	54
4.6 Species	18	A Summary of Notation	54
4.7 Parameters	20	B Differences between Level 1 Version 2 and Level	
4.8 Rules	21	2	54
4.9 Reactions	25	C XML Schema for SBML	56
4.10 Events	30	References	64

1 Introduction

We present the Systems Biology Markup Language (SBML) Level 2, a model representation formalism for systems biology. SBML is oriented towards describing systems of biochemical reactions of the sort common in research on a number of topics, including cell signaling pathways, metabolic pathways, biochemical reactions, gene regulation, and many others. SBML is defined in a neutral fashion with respect to programming languages and software encoding; however, it is primarily oriented towards allowing models to be encoded using XML, the eXtensible Markup Language (Bosak and Bray, 1999; Bray et al., 2000). This document contains many examples of SBML models written in XML, as well as an XML Schema (Biron and Malhotra, 2000; Fallside, 2000; Thompson et al., 2000) that defines SBML Level 2. A downloadable copy of the XML Schema and other related documents and software are also available from the SBML project web site, http://www.sbml.org/.

Major releases of SBML are termed *levels*. SBML Level 2 evolved out of SBML Level 1 (Hucka et al., 2001, 2003). All of the structures of Level 1 can be mapped in a straightforward fashion to Level 2. In addition, a large subset of the structures in Level 2 can be mapped to Level 1. However, the levels remain distinct; a valid SBML Level 1 document is not a valid SBML Level 2 document, and likewise, a valid SBML Level 2 document is not a valid SBML Level 1 document. Appendix B lists the differences between Level 1 and Level 2.

The Systems Biology Markup Language was originally created in collaboration with the authors of the following systems: BioSpice (Arkin, 2001), Cellerator (Shapiro et al., 2001, 2003), COPASI (Mendes, 2000), DBSolve (Goryanin, 2001; Goryanin et al., 1999), E-Cell (Tomita et al., 1999, 2001), Gepasi (Mendes, 1997, 2001), Jarnac (Sauro, 2000; Sauro and Fell, 1991), JDesigner (Sauro, 2001), JigCell (Vass et al., 2003), NetBuilder (Schilstra and Bolouri, 2002), ProMot/DIVA (Stelling et al., 2001), StochSim (Bray et al., 2001; Morton-Firth and Bray, 1998), and Virtual Cell (Schaff et al., 2000, 2001). SBML Level 2 was developed with the help of these packages' authors, as well as help and collaboration from the creators of CellML (Hedley et al., 2001) and many other individuals listed in the Acknowledgments (Section 6.2).

1.1 Scope and Limitations

SBML Level 2 is meant to support basic biochemical network models and the kinds of operations that are possible in existing analysis/simulation tools. Future software tools will undoubtedly require further evolution of SBML, and we expect that higher SBML levels will add structures and facilities on top of Level 2 after the simulation community has had time to gain experience with the current language definition. In Section 6.1, we discuss extensions that will likely be included in SBML Level 3.

The definition of the model description language presented here does not specify *how* programs should communicate or read/write SBML. We assume that for a simulation program to communicate a model encoded in SBML, the program will have to translate its internal data structures to and from SBML, use a suitable transmission medium and protocol, etc., but these issues are outside of the scope of this document.

1.2 Notational Conventions

We define SBML using a graphical notation based upon UML, the Unified Modeling Language (Eriksson and Penker, 1998; Oestereich, 1999). This UML-based definition in turn is used to define an XML Schema (Biron and Malhotra, 2000; Fallside, 2000; Thompson et al., 2000) for SBML. There are three main advantages to using UML as a basis for defining SBML data structures. First, compared to using other notations or a programming language, the UML visual representations are generally easier to grasp by readers who are not computer scientists. Second, the visual notation is implementation-neutral: the defined structures can be encoded in any concrete implementation language—not just XML, but C or Java as well. Third, UML is a de facto industry standard that is documented in many sources. Readers are therefore more likely to be familiar with it than other notations.

Our notation and our approach for mapping it to XML Schemas is explained in a separate document (Hucka, 2000). A summary of the essential points is presented in Appendix A, and examples throughout this docu-

ment illustrate the approach. All types in SBML follow XML schema datatype definitions and conventions. We also follow certain naming and typographical conventions throughout this document. Specifically, the names of data structure attributes or fields begin with a lowercase letter, and the names of data structures and types begin with an uppercase letter. Keywords (names of types, XML elements, etc.) are written in a typewriter-style font; for example, Compartment is a type name and compartment is a field name. Likewise, literal XML examples are also written in a typewriter-style font.

2 Overview of SBML

The following is an example of a simple, hypothetical network of biochemical reactions that can be represented in SBML:

$$\begin{array}{ccc} S_1 & \underbrace{k_1[S_1]/([S_1]+k_2)} & S_2 \\ \\ S_2 & & \underbrace{k_3[S_2]} & S_3+S_4 \end{array}$$

Broken down into its constituents, this model contains a number of components: reactant species, product species, reactions, rate laws, and parameters in the rate laws. To analyze or simulate this network, additional components must be made explicit, including compartments for the species, and units on the various quantities. The top level of an SBML model definition simply consists of lists of these components:

beginning of model definition
list of function definitions (optional)
list of unit definitions (optional)
list of compartments (optional)
list of species (optional)
list of parameters (optional)
list of rules (optional)
list of reactions (optional)
list of events (optional)
end of model definition

The meaning of each component is as follows:

Function definition: A named function that may be used throughout the rest of the model.

Unit definition: A name for a unit used in the expression of quantities in a model. Units may be supplied in a number of contexts in an SBML model, and it is convenient to have a facility for both setting default units and for allowing combinations of units to be given abbreviated names.

Compartment: A container of finite size for substances. In SBML Level 2, a compartment is primarily a topological structure with size but no geometric qualities.

Species: A substance or entity that takes part in a reaction. Some example species are ions such as Ca²⁺ and molecules such as glucose or ATP. The primary qualities associated with a chemical species in SBML Level 2 are its initial amount and the compartment in which it is located.

Parameter: A quantity that has a symbolic name. SBML Level 2 provides the ability to define parameters that are global to a model as well as parameters that are local to a single reaction.

Rule: In SBML, a mathematical expression that is used in combination with the differential equations constructed based on the set of reactions; it can be used to establish constraints between variables in a model, define how a variable can be calculated from other variables, or used to define the rate of change of a variable.

Reaction: A statement describing some transformation, transport or binding process that can change the amount of one or more species. For example, a reaction may describe how certain entities (reactants)

are transformed into certain other entities (products). Reactions have associated rate laws describing how quickly they take place.

Event: A statement describing an instantaneous, discontinuous transformation of a set of variables of any type (species concentration, compartment size or parameter value) when some triggering condition is satisfied.

A software package can read an SBML model description and translate it into its own internal format for model analysis. For example, a package might provide the ability to simulate the model by constructing differential equations representing the network and then perform numerical time integration on the equations to explore the model's dynamic behavior.

SBML allows models of arbitrary complexity to be represented. Each type of component in a model is described using a specific type of data structure that organizes the relevant information. The data structures determine how the resulting model is encoded in XML.

In the sections that follow, the various constructs in SBML and their uses are described in detail. Section 3 first introduces a few basic structures that are used throughout SBML Level 2, then Section 4 provides details on each of the main components. Section 5 provides a number of complete examples of models encoded in XML using SBML Level 2. Section 6 contains a list of anticipated enhancements that will be made in SBML Level 3 and a discussion of other efforts related to SBML. Appendix A summarizes the UML-based notation used in this document. Appendix B describes the differences between SBML Level 1 Version 2 and SBML Level 2 as described in this document and Appendix C provides the complete XML Schema for SBML Level 2.

3 Preliminary Definitions

This section covers certain concepts and constructs that are used repeatedly in the rest of SBML Level 2 and are useful to discuss before diving into the details of the data structures provided in SBML Level 2.

3.1 Type SBase and the SBML Type Inheritance Hierarchy

Every structure composing an SBML Level 2 model definition has a specific data type that is derived directly or indirectly from a single abstract type called SBase. This base type is designed to allow a modeler or a software package to attach arbitrary information to each major structure or list in an SBML model. The definition of SBase is presented in Figure 1.

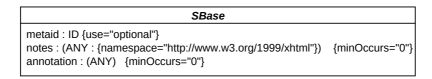


Figure 1: The definition of SBase. Text enclosed in braces next to attribute types (e.g., {minOccurs="0"}) indicates constraints on the possible attribute values. We use XML Schema language to express constraints since we are primarily interested in the XML encoding of SBML. The constraint expression use="optional" means that the indicated field is optional and may be omitted in a particular instance in a model. The constraint expression minOccurs="0" likewise means that the indicated field is optional; this alternate form of expression must be used in XML Schema for those fields that are containers (i.e., fields that are encoded as subelements in XML).

SBase contains three fields, all of which are optional: metaid, notes and annotation.

3.1.1 The metaid Field

The metaid field is present for supporting metadata annotations using RDF. It has a data type of ID (the XML identifier type), and serves as anchors for metadata references. Metadata expressed using RDF can

be placed anywhere within an sbml element and its subelements, except within MathML elements. The metadata elements can include RDF description elements in which the RDF describes attributes contain the values of the metaid fields of SBML elements in the model. The form of the RDF element content in SBML should follow the form described in the CellML Metadata Specification (Cuellar et al., 2002).

3.1.2 The notes Field

The field notes in SBase is a container for XHTML content. It is intended to serve as a place for storing optional information intended to be seen by humans. An example of the kind of information that would be appropriate to place inside notes is user comments about a particular component of the model. Every data object derived directly or indirectly from type SBase can have a separate value for notes, allowing users considerable freedom when adding comments to their models.

3.1.3 The annotation Field

Finally, SBase includes the field called annotation to provide a container for software-generated annotations that are *not* intended to be seen by humans. This field is a container for arbitrary data (XML type any). As with the user-visible notes field, every data object can have its own value for annotation. Guidelines for using this field are given in Section 3.2.

3.1.4 The SBML Type Inheritance Hierarchy

The overall SBML inheritance hierarchy is depicted in Figure 2. In addition to the relationships shown, all substructures such as trigger on Event and the listOf______ lists are also derived from SBase. (However, the notes and annotation elements contained inside SBase are not derived from SBase.)

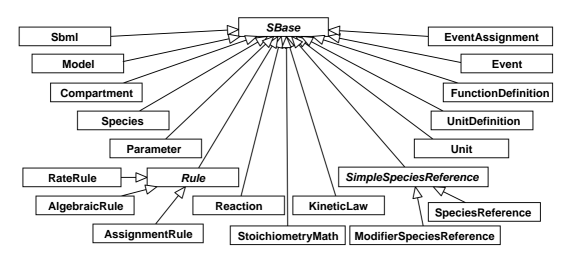


Figure 2: A UML diagram of the inheritance hierarchy of major data types in SBML. Open arrows indicate inheritance, pointing from inheritors to their parents (Eriksson and Penker, 1998; Oestereich, 1999).

3.2 Guidelines for the Use of the annotation Field in SBase

The annotation field in the definition of SBase is formally unconstrained in order that software developers may attach any information they need to the structures in an SBML model. However, it is important that this facility not be misused accidentally. In particular, it is critical that information essential to a model definition is *not* stored in annotation. Parameter values, functional dependencies between model structures, etc., should not be recorded as annotations.

Here are examples of the kinds of data that may be appropriately stored in annotation: (a) information about the graphical layout of model components; (b) application-specific processing instructions that do not change the essence of a model; (c) identification information for cross-referencing components in a model

with items in a database.

Different applications may use XML Namespaces (Bray et al., 1999) to specify the intended vocabulary of a particular annotation. Here is an example. Suppose a particular application needs to annotate data structures in an SBML model definition with screen layout information and a time stamp. The application's developers should choose a URI (*Universal Resource Identifier*; Harold and Means 2001; W3C 2000a) reference that uniquely identifies the vocabulary the application will use for such annotations, and a prefix string for the annotations. For illustration purposes, let us say the URI reference is "http://www.mysim.org/ns" and the chosen prefix is mysim. An example of an annotation might then be as follows:

The namespace prefix mysim is used to qualify the XML elements mysim:nodecolors and mysim:timestamp; presumably these symbols have meaning to the application. This example places the XML Namespace information on annotation itself rather than on a higher-level enclosing construct or the enclosing document level, but other placements would be valid as well (Bray et al., 1999).

The use of XML Namespaces permits multiple applications to place annotations on XML elements of a model without risking interference or element name collisions. Annotations stored by different simulation packages can thus coexist in the same model definition. Although XML Namespace names must be URI references, an XML Namespace name is not required to be directly usable in the sense of identifying an actual, retrieval document or resource on the Internet (Bray et al., 1999). "http://www.mysim.org/" is a namespace name or URI in the example above. The name is simply intended to enable unique identification of constructs, and using URIs is a common and simple way of creating a unique name string. For the convenience of developers of simulation and analysis tools, we reserve certain namespace names for use with annotations in SBML. These reserved names are listed in Table 1.

Note that the namespaces being referred to here are XML Namespaces specifically in the context of the annotation field on SBase. The namespace issue here is unrelated to the namespaces discussed in Section 3.5 in the context of SId and symbols in SBML.

3.3 The id and name Fields on SBML Components

As will become apparent below, most structures in SBML include two common fields: id and name. The id field is usually required for most structures and is used to identify a component within the model definition.

```
http://www.sbml.org/2001/ns/basis
                                           http://www.sbml.org/2001/ns/jdesigner
http://www.sbml.org/2001/ns/biocharon
                                           http://www.sbml.org/2001/ns/jigcell
http://www.sbml.org/2001/ns/bioreactor
                                           http://www.sbml.org/2001/ns/jsim
http://www.sbml.org/2001/ns/biosketchpad
                                           http://www.sbml.org/2001/ns/libsbml
http://www.sbml.org/2001/ns/biospice
                                           http://www.sbml.org/2001/ns/mathsbml
                                           http://www.sbml.org/2001/ns/mcell
http://www.sbml.org/2001/ns/celldesigner
http://www.sbml.org/2001/ns/cellerator
                                           http://www.sbml.org/2001/ns/netbuilder
                                           http://www.sbml.org/2001/ns/pathdb
http://www.sbml.org/2001/ns/copasi
http://www.sbml.org/2001/ns/cytoscape
                                           http://www.sbml.org/2001/ns/promot
                                           http://www.sbml.org/2001/ns/sbedit
http://www.sbml.org/2001/ns/dbsolve
http://www.sbml.org/2001/ns/ecell
                                           http://www.sbml.org/2001/ns/sigpath
http://www.sbml.org/2001/ns/gepasi
                                           http://www.sbml.org/2001/ns/stochsim
http://www.sbml.org/2001/ns/isys
                                           http://www.sbml.org/2001/ns/vcell
http://www.sbml.org/2001/ns/jarnac
                                           http://www.sbml.org/2001/ns/winscamp
```

Table 1: Reserved XML Namespace names in SBML Level 2.

Other SBML structures can refer to the component using this identifier. Section 3.4 defines the data type SId used for the id field, and Section 3.5 describes the scoping and namespace rules for these identifiers. The equality of SId values is determined by an exact character sequence match, i.e. in a case sensitive manner, this applies to all uses of SId including the identification of unit definitions.

In contrast to the id field, the name field is optional and is not intended to be used for cross-referencing purposes within a model. Its purpose instead is to provide a human-readable label for the component. The data type of the name field is the type string defined in XML Schema (Biron and Malhotra, 2000; Thompson et al., 2000). This type includes all Unicode characters (Unicode Consortium, 1996) except for two delimiter characters, 0xFFFE and 0xFFFF (Biron and Malhotra, 2000). The ampersand (&) character must be escaped (using the entity &). The apostrophe (") or single-quote character (') characters must be escaped (using the entities ' or " respectively) when those characters are used to delimit a string attribute value. Other XML built-in character or entity references e.g < or &x1A; can be used. No restrictions as to the content of name fields are imposed by SBML beyond those defined by the string type of XML Schema.

The recommended practice for handling name is as follows. If a software tool has the capability for displaying the content of name fields, it should display this content to the user as a component's label instead of the component's id field. If the user interface does not have this capability (e.g., because it cannot display or use special characters in symbol names), or if the name field is missing on a given component, then the user interface should display the value of the id field instead. (Script language interpreters are especially likely to display id fields instead of name fields.)

As a consequence of the above, authors of systems that automatically generate the values of id fields should be aware some systems may display the id's to the user. Authors therefore may wish to take some care to have their software create id values that are easy for humans to type and read.

An additional point worth mentioning is although there are restrictions on the uniqueness of id values (see Section 3.5 below), there are no restrictions on the uniqueness of name values in a model. This allows a software package more leeway in assigning component identifiers. For example, a species in an SBML model must be located in a compartment, which means that if the same species appears in multiple compartments (e.g., in the context of a transport reaction), they must be given different identifiers. It is currently the case that users and software differ sharply in philosophy about how to treat this situation: some treat these as different species, and others treat them as the same species located in different places. Those in the latter group often want to use the same name but have different id values for the differently-localized "instances" of the species. The freedom from restrictions on name values enables SBML to accommodate both philosophies.

3.4 Type SId

The type SId is the type of the id field found on the majority of SBML components. SId is a data type derived from the basic XML type string, but with restrictions about the types of characters permitted and the sequence in which they may appear. Its definition is shown in Figure 3.

```
letter ::= 'a'..'z','A'..'Z'
digit ::= '0'..'9'
nameChar ::= letter | digit | '_'
name ::= ( letter | '_' ) nameChar*
```

Figure 3: The definition of the type SId expressed in the variant of BNF used by the XML 1.0 specification (Bray et al., 2000). The characters (and) are used for grouping, and the character * indicates "zero or more times".

The SId is purposefully not derived from the XML ID type. Using XML's ID would force all SBML identifiers to exist in a single global namespace, which would affect not only the form of local parameter definitions but also future extensions for supporting model/submodel composition. Further, the use of the ID type for SBML identifiers would have limited utility because MathML ci elements are not of the type IDREF (see Section 3.6). If the IDREF-ID linkage cannot be exploited in MathML constructs, the utility of the XML ID type is greatly reduced.

3.5 Component Identifiers and Namespaces in SBML

A biochemical network model can contain a large number of components representing different parts of a model. This leads to a problem in deciding the scope of an identifier: in what contexts does a given identifier X represent the same thing? The approaches used in existing simulation packages tend to fall into two categories that we may call global and local. The global approach places all identifiers into a single global namespace, so that an identifier X represents the same thing wherever it appears in a given model definition. The local approach places symbols in different namespaces depending on the context, where the context may be, for example, individual rate laws. The latter approach means that a user may use the same identifier X in different rate laws and have each instance represent a different quantity.

The fact that different simulation programs may use different rules for identifier resolution poses a problem for the exchange of models between simulation tools. Without careful consideration, a model written out in SBML format by one program may be misinterpreted by another program. SBML Level 2 must therefore include a specific set of rules for treating identifiers and namespaces.

The namespace rules in SBML Level 2 are relatively straightforward and are intended to avoid this problem with a minimum of requirements on the implementation of software tools:

- The identifiers (i.e., the values of the field id) of functions, compartments, species, reactions, events and model-level parameters reside in the same global namespace. This means, for example, that a reaction and a species definition cannot both have the same identifier.
- Each reaction definition (see Section 4.9) establishes a private local namespace for local parameter identifiers. Within the definition of a given reaction, local parameter identifiers introduced in that reaction override (shadow) identical identifiers in the global namespace.
- Unit identifiers (the values of the field id in the UnitDefinition structure) exist in a separate global namespace distinct from other identifiers.

The set of rules above can enable software packages using either local or global namespaces for parameters to exchange SBML model definitions. In particular, software environments using local namespaces for parameters internally should be able to accept SBML model definitions without needing to change component identifiers. Environments using a global namespace for parameters internally can perform a simple manipulation of the identifiers of local parameter elements within reaction definitions to avoid name collisions. (An example approach for the latter would be the following: when receiving an SBML-encoded model, prefix each parameter identifier inside each reaction with a string constructed from the reaction's identifier; when writing an SBML-encoded model, strip off the prefix.)

The namespace rules described here will hopefully provide a clean transition path to future levels of SBML, when submodels are introduced (Section 6.1). Submodels will provide the ability to compose one model from a collection of other models. This capability will have to be built on top of SBML Level 2's namespace organization. A straightforward approach to handling namespaces is to make each submodel's space be private. The rules governing namespaces within a submodel can simply be the Level 2 namespace rule described here, with each submodel having its own (to itself, global) namespace.

3.6 Mathematical Formulas in SBML Level 2

Mathematical expressions in SBML Level 2 are represented using MathML 2.0 (W3C, 2000b), the XML standard for describing mathematics in machine-readable format. It is used in the definitions of functions (Section 4.3), rules (Section 4.8), kinetic laws (Section 4.9.7), and stoichiometries (Section 4.9). The KineticLaw and Rule structures each have a single MathML math subelement, and a function definition has a single lambda subelement.

The XML namespace URI for all MathML elements is "http://www.w3.org/1998/Math/MathML". [See the W3C document by Bray et al. (1999) for more information about using XML namespaces.] The examples below illustrate the use of this namespace and MathML in SBML.

3.6.1 Subset of MathML Used in SBML Level 2

The subset of MathML elements used in SBML Level 2 is similar to that used by CellML and is itemized below:

- token: cn, ci, csymbol, sep
- basic content: apply, piecewise, piece, otherwise
- relational operators: eq, neq, gt, lt, geq, leq
- arithmetic operators: plus, minus, times, divide, power, root, abs, exp, ln, log, floor, ceiling, factorial
- logical operators: and, or, xor, not
- qualifiers: degree, bvar, logbase
- trigonometric operators: sin, cos, tan, sec, csc, cot, sinh, cosh, tanh, sech, csch, coth, arcsin, arccos, arctan, arcsec, arccsc, arccot, arcsinh, arccosh, arctanh, arcsech, arcsch, arccoth
- constants: true, false, notanumber, pi, infinity, exponentiale
- annotation: semantics, annotation, annotation-xml

The inclusion of logical operators, relational operators, piecewise, piece, and otherwise elements facilitates the encoding of discontinuous expressions. Elements for representing partial differential calculus are not included. We anticipate that the requirements for partial differential calculus will be addressed in proposals for SBML Level 3 geometry representations (see Section 6.1).

Only the following attributes of MathML elements can occur in SBML documents:

- style, class and id on any element;
- encoding and definitionURL on csymbol elements; and
- type on cn elements.

The MathML default values for missing attributes should be assumed in SBML. These restrictions on attributes are designed to confine the MathML elements to their default semantics and to avoid conflicts in the interpretation of the type of token elements.

3.6.2 Use of cn elements in MathML Expressions in SBML

The type attribute on MathML cn elements should only have the following values in SBML documents: e-notation, real, integer, and rational. The type attribute can default to real.

3.6.3 Use of ci elements in MathML Expressions in SBML

The content of a ci element must obey MathML whitespace rules and contain an identifier that is declared elsewhere in the model. The set of possible identifiers that can appear in a ci element depends on the containing structure in which the ci is used:

- If ci appears in the body of a function definition, the referenced identifier must be either (i) one of the declared arguments to the function, or (ii) the identifier of a previously defined function.
- In all other situations, the referenced identifier must be the identifier of a species, compartment, parameter or function declared in the model. The following are the only possible interpretations of using such an identifier in SBML:

- Species identifier: When a species identifier occurs in a ci element, it represents the quantity of that species in terms of its concentration (where "concentration" is meant somewhat broadly).
 The units associated with a species identifier are the units of the species which are defined in Section 4.6.4.
- Compartment identifier: When a compartment identifier occurs in a ci element, it represents the size of the compartment. The units associated with the size of the compartment are those given on the Compartment structure that declares the identifier (see Section 4.5.4)
- Parameter identifier: When a parameter identifier occurs in a ci element, it represents the value assigned to that parameter. The units associated with the parameter value are the units assigned in its instance of a Parameter structure; see Section 4.7.3.
- Function identifier: When a function identifier occurs in a ci element, it represents a call to that function. Function references in MathML occur in the context of using MathML's apply and often involve supplying arguments to the function; see Section 4.3.

3.6.4 Use of csymbol elements in MathML Expressions in SBML

SBML Level 2 uses the MathML csymbol element to denote certain built-in mathematical entities without introducing reserved names into the component identifier namespace. The encoding field of csymbol should be set to text. The definitionURL should be set to one of the following predefined SBML symbol URLs:

- http://www.sbml.org/sbml/symbols/time. This represents the current simulation time. The units of the current time entity are determined from the built-in time of Table 3 on page 15.
- http://www.sbml.org/sbml/symbols/delay. This represents a delay function. The delay function has the form delay(x, d), taking two arguments. Its value is the value of argument x at d time units before the current time. The units of the d parameter are determined from the built-in time. The delay function is useful for representing biological processes having a delayed response, but where the detail of the processes and delay mechanism is not relevant to the operation of a given model.

The following examples demonstrates these concepts. The XML fragment below encodes the formula x + t, where t is the built-in symbol for time.

As a further example, the following XML fragment encodes the equation k + delay(x, 0.1) or alternatively $k_t + x_{t-0.1}$:

Note that it is not necessary for a parser to access the resource pointed to by the "definitionURL:"; in this context, the URL should be interpreted as a URI. Also, the content of the csymbol element is for rendering purposes only and can be ignored by the parser.

Section 5.8 contains a complete model which uses a delay function.

4 SBML Components

In this section, we define each of the major data structures in SBML. To provide illustrations of their use, we give partial model definitions in XML. Section 5 provides many full examples of SBML in XML.

4.1 The SBML Container

The outermost portion of an SBML Level 2 model definition consists of a single Sbml structure enclosing a single Model structure (see next Section). The definition of Sbml is shown in Figure 4.

```
Sbml

level : positiveInteger { use="required" fixed = "2"}

version : positiveInteger { use="required" fixed = "1"}
```

Figure 4: The definition of Sbml. Additional fields are inherited from SBase but are not shown here.

The XML namespace URI for SBML Level 2 is "http://www.sbml.org/sbml/level2". All SBML Level 2 elements should be encoded using this URI by assigning this URI to either the default namespace or a tag prefix. The character encoding for SBML is UTF-8. SBML documents should include the encoding attribute with the value UTF-8 in the XML prologue.

In the transformation of UML to XML used in this document, the Sbml structure is turned into an element named sbml. The element has two required attributes: level and version. For SBML Level 2 Version 1, these attributes must be set to "2" and "1", respectively. (The version attribute is present in case SBML Level 2 must be revised in the future to correct errors.)

The following is an abbreviated example of the outermost content of an SBML model definition in XML:

4.2 Models

The Model structure is the highest-level construct in an SBML data stream or document. The UML definition of Model is shown in Figure 5 on the next page. Only one component of type Model is allowed per instance of an SBML document or data stream, although it does not necessarily need to represent a single biological entity.

Model serves as a container for FunctionDefinition, UnitDefinition, Compartment, Species, Parameter, Rule, Reaction and Event components. All of these components are optional; that is, the lists in each of the respective fields are permitted to have zero length. (However, there are dependencies between components, such that defining some requires defining others. For example, as explained in other sections below, defining a species requires defining a compartment, and defining a reaction requires defining a species.)

The Model structure has an optional field, id, used to give the model an identifier. The identifier must be a text string conforming to the syntax permitted by the SId data type described in Section 3.4. Model also has an optional name field, of type string. The name and id fields should be used as described in Section 3.3.

In the XML encoding of an SBML model, the lists of species, compartments, unit definitions, parameters, reactions, function definitions, rules and events are translated into lists of XML elements enclosed within

Model

id : SId {use="optional"}
name : string {use="optional"}

functionDefinition: FunctionDefinition[0..*]
unitDefinition: UnitDefinition[0..*]
compartment: Compartment[0..*]

species : Species[0..*]
parameter : Parameter[0..*]

rule : Rule[0..*]
reaction : Reaction[0..*]
event : Event[0..*]

Figure 5: The definition of Model. Additional fields are inherited from SBase.

elements of the form listOf____s, where the blank is replaced by the name of the component type (e.g., "Reaction"). The resulting XML data object has the form illustrated by the following skeletal model:

```
<model id="My_Model">
   <listOfFunctionDefinitions>
   <listOfUnitDefinitions>
   </listOfUnitDefinitions>
   <listOfCompartments>
   </or>
   <listOfSpecies>
   </listOfSpecies>
   <listOfParameters>
   </listOfParameters>
   tOfRules>
   </listOfRules>
   <listOfReactions>
   </listOfReactions>
   <listOfEvents>
   </listOfEvents>
</model>
```

Readers may wonder about the motivations for the listOf____s notation. A simpler approach to creating the lists of components would be to place them all directly at the top level under <model> . . . </model>. We chose instead to group them within XML elements named after listOf____s, because we believe this helps organize the components and makes visual reading of model definitions easier. Further details of how listOf____ elements implement UML lists is described in Appendix A.

4.3 Function Definitions

The FunctionDefinition structure associates an identifier with a function definition. The identifier can then be used in any subsequent MathML apply elements. FunctionDefinition is shown in Figure 6 on the following page.

The FunctionDefinition structure has three fields, id, name and math.

4.3.1 The id and name fields

The id and name fields have types SId and string, respectively, and operate in the manner described in Section 3.3. MathML ci elements can refer to the function defined by a FunctionDefinition using the

```
FunctionDefinition

id : SId
name : string {use="optional"}
math : (lambda:Lambda) {namespace="http://www.w3.org/1998/Math/MathML" }
```

Figure 6: The definition of FunctionDefinition. Fields inherited from SBase are omitted here but are assumed.

value of its id field.

4.3.2 The math field

The math field is a container for MathML content that defines the function. The content of this field can only be a MathML lambda element. This is the only place in SBML where a lambda element can be used. The function is only available for use in other MathML elements that follow the FunctionDefinition structure in an SBML model. (These restrictions prevent recursive and mutually-recursive functions from being expressed.) The lambda element can contain any of the elements in the MathML subset listed in Section 3.6.1 but not any further lamdba elements.

The following abbreviated SBML example shows a FunctionDefinition structure defining pow3(x) as representing x^3 :

4.4 Unit Definitions

Units may be supplied in a number of contexts in an SBML model. A facility for defining units is convenient to have, so that combinations of units can be given abbreviated names and so that the default units can be redefined. These are the motivations for the provision of the UnitDefinition and Unit structures in SBML, whose definitions are shown in Figure 7.

```
UnitDefinition

id : SId
name : string { use="optional" }
unit : Unit[1..*]

kind : UnitKind
exponent : integer { use="optional" default="1"}
scale : integer { use="optional" default="0"}
multiplier : double { use=optional default="1"}
offset : double { use=optional default="0"}
```

Figure 7: The definition of UnitDefinition and Unit.

4.4.1 The UnitDefinition structure

An instance of a UnitDefinition consists of an id field of type SId, an optional string field name and an optional list of structures of type Unit. As mentioned in Section 3.5, unit identifiers defined by the id field are considered to be in a separate global namespace distinct from the namespace of other identifiers in a model; thus, unit identifiers cannot collide with the identifiers of species, compartments, reactions etc. The approach to defining units in SBML is compositional; for example, $meter\ second^{-2}$ is constructed by combining, within the same UnitDefinition Unit list, a Unit representing meter with a Unit representing $second^{-2}$.

4.4.2 The Unit structure

The Unit data structure has one required field, kind, whose value must be taken from UnitKind, an enumeration of base units. The possible values of UnitKind are given in Table 2.

ampere	farad	joule	lux	radian	volt
becquerel	${\tt gram}$	katal	metre	second	watt
candela	gray	kelvin	mole	siemens	weber
Celsius	henry	kilogram	newton	sievert	
coulomb	hertz	litre	ohm	steradian	
<u>dimensionless</u>	item	lumen	pascal	tesla	

Table 2: The possible values of kind in a UnitKind structure. All are names of base or derived SI units (Bureau International des Poids et Mesures, 2000), except for "dimensionless" and "item", which are SBML additions important for handling certain common situations. "Dimensionless" is intended for cases where a quantity does not have units, and "item" for expressing such things as "N items" (e.g., "100 molecules"). Although "Celsius" is capitalized, for simplicity, SBML requires that these unit names be treated in a case-insensitive manner. Also, note that the gram and litre are not strictly part of SI; however, they are so commonly used in SBML's areas of application that they are included as predefined unit names. (The standard SI unit of mass is in fact the kilogram, and volume is defined in terms of cubic meters.)

Note that the set of acceptable values for the field kind does not include units defined by UnitDefinition structures. This means that the units definition feature in SBML is not hierarchial—user-defined units cannot be built on top of other user-defined units, only on top of base units. SBML differs from CellML in this respect; CellML does allow the construction of hierarchial unit definitions.

An instance of a Unit definition represents a (possibly transformed) reference to a base unit chosen from UnitKind. The formula for a single transformation represented by Unit definition is as follows (where u is the original base unit and u_{new} is the new unit):

$$u_{new} = (multiplier \times 10^{scale} \times u^{exponent}) + offset$$

The optional exponent field on Unit represents an exponent on the unit. Its default value is "1" (one). For the example mentioned at the beginning of this section, $second^{-2}$ would be obtained by using kind="second" and exponent="-2". A Unit structure also has an optional scale field; its value must be an integer exponent for a power of ten multiplier used to set the scale of the unit. For example, a unit that has a kind value of "gram" and a scale value of "-3" signifies $10^{-3} * gram$, or milligrams. The default value of scale is "0" (zero), because $10^0 = 1$.

The optional multiplier field can be used to multiply the kind unit by a real-numbered factor; this enables the definition of units that are not power-of-ten multiples of SI units. For instance, a multiplier of 0.3048 could be used to define "foot" as a measure of length in terms of a metre. The multiplier field has a default value of "1" (one). Finally, the offset field is used to represent the addition of a constant in the transformation the kind unit. For example, an offset value of "32.0" would be needed to define Fahrenheit in terms of degrees Celsius. The offset field has a default value of "0" (zero).

The composition of n Unit structures within a UnitDefinition to create more complex units involves a

linear product according to the following formula:

```
u_{new} = m_1 \times \ldots \times m_n \times 10^{s_1} \times \ldots \times 10^{s_n} \times u^{e_1} \times \ldots \times u^{e_n}
```

The following example illustrates the definition of an abbreviation named "mmls" for the units $mmol\ l^{-1}\ s^{-1}$:

The following example defines Fahrenheit:

4.4.3 Built-in Units

There are five special unit names in SBML, listed in Table 3, corresponding to the five types of quantities or built-in units that play roles in biochemical reactions: amount of substance, volume, area, length and time. SBML defines defaults for these built-in units, see the 3rd column of Table 3, all with default scale and offset values of zero and default multiplier values of one. The various components of a model, such as parameters, can use only the predefined units from Table 2 on the preceding page, new units defined in unit definitions, or the five predefined names of built-in units "substance", "volume", "area", "length", and "time" from Table 3.

Name	Possible Scalable Units	Default Units
substance	mole, item	mole
volume	litre, cubic metre	litre
area	square metre	square metre
length	metre	metre
time	second	second

Table 3: SBML's built-in units.

Within certain limits, a model may change the built-in units by reassigning the keywords "substance", "length", "area", "time", and "volume" in a UnitDefinition. The second column in Table 3 lists the set of units that should be used in redefining a given built-in unit.

The following example illustrates how to change the built-in units of volume to be milliliters. If this definition appeared in a model, the units of volume on all components that did not explicitly specify different units would be changed to milliliters.

Software developers are asked to pay special attention to the units used in an SBML model. Different users and developers sometimes make different assumptions about units, and these assumptions may not correspond to what is defined in SBML. Sections 3.6.3, 4.6 and 4.9.7 have particularly important notes about the usage of units in SBML.

4.5 Compartments

A compartment in SBML represents a bounded space in which species are located. Compartments do not necessarily have to correspond to actual structures inside or outside of a cell, although models are often designed that way. The definition of Compartment is shown in Figure 8.

id: SId name: string {use="optional"} spatialDimensions: integer { maxInclusive="3" minInclusive="0" use="optional" default="3"} size: double {use="optional"} units: SId {use="optional"} outside: SId {use="optional"} constant: boolean {use="optional" default="true"}

Figure 8: The definition of Compartment. Fields inherited from SBase are omitted here but are assumed.

It is worth pointing out that, although compartments are optional in the overall definition of Model (see Section 4.2), every species in an SBML model must be located in a compartment. This in turn means that if a model declares any species, that model must also declare at least one compartment.

4.5.1 The id and name Fields

Compartment has one required field, id, of type SId, to give the compartment a unique identifier by which other parts of an SBML model definition can refer to it. A compartment can also have an optional name field of type string. Identifiers and names should be used according to the guidelines described in Section 3.3.

4.5.2 The spatialDimensions Field

A Compartment structure has an optional field spatialDimensions, whose value must be a positive integer indicating the number of spatial dimensions possessed by the compartment. The maximum value of spatialDimensions is "3", meaning a three-dimensional structure (a volume). Other permissible values are "2" (for a two-dimensional area), "1" (for a one-dimensional curve), and "0" (for a point). The default value is "3"

4.5.3 The size Field

Each compartment has an optional floating-point field named size, representing the total size of the compartment. The size field enables concentrations of species to be calculated in the absence of geometry information. A missing value for size for a given compartment signifies that the value is either unknown, not required for analysis, or available from an external data source. Note in particular that in SBML Level 2, a missing size value does *not* imply that the compartment size is 1. (This is unlike the definition of compartment volume in SBML Level 1.). The size field must not be present if the spatialDimensions field is 0.

4.5.4 The units Field

The units associated with the compartment's size value may be explicitly set using the optional field units. The value chosen for this field must be either one of the base units from Table 2 on page 14, or the built-in units "volume", "area", "length" or "dimensionless", or a new unit defined by a unit definition in the

enclosing model. The type of units assigned to the units field must also agree with the number of spatial dimensions of the compartment; that is, they must be units of volume if the value of spatialDimensions is "3"; they must be units of area if the value of spatialDimensions is "2"; they must be units of length if the value of spatialDimensions is "1"; and they must be "dimensionless" if the value of spatialDimensions is "0". The default units depends on the value of the compartment's spatialDimensions attribute according to the following rule: for spatial dimensions of 3, 2, 1 or 0, the compartment has the default units of volume, area, length and dimensionless, respectively. (See Table 3 on page 15 and Table 2 on page 14.)

The units of the compartment size, as defined by the units attribute, are used as in the following ways.

- The value of the units field is used as the units of size field of the compartment structure (see Section 4.5.3).
- The value of the units field is used as the units of the compartment identifier when it appears as a numerical quantity in a mathematical formula expressed in MathML (discussed in Section 3.6.3).
- The value of the units field is used as the units of the math field of the AssignmentRule structures determining the compartment's size (see Section 4.8.2).
- In RateRule structures that set the rate of change of the compartment's size (Section 4.8.3), the units on the rule's math field are those in the compartment's units field divided by the default *time* units. (In other words, the units for the rate of change of compartment size are *compartment size/time* units.)

4.5.5 The constant Field

A Compartment also has an optional boolean field called constant which indicates whether the compartment's size stays constant or can vary during a simulation. A value of "false" indicates that the compartment's size can be determined by rules (see Section 4.8), and the value of the size field should be taken as being the initial size of the compartment. The default value for the constant field is "true" because in the most common modeling scenarios at the time of this writing, compartment sizes remain constant. The constant field must default to or be set to "true" if the spatialDimensions field is 0.

4.5.6 The outside Field

The optional field outside of type SId can be used to express containment relationships between compartments. If present, the value of outside for a given compartment must be the name of another compartment enclosing it, or in other words, the compartment that is "outside" of it. This enables the representation of simple topological relationships between compartments, for those simulation systems that can make use of the information (e.g., for drawing simple diagrams of compartments).

Although containment relationships are partly taken into account by the compartmental localization of reactants and products, it is not always possible to determine purely from the reaction equations whether one compartment is meant to be located within another. In the absence of a value for outside, compartment definitions in SBML Level 2 do not have any implied spatial relationships between each other. For many modeling applications, the transfer of substances described by the reactions in a model sufficiently express the relationships between the compartments. (As discussed in Section 6.1, we expect that SBML Level 3 will introduce the ability to define geometries and spatial qualities.)

4.5.7 Examples

The following example illustrates two compartments in an abbreviated SBML example of a model definition:

The following is an example of using outside to model a cell membrane. To express that a compartment named B has a membrane that is modeled as another compartment M, which in turn is located within another compartment A, one would write:

4.6 Species

The term *species* refers to chemical entities that take part in reactions. These include simple ions (e.g., protons, calcium), simple molecules (e.g., glucose, ATP), large molecules (e.g., RNA, polysaccharides, and proteins), and others. The Species data structure is intended to represent these entities. Its definition is shown in Figure 9.

```
id: SId
name: string {use="optional"}
compartment: SId
initialAmount: double {use="optional"}
initialConcentration: double {use="optional"}
substanceUnits: SId {use="optional"}
spatialSizeUnits: Sid {use="optional"}
boundaryCondition: boolean {use="optional" default="false"}
charge: integer {use="optional"}
constant: boolean {use="optional" default="false"}
```

Figure 9: The definition of Species. As usual, fields inherited from SBase are omitted here but are assumed.

4.6.1 The id and name attributes

As with other major structures in SBML, Species has an mandatory field, id, used to give the species an identifier. The identifier must be a text string conforming to the syntax permitted by the SId data type described in Section 3.4. Species also has an optional name field, of type string. The name and id fields should be used as described in Section 3.3.

4.6.2 The compartment Field

The required field compartment, also of type SId, is used to identify the compartment in which the species is located. The field's value must be the identifier of an existing Compartment structure. It is important to note that there is no default value for the compartment field on Species: every species in an SBML model must be assigned a compartment, and consequently, a model must define at least one compartment if that model contains any species.

4.6.3 The initial Amount and initial Concentration Fields

The optional fields initialAmount and initialConcentration, both having a data type of double, are used to set the initial quantity of the species in the named compartment. These fields are mutually exclusive i.e. only one can have a value on any given instance of a Species structure. Also, initialConcentration must not have a value if the species' compartment has a spatialDimensions value of "0". (The reason is simply that there cannot be a concentration defined when the compartment has no spatial extent.) Missing initialAmount or initialConcentration values implies that their values are either unknown, set by an

assignment rule, not required for analysis or available from an external data source.

The units of the value in the initialAmount field is that given by the substanceUnits field of the species structure. The units of the value in the initialConcentration field are the *units of the species* see Section 4.6.4.

4.6.4 The substanceUnits and spatialSizeUnits Fields

The units associated with species referred to as the *units of the species* are determined via the optional fields substanceUnits and spatialSizeUnits. The *units of the species* can be thought of as the concentration units of the species if one adopts a broad definition of concentration. The *units of the species* are of the form *substance/size* units if the compartment's spatialDimensions are non-zero or are of the form *substance* if spatialDimensions are zero. The units of *substance* are those defined in the substanceUnits, and the *size* units are those given in the spatialSizeUnits attribute.

In the case of both substanceUnits and spatialSizeUnits fields the value chosen must be either a base unit from Table 2 on page 14, a built-in unit from Table 3 on page 15, or a new unit defined by a unit definition in the enclosing model. The chosen units for substanceUnits must be a variant of mole or item units. The substanceUnits field defaults to the built-in unit "substance" shown in Table 3 on page 15.

The type of units assigned to the spatialSizeUnits field must agree with the number of spatial dimensions of the species' compartment; that is, they must be units of volume if the value of the compartment's spatialDimensions is "3"; they must be units of area if the value of spatialDimensions is "2"; and they must be units of length if the value of spatialDimensions is "1". The spatialSizeUnits must not have a value if the value of spatialDimensions is "0". The default value of the spatialSizeUnits is the value of the units field of the species' compartment.

The units of the species are used in the following ways.

- The species initialConcentration field has these units. (see Section 4.6.3).
- The species identifier has these units when it appears as a numerical quantity in a mathematical formula expressed in MathML (discussed in Section 3.6.3).
- The math field of the classAssignmentRule structures determining the species' concentration (see Section 4.8.2) has these units.
- In RateRule structures that set the rate of change of the species' concentration (Section 4.8.3), the units on the rule's math field are the *units of the species* divided by the default *time* units.

4.6.5 The constant and boundaryCondition Fields

The Species structure has optional boolean field named constant which is used to indicate whether the concentration of that species can vary during a simulation. The default value is "false", indicating that the species' concentration can be determined by rules and reactions.

Another optional field defined for Species is boundaryCondition. By default, when a species is a product or reactant of one or more reactions, its concentration is determined by those reactions. In SBML, it is possible to indicate that a given species' concentration is not determined by the set of reactions even when that species occurs as a product or reactant; i.e., the species is on the boundary of the reaction system but is a component of the rest of the model. The boolean field boundaryCondition can be used to indicate this. The value of the field defaults to "false", indicating that by default, the species is part of the reaction system. Table 4 on the next page shows how to interpret the combined values of the boundaryCondition and constant fields. In practice, a boundaryCondition value of "true" means that a differential equation derived from the reaction definitions should not be generated for the species. The example model in section 5.5 contains all four possible combinations of the boundaryCondition and constant attributes on species elements.

constant value	boundaryCondition value	${\it assignment}$	can be reactant or product	concentration is changed by
true	true	no	yes	never changes
false	true	yes	yes	rule
true	false	no	no	never changes
false	false	yes	yes	reactions or rule but not both

Table 4: How to interpret the values of the constant and boundary Condition fields of the Species structure.

4.6.6 The charge Field

Finally, the optional field charge on Species takes an integer indicating the charge on the species (in terms of electrons, not the SI unit Coulombs). This may be useful when the species involved is a charged ion such as calcium (Ca^{2+}) .

4.6.7 Example

The following example shows two species definitions within an abbreviated SBML model definition. The example shows that species are listed under the heading listOfSpecies in the model:

4.7 Parameters

A Parameter structure is used to declare a variable for use in mathematical formulas in an SBML model definition. By default, parameters have constant value for the duration of a simulation and for this reason are called "parameters" instead of variables in SBML. The definition of Parameter is shown in Figure 10.

```
Parameter

id : SId
name : string {use="optional"}
value : double {use="optional"}
units : SId {use="optional"}
constant : boolean {use="optional" default="true"}
```

Figure 10: The definition of Parameter. Fields inherited from SBase are omitted here but are assumed.

4.7.1 The id and name Fields

Parameter has one required field, id, of type SId, to give the parameter a unique identifier by which other parts of an SBML model definition can refer to it. A parameter can also have an optional name field of type string. Identifiers and names should be used according to the guidelines described in Section 3.3.

4.7.2 The value Field

The optional field value determines the value (of type double) assigned to the identifer. A missing value implies that the value is either unknown, not required for analysis or available from an external data source.

4.7.3 The units Field

The units associated with the value of the parameter are specified by the field units. These units are used when the parameter identifier appears in MathML expressions and in AssignmentRule structures setting the value of the parameter. A RateRule structure which may determine the value of the parameter has units parameter units/time, where parameter units are the units assigned to the parameter and time is the built-in time units. The value assigned to the parameter's units attribute must be chosen from one of the following possibilities: one of the base unit names from Table 2 on page 14; one of the built-in unit names appearing in first column of Table 3 on page 15); or the name of a new unit defined in the list of unit definitions in the enclosing Model structure. There are no constraints on which units can be chosen from these sets.

4.7.4 The constant Field

The Parameter structure has an optional boolean field named constant which indicates whether the parameter's value can vary during a simulation. The field's default value is "true"; a value of "false" indicates that the parameter's value can be changed by rules (see Section 4.8) and that the value is actually intended to be the initial value of the parameter.

4.7.5 Local and Global Parameters

Parameters can be defined in two places in SBML: in lists of parameters defined at the top level in a Model-type structure, and within individual reaction definitions (as described in Section 4.9). Parameters defined at the top level are *global* to the whole model; parameters that are defined within a reaction are local to the particular reaction and (within that reaction) *override* any global parameters having the same names. (See Section 3.5 for further details.) Parameters local to a reaction cannot be changed by rules and therefore are implicitly always constant; thus, parameter definitions within Reaction structures should not have a constant attribute value.

4.7.6 Example

The following is an example of parameters defined at the Model level:

4.8 Rules

In SBML, rules provide a way to create constraints on variables for cases in which the constraints cannot be expressed using reactions (Section 4.9) nor the assignment of an initial value to a component in a model. There are two orthogonal dimensions by which rules can be described. First, there are three different possible functional forms, corresponding to the following three general cases (where x is a variable, f is some arbitrary function, and W is a vector of variables that may include x):

```
\begin{array}{ll} \textit{Algebraic} & \text{left-hand side is zero:} & 0 = f(W) \\ \textit{Assignment} & \text{left-hand side is a scalar:} & x = f(W) \\ \textit{Rate} & \text{left-hand side is a rate-of-change:} & dx/dt = f(W) \end{array}
```

The second dimension concerns the role of variable x in the equations above: x can be the identifier of a compartment (to set its size), a species (to set its concentration), or a parameter (to set its value).

In their general form given above, there is little to distinguish between assignment and algebraic rules. They are treated as separate cases for the following reasons:

- Assignment rules can simply be evaluated to calculate intermediate values for use in numerical methods;
- Some simulators do not contain numerical solvers capable of solving unconstrained algebraic equations;
- Those simulators that *can* solve these *algebraic* equations make a distinction between the different categories listed above; and
- Some specialized numerical analyses of models may only be applicable to models that do not contain algebraic rules.

The approach taken to covering these cases in SBML is to define an abstract Rule structure that contains only one field, math, to hold the right-hand side expression, then to derive subtypes of Rule that add fields to distinguish the cases of algebraic, assignment and rate rules. Figure 11 gives the definitions of Rule and the subtypes derived from it. The figure shows there are three subtypes, AlgebraicRule, AssignmentRule and RateRule defined directly from Rule. These correspond to the cases Algebraic, Assignment and Rate described above respectively.

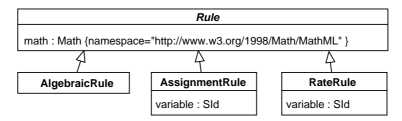


Figure 11: The definition of Rule and derived types.

4.8.1 AlgebraicRule

The rule type AlgebraicRule is used to express equations that are neither assignments of model variables nor rates of change. AlgebraicRule does not add any fields to the basic Rule; its role is simply to distinguish this case from the other cases. An example of the use of AlgebraicRule structures is given in Section 5.4.

4.8.2 AssignmentRule

The rule type AssignmentRule is used to express equations that set the values of variables. The left-hand side (the variable field of an assignment rule) of a rule can refer to the identifier of a species, compartment, or parameter. Two or more RateRule or AssignmentRule structures cannot have the same left-hand side or variable attribute value in an SBML model definition. In all cases, as would be expected, the units of the formula representing the right hand side, the math field, are identical to the units associated with the left hand side, the variable field, when that variable appears in other formulae.

The effects of an AssignmentRule structure are in general terms the same, but differ in the precise details depending on which variable is being set:

• In the case of a species, an AssignmentRule sets the referenced species' concentration to the value determined by the formula in math. The units of the formula are the units of the species as defined in Section 4.6.4. (These units can be thought of as concentration units if a broad definition of concentration is adopted.)

Restrictions: In a given SBML model, there cannot be both a AssignmentRule variable attribute and a SpeciesReference species attribute that have the same value. (See Section 4.9 for the definition of

SpeciesReference.) This means that an assignment rule cannot be defined for a species that is created or destroyed in a reaction. The only exception is when the given species is a boundary condition; i.e., on the Species structure that defines the species the boundaryCondition field is set to "true".

- In the case of a compartment, an AssignmentRule sets the referenced compartment's size to the size determined by the formula in math. The overall units of the formula must match the units specified for the size of the compartment identified by the value of the AssignmentRule's variable field. (See Section 4.5.4 for an explanation of how the units of the compartment's size are determined.)
- In the case of a parameter, an AssignmentRule sets the referenced parameter's value to that determined by the formula in math. The overall units of the formula must match the units defined for the parameter identified by the value of the AssignmentRule's variable field. (See Section 4.7.3 for an explanation of how the units of the parameter are determined.)

4.8.3 RateRule

The rule type RateRule is used to express equations that determine the rates of change of variables. The left-hand side (the variable of a rate rule) can refer to the identifier of a species, compartment, or parameter. Two or more RateRule or AssignmentRule structures cannot have the same left-hand side or variable attribute value in an SBML model definition. In all cases, as would be expected, the units of the formula representing the right hand side, in the math field, are of the form x/time where x are the same units as associated with the symbol in the variable field, when that variable appears in other formulae. time is a built-in unit (see Section 4.4). The effects of a RateRule are in general terms the same, but differ in the precise details depending on which variable is being set:

- In the case of a species, a RateRule sets the rate of change of the species's concentration to the value determined by the formula in math. The overall units of the formula must be concentration/time, where the time units are the built-in units of time described in Section 4.4 and the concentration units are the units of the species as defined in Section 4.6.4.
 - Restrictions: In a given SBML model, there cannot be both a RateRule variable attribute and a SpeciesReference species attribute that have the same value. (See Section 4.9 for the definition of SpeciesReference.) This means that an assignment rule cannot be defined for a species that is created or destroyed in a reaction. The only exception is when the given species is a boundary condition; i.e., on the Species structure that defines the species the boundaryCondition field is set to "true".
- In the case of a compartment, a RateRule sets the rate of change of the compartment's size to the value determined by the formula in math. The overall units of the formula must be size/time, where the time units are the built-in units of time described in Section 4.4 and the size units are the units of size on the compartment identified by the value of the RateRule's variable field. (See Section 4.5.4 for an explanation of how the units of the compartment's size are determined.)
- In the case of a parameter, a RateRule sets the rate of change of the parameter's value to that determined by the formula in math. The overall units of the formula must be of the form x/time where x is the units defined for the parameter identified by the value of the AssignmentRule's variable field and the time units are the built-in units of time described in Section 4.4. (See Section 4.7.3 for an explanation of how the units of the parameter are determined.)

4.8.4 Constraints on rules

SBML specifically does not stipulate the form of the algorithms that can be applied to rules and reactions. For example, SBML does not specify when or how often rules should be evaluated. The constraints described by rules and kinetic rate laws are meant to apply collectively to the set of variable values for a specific time.

The ordering of assignment rules is significant: they are always evaluated in the order given in SBML.

No more than one assignment or rate rule can be defined for a given identifier. No assignment or rate rule can be defined for an identifier whose corresponding structure has the field constant set to true.

An assignment rule for a given identifier overrides the initial value assigned to that identifier; i.e., the initial value should be ignored. This does not mean that any structure declaring an identifier can be omitted if there is an assignment rule for that identifier. For example, there must be a Parameter structure for a given parameter if there is a rule for that parameter.

The math field of an assignment rule structure can contain any identifier in a MathML ci element except for the following: (a) identifiers for which there exists a subsequent assignment rule, and (b) the identifier for which the rule is defined. These constraints are designed to eliminate algebraic loops among the scalar rules; eliminating algebraic loops ensures that assignment rules can be evaluated any number of times without the result of those evaluations changing. As an example, consider the following equations, in the order shown:

$$x = x + 1$$
, $y = z + 200$, $z = y + 100$

If this set of equations were interpreted as a set of assignment rules, it would be invalid because the rule for x refers to x and the rule for y refers to z before z is defined.

4.8.5 Example of Rule Use

This section contains an example set of rules. Consider the following set of equations:

$$k = \frac{k_3}{k_2}, \quad s_2 = \frac{k x}{1 + k_2}, \quad A = 0.10 x$$

This can be encoded by the following scalar rule set (where the definitions of x, s, k, k2, k3 and A are assumed to be located elsewhere in the model and not shown in this abbreviated example):

```
<model>
    <listOfRules>
        <assignmentRule variable="k">
            <notes>
                <xhtml:p>
                    k = k3/k2
                </xhtml:p>
            </notes>
            <math xmlns="http://www.w3.org/1998/Math/MathML">
                <apply>
                    <divide/>
                    <ci> k3 </ci>
                    <ci> k2 </ci>
                </apply>
            </assignmentRule>
        <assignmentRule variable="s2">
            <notes>
                <xhtml:p>
                    s2 = (k * x)/(1 + k2)
                </xhtml:p>
            </notes>
            <math xmlns="http://www.w3.org/1998/Math/MathML">
                <apply>
                    <divide/>
                    <apply>
                        <times/>
                        <ci> k </ci>
                        <ci> x </ci>
                    </apply>
                    <apply>
                        <plus/>
                        <cn> 1 </cn>
                        <ci> k2 </ci>
                    </apply>
                </apply>
            </assignmentRule>
```

```
<assignmentRule variable="A">
            <notes>
                <xhtml:p>
                    A = 0.10 * x
                </xhtml:p>
            </notes>
            <math xmlns="http://www.w3.org/1998/Math/MathML">
                <apply>
                    <times/>
                    <cn> 0.10 </cn>
                    <ci> x </ci>
                </apply>
            </assignmentRule>
   </listOfRules>
</model>
```

4.9 Reactions

A reaction represents any transformation, transport or binding process, typically a chemical reaction, that can transform or change the amount of one or more species. In SBML, a reaction is defined primarily in terms of the participating reactants and products (and their corresponding stoichiometries), along with optional modifier species, an optional kinetic law describing the rate at which the reaction takes place, and optional parameters entering into the kinetic law. These various parts of a reaction are recorded in the SBML Reaction type defined in Figure 12 on the following page.

4.9.1 The id and name Fields

As with most other main structures in SBML, the Reaction data structure includes a required id and an optional name. These must be used according to the guidelines described in Section 3.3.

4.9.2 The reactant, product and modifier Fields

The reactant species, product species and modifier species in a reaction are described using the fields reactant, product and modifiers, respectively. These fields are optional lists of SpeciesReference and ModifierSpeciesReference structures, as shown in Figure 12 on the next page. They are described in more detail in Sections 4.9.5 and 4.9.6 below. The abstract type SimpleSpeciesReference is shown simply to demonstrate the common attribute, species, of the SpeciesReference and ModifierSpeciesReference structures. In future levels of SBML it is anticipated that SimpleSpeciesReference will have addition attributes.

4.9.3 The reversible Field

The reversible optional boolean field indicates whether the reaction is reversible. The field is optional, and if left unspecified in a model, it defaults to a value of "true". Although the reversibility of a reaction is determined by its rate law, the need to allow rate expressions in SBML to be optional leads to the need for a flag indicating reversibility. Information about reversibility in the absence of a KineticLaw in a Reaction is useful in certain kinds of structural analyses such as elementary mode analysis. It is true that the presence of this information in two places (i.e., the rate expression and the flag reversible) leaves open the possibility of a model containing contradictory information, but the creation of such a model would indicate an error on the part of the software generating it. Software developers must take care to ensure against logical contradictions in the definitions of reactions.

4.9.4 The fast Field

The optional boolean field fast is another boolean attribute in the Reaction data structure; a value of "true" signifies that the given reaction is a "fast" one compared to others in the system being modeled. This may be

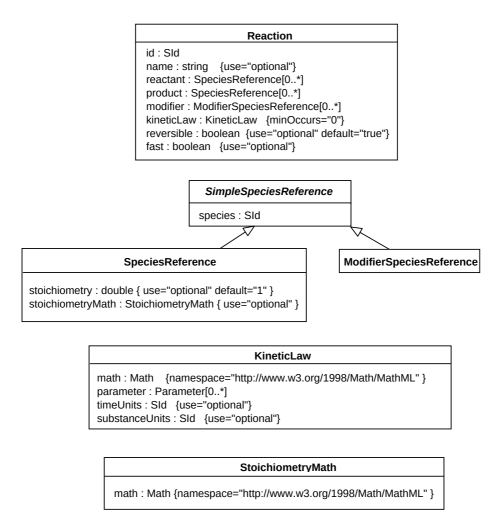


Figure 12: The definitions of Reaction, KineticLaw, SpeciesReference and ModiferSpeciesReference.

relevant when computing equilibrium concentrations of rapidly equilibrating reactions. Simulation/analysis packages may chose to use this information to reduce the number of ODEs required and thereby optimize such computations. (A simulator/analysis package that has no facilities for dealing with fast reactions can ignore this attribute. In theory, if the choice of which reactions are fast is correctly made, then a simulation performed with them should give the same results as a simulation performed without fast reactions. However, currently there appears to be no single unambiguous method for designating which reactions should be considered fast, and some users may designate a reaction as fast when in fact it is not.) The fast attribute does not have a default value: a missing value indicates that the modeller doesn't know or doesn't wish to specify the rate of the reaction relative to other reactions in the model.

4.9.5 SpeciesReference

Every species that enters into a given reaction must appear in that reaction's lists of reactants, products or modifiers. In an SBML model, all species that participate in any reaction are listed in the listOfSpecies field of the top-level Model data structure (see Section 4.2). Lists of products, reactants and modifiers in Reaction type structures do not introduce new species, but rather, they refer back to those listed in the model's listOfSpecies. For reactants and products, the connection is made using the SpeciesReference type data structure defined in Figure 12.

In SpeciesReference, the field species of type SId must refer to the name of an existing species defined in the enclosing Model-type structure. The stoichiometry for the product or reactant can be specified using either the stoichiometry or stoichiometryMath fields of the SpeciesReference structure. The stoichiometry field is of type double and defaults to 1. It is recommended for maximal interoperability that this field is used in preference to the stoichiometryMath field and should contain an integer value (see the following example). The stoichiometryMath field is implemented as an element containing a MathML math expression in dimensionless units. Only one of the stoichiometry and stoichiometryMath fields should be used on a given SpeciesReference structure.

The following is a simple example of a species reference for species "X0", with stoichiometry 2, in a list of reactants within a reaction named "J1":

The following is a more complex example of a species reference for species "X0", with a stoichiometry expression containing the parameter x:

A reaction can contain an empty list of reactants or an empty list of products but must have at least one reactant or product. Also note that whether a given species is allowed to appear as a reactant or product is dictated by certain flags on the structure describing the species in the Model; see Table 4 on page 20 for more information.

A species can occur more than once in the lists of reactants and products of a given reaction. Such a Reaction structure represents the combined effect of all the SpeciesReference structures that are contained in it. The effective stoichiometry for a species in a reaction is the sum of the stoichiometry values given on the SpeciesReference structures in the list of products minus the sum of stoichiometry values given on the SpeciesReference structures in the list of reactants. By convention a positive value indicates that the species is effectively a product and a negative value indicates that the species is effectively a reactant. SBML places no restrictions on the effective stoichiometry of a species in a reaction, for example, it can be zero. The following two SBML fragments containing reaction are exactly equivalent, having the same effective stoichiometry for all their species:

```
<reaction id="x">
         <listOfReactants>
             <speciesReference species="a"/>
              <speciesReference species="a"/>
              <speciesReference species="b"/>
         </listOfReactants>
         <listOfProducts>
             <speciesReference species="c"/>
              <speciesReference species="b"/>
         </listProducts>
     </reaction>
is equivalent to:
     <reaction id="x">
         <listOfReactants>
              <speciesReference species="a" stoichiometry="2"/>
         </listOfReactants>
         t0fProducts>
             <speciesReference species="c"/>
         </listProducts>
     </reaction>
```

4.9.6 ModifierSpeciesReference

In some cases, a species may act as a catalyst or inhibitor of a reaction and would not appear in the list of reactants or products because it is neither created nor destroyed in that reaction. In these cases, the species is known as a modifier. (That same species may still be a reactant or product of another reaction.)

The Reaction structure provides a way to express which species act as modifiers in a given reaction. This is the purpose of the modifier field in Reaction; this field is a list of ModifierSpeciesReference structures defined in Figure 12 on page 26. The ModifierSpeciesReference structure has only one field, species, of type SId; its value must be the identifier of a species defined in the enclosing Model.

4.9.7 KineticLaw

A kineticLaw structure, enclosed in a Reaction structure, describes the rate at which the reaction takes place. The inclusion of a KineticLaw structure in an instance of a Reaction component is optional; however, in general there is no useful default that can be substituted in place of a missing rate law definition in a reaction.

It is important to make clear at the outset that a "kinetic law" in SBML is not equivalent to a traditional rate law. The reason is that SBML must support multi-compartment models, and the units used in textbook rate laws as well as some conventional single-compartment modeling packages are problematic when used for defining reactions between multiple compartments. Rate expressions in SBML are expressed in terms of substance/time, rather than the more typical concentration/time. Converting between these two conventions is a simple matter of dividing the substance/time expression by the size of the compartment where a given species is located. To put this in slightly more precise terms, suppose that there are two species S and P in a reaction $S \to P$, where S is located in a compartment A having volume V_A , P is located in a compartment B having volume V_B , and the kinetic law expression gives the rate of the reaction as being B. Then the rate of change in the concentration of S is given by $d[S]_A/dt = -R/V_A$ and the rate of change in the concentration of P is $d[P]_B/dt = R/V_B$.

As stated in Section 3.6.3 the symbols for a species in a kinetic law formula represents the concentration of the species if the spatial dimensions of the species' compartment is non-zero and represents the substance amount of the species if the spatial dimensions of the species' compartment is 0.

As shown in Figure 12 on page 26, the KineticLaw structure has a field called math; this is a MathML expression that defines the rate of the reaction. The overall rate expression in math must have the units of substance/time. The optional fields substanceUnits and timeUnits determine the units of substance and time for the reaction. The values of these fields must be chosen from one of the following possibilities: one of the base unit names from Table 2 on page 14; one of the built-in unit names appearing in first column of

Table 3 on page 15); or the name of a new unit defined in the list of unit definitions in the enclosing Model structure. In the case of substanceUnits, the value chosen must be a scaled and/or multiplied variant of moles or item. In the case of timeUnits, the value chosen must be a scaled and/or multiplied variant of seconds. If these fields are not set in a given KineticLaw instance, the units are taken from the defaults defined by the built-in "substance" and "time" of Table 3 on page 15.

The expression in math may refer to species identifiers; in that case, as discussed in Section 3.6.3, the units of each species reference are those of *concentration*. The only species identifiers that can be used in math are those listed in the reactant, product and modifier fields of the Reaction structure.

An instance of a KineticLaw type structure can contain zero or more parameter structures (Section 4.7) that define symbols that can be used in the math element. As discussed in Section 3.5, reactions introduce local namespaces for parameter identifiers. Within a KineticLaw structure inside a reaction definition, a local parameter whose identifier is identical to a global parameter defined in the enclosing Model-type structure takes precedence over that global parameter.

The following is an example of a Reaction structure that defines a reaction named J_1 , in which $X_0 \to S_1$ at a rate given by kX_0S_2 , where S_2 is a catalyst and k is a parameter. It demonstrates the use of species references and the KineticLaw structure. The units on the species here are the defaults of substance/volume (see Section 4.6), and so the units of the parameter k are set appropriately to make the rate expression have final units of substance/time.

```
<model>
    <listOfUnitDefinitions>
        <unitDefinition id="vol_per_concent_per_time">
            tOfUnits>
                <unit kind="litre" exponent="2"/>
                 <unit kind="mole"</pre>
                                      exponent="-1"/>
                 <unit kind="second" exponent="-1"/>
            </listOfUnits>
        </unitDefinition>
    </listOfUnitDefinitions>
    <listOfSpecies>
        <species id="S1" compartment="c1" initialConcentration="2.0"/>
        <species id="S2" compartment="c1" initialConcentration="0.0"/>
<species id="E1" compartment="c1" initialConcentration="1.0"/>
    </listOfSpecies>
    <listOfReactions>
        <reaction id="J1">
            <listOfReactants>
                 <speciesReference species="S1"/>
            </listOfReactants>
            <listOfProducts>
                <speciesReference species="S2"/>
            </listOfProducts>
            <listOfModifiers>
                 <modifierSpeciesReference species="E1"/>
            </listOfModifiers>
            <kineticLaw>
                <math xmlns="http://www.w3.org/1998/Math/MathML">
                     <apply>
                         <times/>
                         <ci> k </ci>
                         <ci> S1 </ci>
                         <ci> E1 </ci>
                     </apply>
                 <parameter id="k" value="0.1" units="vol_per_concent_per_time"/>
                 </kineticLaw>
        </reaction>
```

```
</listOfReactions>
...
</model>
```

4.10 Events

Model has an optional list of Event structures that describe the time and form of explicit instantaneous discontinuous state changes in the model. For example, an event may describe that one species concentration is halved when another species concentration exceeds a given threshold value.

An Event structure defines when the event can occur, the variables that are affected by the event, and how the variables are affected. The effect of the event can optionally be delayed after the occurrence of the condition which invokes it. The operation of an Event structure is divided into two phases (even when the event is not delayed): one when the event is *fired* and the other when the event is *executed*. The Event type is defined in Figure 13. Both Event and EventAssignment are derived from SBase (see Section 3.1). An example of a model which uses events is given below.

id : SId { use="optional" } name : string { use="optional" } trigger : (math : Math {namespace="http://www.w3.org/1998/Math/MathML"}) delay : (math : Math {namespace="http://www.w3.org/1998/Math/MathML"}) timeUnits : SId { use="optional" } eventAssignment : EventAssignment[1..n]

```
EventAssignment

variable : SId

math : Math {namespace="http://www.w3.org/1998/Math/MathML"}
```

Figure 13: The definitions of Event and EventAssignment

4.10.1 The id and name Fields

These optional fields are available to enable to support external references to event structures. These fields operate in the manner described in Section 3.3 except that the id field is optional.

4.10.2 The trigger Field

The trigger field defines when the Event structure has an effect on the model. The trigger field contains a MathML boolean expression. The exact instant that the expression evaluates to true is the time point when the Event is *fired*. The event only fires when the trigger makes the transition from false to true. The event will fire at any further time points when the trigger make this transition.

4.10.3 The delay Field

The optional delay field defines the length of time after the event has *fired* that the event is *executed*. The delay field is another MathML expression. This expression should be evaluated when the rule is *fired*. The default value for the delay field is 0. The value of the delay field should always be positive.

4.10.4 The timeUnits Field

The optional field timeUnits determines the units of time that apply to the delay field. If not set, the units are taken from the defaults defined by the built-in "time" of Table 3 on page 15.

4.10.5 The eventAssignment Field

The eventAssignment field consists of a non-empty list of eventAssignment structures. This field is implemented as a listOfEventAssignments element containing one or more eventAssignment elements. The EventAssignment structures represent variable assignments which have effect when the event is executed. The Assignment structure is shown in Figure 13. The variable field is of type SId and contains the identifier of a variable i.e. a compartment, species or parameter. The structures referenced by the variable field must have their constant fields set to "false". The math field contains a MathML expression which defines the new value of the variable. This expression is evaluated when the Event is fired but the variable only acquires the result or new value when the Event is executed. The order of the EventAssignment structures is not significant (unlike assignment rules), the effect of one assignment cannot affect the result of another assignment. The identifiers occurring in the MathML ci fields of the EventAssignment structures represent the value of the identifier at the point when the Event is fired.

4.10.6 Example Event structure

A example of an Event structure follows. This structure makes the assignment $k_2 = 0$ at the point when $P_1 \leq t$:

```
<event>
    <trigger>
        <math xmlns="http://www.w3.org/1998/Math/MathML">
            <apply>
                 <leq/>
                 <ci> P1 </ci>
                 <ci> t </ci>
            </apply>
        </trigger>
    <listOfEventAssignments>
        <eventAssignment variable="k2">
             <math xmlns="http://www.w3.org/1998/Math/MathML">
                 \langle cn \rangle 0 \langle cn \rangle
            </eventAssignment>
    <listOfEventAssignments>
</event>
```

A complete example model that uses events is given in Section 5.9

4.10.7 Detailed semantics of events

The description of events above describes the action of events in isolation from each other. This section describes how events interact. Events whose trigger expression is true at the start of a simulation do not fire at the start of the simulation. Events fire only when the trigger becomes true i.e. the trigger expression transitions from false to true. It is possible for events to fire other events, i.e. an event assignment can cause an event to fire, therefore it is possible for model to be entirely encoded in Event structures.

It is entirely possible for two events to be *executed* simultaneously in simulated time. It is assumed that, although the precise time at which these events are *executed* is not resolved beyond the given point in simulated time, the order in which the events occur is resolved. This order can be significant in determining the overall outcome of a given simulation. SBML Level 2 does not define the algorithm for determining this order (the tie-breaking algorithm). As a result the results of simulations involving events may vary when simultaneous events occur during simulation. It is anticipated that future versions or levels of SBML will define a specific set of tie-breaking algorithms and a mechanism for models to indicate which algorithm should be applied during simulation.

Despite the absence of a specific tie-breaking algorithm SBML event simulation is constrained as follows. When an event X fires another event Y and event Y has zero delay then event Y is added to the existing set of simultaneous events that are pending execution. Events such as Y do not have a special priority or ordering within the tie-breaking algorithm. Events X and Y form a cascade of events at the same point in

simulation time. All events in a model are open to being in a cascade. The position of an event in the event list doesn't affect whether it can be in the cascade: Y can be triggered whether it is before or after X in the list of events. A cascade of events can be infinite (never terminate). When this occurs a simulator should indicate this has occurred i.e. it is incorrect for the simulator to arbitrarily break the cascade and continue the simulation without at least indicating the infinite cascade occurred. Given that a variable can change more than once when processing simultaneous events at simulation time t then the model behavior (output) for that variable at time t is the value of the variable at the end of processing all the simultaneous events at time t.

5 Example Models Expressed in XML Using SBML

In this section, we present several examples of complete models encoded in XML using SBML Level 2.

5.1 A Simple Example Application of SBML

Consider the following hypothetical branched system:

$$X_0 \xrightarrow{k_1 X_0} S_1 \xrightarrow{k_2 S_1} X_1$$

$$X_2$$

The following is an XML document that encodes the model shown above:

```
<?xml version="1.0" encoding="UTF-8"?>
<sbml xmlns="http://www.sbml.org/sbml/level2" level="2" version="1">
   <model id="Branch">
       <notes>
            <body xmlns="http://www.w3.org/1999/xhtml">
                Simple branch system.
                The reaction looks like this:
                                X0 -> S1; k1*X0;
S1 -> X1; k2*S1;
S1 -> X2; k3*S1;
                reaction-1:
                reaction-2:
                reaction-3:
            </body>
        </notes>
        <listOfCompartments>
            <compartment id="compartmentOne" size="1"/>
        </listOfCompartments>
        <listOfSpecies>
            <species id="S1" initialConcentration="0" compartment="compartmentOne"</pre>
                     boundaryCondition="false"/>
            <species id="X0" initialConcentration="0" compartment="compartmentOne"</pre>
                     boundaryCondition="true"/>
            <species id="X1" initialConcentration="0" compartment="compartmentOne"</pre>
                     boundaryCondition="true"/>
            <species id="X2" initialConcentration="0" compartment="compartmentOne"</pre>
                     boundaryCondition="true"/>
        </listOfSpecies>
        <listOfReactions>
            <reaction id="reaction_1" reversible="false">
                <listOfReactants>
                    <speciesReference species="X0"/>
                <listOfProducts>
                    <speciesReference species="S1"/>
                <kineticLaw>
                    <math xmlns="http://www.w3.org/1998/Math/MathML">
                        <apply>
                            <times/>
                            <ci> k1 </ci>
```

```
<ci> X0 </ci>
                        </apply>
                    <listOfParameters>
                        <parameter id="k1" value="0"/>
                    </listOfParameters>
                </kineticLaw>
            </reaction>
            <reaction id="reaction_2" reversible="false">
                <listOfReactants>
                    <speciesReference species="S1"/>
                </listOfReactants>
                <listOfProducts>
                    <speciesReference species="X1"/>
                </listOfProducts>
                <kineticLaw>
                    <math xmlns="http://www.w3.org/1998/Math/MathML">
                        <apply>
                            <times/>
                            <ci> k2 </ci>
                            <ci> S1 </ci>
                        </apply>
                    <listOfParameters>
                        <parameter id="k2" value="0"/>
                    </listOfParameters>
                </kineticLaw>
            </reaction>
            <reaction id="reaction_3" reversible="false">
                <listOfReactants>
                    <speciesReference species="S1"/>
                </listOfReactants>
                <listOfProducts>
                    <speciesReference species="X2"/>
                </listOfProducts>
                <kineticLaw>
                    <math xmlns="http://www.w3.org/1998/Math/MathML">
                        <apply>
                            <times/>
                            <ci> k3 </ci>
                            <ci> S1 </ci>
                        </apply>
                    <listOfParameters>
                        <parameter id="k3" value="0"/>
                    </listOfParameters>
                </kineticLaw>
            </reaction>
       </listOfReactions>
   </model>
</sbml>
```

In this example, the model has the identifier "Branch". The model contains one compartment, four species, and three reactions. The elements in the st0fReactants> and tofProducts> in each reaction refer to the names of elements listed in the tofSpecies>. The correspondences between the various elements is explicitly stated by the <speciesReference> elements.

The model also includes a <notes> annotation that summarizes the model in text form, with formatting encoded in XHTML. This may be useful for a software package that is able to read such annotations and, for example, render them in HTML in a graphical user interface.

5.2 Example Involving Units

The following model uses the units features of SBML Level 2. In this model, the default value of substance is changed in the list of unit definitions to be mole units with a scale factor of -3, or millimoles. This sets the default substance units in the model; components can override this scale locally. The size and time

built-ins are left to their defaults, ensuring that size is in liters and time is in seconds. The result is that, in this model, kinetic law formulas define rates in millimoles per second and the species symbols in them represent concentration values in millimoles per liter. All the species elements set the initial amount of every given species to 1 millimole. The parameters Vm and Km are defined to be in millimoles per liter per second, and milliMolar, respectively.

```
<?xml version="1.0" encoding="UTF-8"?>
<sbml xmlns="http://www.sbml.org/sbml/level2" level="2" version="1"</pre>
      xmlns:html="http://www.w3.org/1999/xhtml">
        <listOfUnitDefinitions>
             <unitDefinition id="substance">
                 tOfUnits>
                      <unit kind="mole" scale="-3"/>
                 </listOfUnits>
             </unitDefinition>
             <unitDefinition id="mmls">
                 stOfUnits>
                      <unit kind="mole" scale="-3"/>
                      <unit kind="litre" exponent="-1"/>
                      <unit kind="second" exponent="-1"/>
                 </listOfUnits>
             </unitDefinition>
             <unitDefinition id="mm">
                 stOfUnits>
                      <unit kind="mole" scale="-3"/>
                 </listOfUnits>
             </unitDefinition>
        </listOfUnitDefinitions>
        <listOfCompartments>
             <compartment id="cell"/>
        </listOfCompartments>
        tOfSpecies>
            <species id="x0" compartment="cell" initialConcentration="1"/>
<species id="x1" compartment="cell" initialConcentration="1"/>
<species id="s1" compartment="cell" initialConcentration="1"/>
             <species id="s2" compartment="cell" initialConcentration="1"/>
        </listOfSpecies>
        <listOfParameters>
             <parameter id="vm" value="2" units="mmls"/>
             <parameter id="km" value="2" units="mm"/>
        </listOfParameters>
        stOfReactions>
             <reaction id="v1">
                 <listOfReactants>
                      <speciesReference species="x0"/>
                 <listOfProducts>
                     <speciesReference species="s1"/>
                 </listOfProducts>
                 <kineticLaw>
                      <notes>
                          <html:p>((vm * s1)/(km + s1))*cell</html:p>
                      <math xmlns="http://www.w3.org/1998/Math/MathML">
                          <apply>
                              <times/>
                              <apply>
                                   <divide/>
                                   <apply>
                                       <times/>
                                       <ci> vm </ci>
                                       <ci> s1 </ci>
                                   </apply>
                                   <apply>
                                       <plus/>
                                       <ci> km </ci>
                                       <ci> s1 </ci>
```

```
</apply>
               </apply>
               <ci> cell </ci>
            </apply>
       </kineticLaw>
</reaction>
<reaction id="v2">
   <listOfReactants>
       <speciesReference species="s1"/>
   </listOfReactants>
   <listOfProducts>
       <speciesReference species="s2"/>
   </listOfProducts>
   <kineticLaw>
        <notes>
            <html:p>((vm * s2)/(km + s2))*cell</html:p>
       </notes>
       <math xmlns="http://www.w3.org/1998/Math/MathML">
           <apply>
               <times/>
                <apply>
                    <divide/>
                    <apply>
                       <times/>
                       <ci> vm </ci>
                       <ci> s2 </ci>
                    </apply>
                    <apply>
                        <plus/>
                       <ci> km </ci>
                       <ci> s2 </ci>
                   </apply>
               </apply>
               <ci>ci> cell </ci>
           </apply>
       </kineticLaw>
</reaction>
<reaction id="v3">
   <listOfReactants>
       <speciesReference species="s2"/>
   <listOfProducts>
       <speciesReference species="x1"/>
   </listOfProducts>
   <kineticLaw>
       <notes>
            <html:p>((vm * x1)/(km + x1))*cell</html:p>
       </notes>
        <math xmlns="http://www.w3.org/1998/Math/MathML">
            <apply>
               <times/>
                <apply>
                    <divide/>
                    <apply>
                        <times/>
                       <ci> vm </ci>
                       <ci> x1 </ci>
                    </apply>
                    <apply>
                       <plus/>
                       <ci> km </ci>
                       <ci> x1 </ci>
                   </apply>
                </apply>
               <ci> cell </ci>
           </apply>
```

5.3 Example Involving Assignment Rules

This section contains a model which simulates a system containing a fast reaction. This model uses rules to express the mathematics of the fast reaction explicitly rather than using the implicit fast field on a reaction element.

The system modeled is

$$X_{0} \xrightarrow{k_{1}X_{0}} S_{1}$$

$$S_{1} \xrightarrow{k_{f}S_{1} - k_{r}S_{2}} S_{2}$$

$$S_{2} \xrightarrow{k_{2}S_{2}} X_{1}$$

$$k_1 = 0.1$$
, $k_2 = 0.15$, $k_f = K_{eq} 10000$, $k_r = 10000$, $K_{eq} = 2.5$.

This can be approximated with the following system:

$$X_0 \xrightarrow{k_1 X_0} T$$

$$T \xrightarrow{k_2 S_2} X_1$$

$$S_1 = \frac{T}{1 + K_{eq}}, \quad S_2 = K_{eq}S_1$$

The following example SBML model encodes the approximate form.

```
<?xml version="1.0" encoding="UTF-8"?>
<sbml xmlns="http://www.sbml.org/sbml/level2" level="2" version="1"</pre>
       xmlns:math="http://www.w3.org/1998/Math/MathML">
          <listOfCompartments>
               <compartment id="cell"/>
          Species>
               <species id="X0" compartment="cell" initialConcentration="1"/>
<species id="X1" compartment="cell" initialConcentration="0"/>
<species id="T" compartment="cell" initialConcentration="0"/>
               <species id="S1" compartment="cell" initialConcentration="0"/>
<species id="S2" compartment="cell" initialConcentration="0"/>
          </listOfSpecies>
          <listOfParameters>
               <parameter id="Keq" value="2.5"/>
          </listOfParameters>
          <listOfRules>
               <assignmentRule variable="S1">
                    <math xmlns="http://www.w3.org/1998/Math/MathML">
                         <apply>
                              <divide/>
                              <ci> T </ci>
                              <apply>
                                    <plus/>
                                   <cn> 1 </cn>
                                   <ci> Keq </ci>
                              </apply>
                         </apply>
```

```
</assignmentRule>
           <assignmentRule variable="S2">
               <math xmlns="http://www.w3.org/1998/Math/MathML">
                   <apply>
                       <times/>
                       <ci> Keq </ci>
                       <ci> S1 </ci>
                   </apply>
               </assignmentRule>
       </listOfRules>
       <listOfReactions>
           <reaction id="in">
               <listOfReactants>
                   <speciesReference species="X0"/>
               <listOfProducts>
                   <speciesReference species="T"/>
               </listOfProducts>
               <kineticLaw>
                   <math xmlns="http://www.w3.org/1998/Math/MathML">
                       <apply>
                           <times/>
                          <ci> k1 </ci>
                          <ci> X0 </ci>
                      </apply>
                   <listOfParameters>
                       <parameter id="k1" value="0.1"/>
                   </listOfParameters>
               </kineticLaw>
           </reaction>
           <reaction id="out">
               <listOfReactants>
                   <speciesReference species="T"/>
               <listOfProducts>
                   <speciesReference species="X1"/>
               </listOfProducts>
               <kineticLaw>
                   <math xmlns="http://www.w3.org/1998/Math/MathML">
                       <apply>
                           <times/>
                          <ci> k2 </ci>
                          <ci> S2 </ci>
                       </apply>
                   <listOfParameters>
                       <parameter id="k2" value="0.15"/>
                   </listOfParameters>
               </kineticLaw>
           </reaction>
       </listOfReactions>
   </model>
</sbml>
```

5.4 Example Involving Algebraic Rules

This section contains an example model which contains an AlgebraicRule structure. The model contains a different formulation of the fast reaction described in Section 5.3.

The system described in Section 5.3 can be approximated with the following system:

$$X_0 \xrightarrow{k_1 X_0} T$$

$$T \xrightarrow{k_2 S_1} X_1$$

$$S_2 = K_{eq} S_1$$

with the constraint:

$$S_1 + S_2 - T = 0$$

The following example SBML model encodes this approximate form.

```
<?xml version="1.0" encoding="UTF-8"?>
<sbml xmlns="http://www.sbml.org/sbml/level2" level="2" version="1">
    <model>
         <listOfCompartments>
              <compartment id="cell"/>
         </listOfCompartments>
         <listOfSpecies>
             <species id="X0" compartment="cell" initialConcentration="1"/>
<species id="X1" compartment="cell" initialConcentration="0"/>
<species id="T" compartment="cell" initialConcentration="0"/>
             <species id="S1" compartment="cell" initialConcentration="0"/>
<species id="S2" compartment="cell" initialConcentration="0"/>
         </listOfSpecies>
         <listOfParameters>
              <parameter id="Keq" value="2.5"/>
         </listOfParameters>
         tOfRules>
              <assignmentRule variable="S2">
                  <math xmlns="http://www.w3.org/1998/Math/MathML">
                       <apply>
                            <times/>
                            <ci> Keq </ci>
                            <ci> S1 </ci>
                       </apply>
                  </assignmentRule>
              <algebraicRule>
                   <math xmlns="http://www.w3.org/1998/Math/MathML">
                       <apply>
                            <minus/>
                            <apply>
                                 <plus/>
                                 <ci> S2 </ci>
                                 <ci> S1 </ci>
                            </apply>
                            <ci> T </ci>
                       </apply>
                   </algebraicRule>
         </listOfRules>
         <listOfReactions>
              <reaction id="in">
                  <listOfReactants>
                       <speciesReference species="X0"/>
                   </listOfReactants>
                  <listOfProducts>
                       <speciesReference species="T"/>
                  </listOfProducts>
                   <kineticLaw>
                       <math xmlns="http://www.w3.org/1998/Math/MathML">
                            <apply>
                                 <times/>
```

```
<ci> k1 </ci>
                            <ci> X0 </ci>
                        </apply>
                    <listOfParameters>
                        <parameter id="k1" value="0.1"/>
                    </listOfParameters>
                </kineticLaw>
           </reaction>
           <reaction id="out">
                <listOfReactants>
                    <speciesReference species="T"/>
                </listOfReactants>
                <listOfProducts>
                    <speciesReference species="X1"/>
                </listOfProducts>
                <kineticLaw>
                    <math xmlns="http://www.w3.org/1998/Math/MathML">
                        <apply>
                            <times/>
                            <ci> k2 </ci>
                            <ci> S2 </ci>
                        </apply>
                    <listOfParameters>
                        <parameter id="k2" value="0.15"/>
                    </listOfParameters>
                </kineticLaw>
            </reaction>
       </listOfReactions>
   </model>
</sbml>
```

5.5 Example Involving Combinations of boundaryCondition and constant Values on Species with RateRule structures

This section contains a model which includes four species each with a different combination of values of for the boundaryCondition and constant attributes.

Consider the following hypothetical system:

$$S_1 + S_2 \quad k_1 S_1 S_2 S_3 \quad S_4$$

where values over time are determined by:

$$\frac{dS_1}{dt} = k_2$$

$$S_2 = 1$$

$$S_3 = 2$$

$$k_1 = 0.5$$

$$k_2 = 0.1$$

and initial values are:

$$S_1 = 0$$

$$S_4 = 0$$

 S_1 and S_2 are reactants but their values are not determined by a kinetic law thus they are both on the boundary of the reaction system and in SBML have boundaryCondition attribute values of true. The

value of S_1 varies over time so in SBML S_1 has a constant attribute with a default value of "false". The values of S_2 and S_3 are fixed so in SBML they have a constant attribute values of "true". S_3 only occurs as a modifier so the value of its boundaryCondition attribute can default to false. S_4 is a product whose value is determined by a kinetic law and therefore in the SBML representation has false values, the default values, for both boundaryCondition and constant attributes.

The following is the XML document that encodes the model shown above:

```
<?xml version="1.0" encoding="UTF-8"?>
<sbml xmlns="http://www.sbml.org/sbml/level2" level="2" version="1">
   <model id="BoundaryCondExampleModel">
       <listOfCompartments>
           <compartment id="compartmentOne" size="1"/>
       <listOfSpecies>
           <species id="S1" initialConcentration="0" compartment="compartmentOne"</pre>
               boundaryCondition="true" />
           <species id="S2" initialConcentration="1" compartment="compartmentOne"</pre>
               boundaryCondition="true" constant="true" />
           <species id="S3" initialConcentration="3" compartment="compartmentOne"</pre>
               constant="true"/>
           <species id="S4" initialConcentration="0" compartment="compartmentOne"/>
       </listOfSpecies>
       <parameter id="k1" value="0.5"/>
           </listOfParameters>
       tOfRules>
           <rateRule variable="S1">
               <math xmlns="http://www.w3.org/1998/Math/MathML">
                   <apply>
                      <ci> k2 </ci>
                   </apply>
               </rateRule>
       </listOfRules>
       <listOfReactions>
           <reaction id="reaction_1" reversible="false">
               <listOfReactants>
                   <speciesReference species="S1"/>
                   <speciesReference species="S2"/>
               <listOfProducts>
                   <speciesReference species="S4"/>
               </listOfProducts>
               <listOfModifiers>
                   <modifierSpeciesReference species="S3"/>
               </listOfModifiers>
               <kineticLaw>
                   <math xmlns="http://www.w3.org/1998/Math/MathML">
                       <apply>
                           <times/>
                           <ci> k1 </ci>
                           <ci> S1 </ci>
                           <ci> S2 </ci>
                           <ci> S3 </ci>
                        </apply>
                   </kineticLaw>
           </reaction>
       </listOfReactions>
   </model>
</sbml>
```

5.6 Example of translation from a multi-compartmental model containing reactions and rate rules to ODEs

This section contains a model with 2 compartments, reactions and rate rules. The rate rules are for a compartment and a species. The model is followed by its complete translation into ODEs. The model is shown in figure 14.

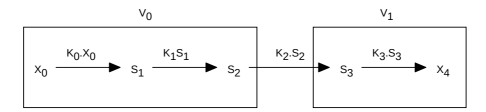


Figure 14: A example multi-compartmental model

Reaction equations are in substance per time units. Species variables are in concentrations. Species X_0 and X_4 are boundary conditions. The volume of compartment V_1 and the concentration of X_0 vary according to rules.

The SBML for this model is

```
<?xml version="1.0" encoding="UTF-8"?>
<sbml xmlns="http://www.sbml.org/sbml/level2" level="2" version="1">
    <model id="ODEExampleModel">
        <listOfCompartments>
            <compartment id="V0" size="10"/>
            <compartment id="V1" size="1" constant="false"/>
        </listOfCompartments>
        Species>
            <species id="X0" initialConcentration="0" compartment="V0"</pre>
                boundaryCondition="true" />
            <species id="S1" initialConcentration="0" compartment="V0"/>
<species id="S2" initialConcentration="0" compartment="V0"/>
            <species id="S3" initialConcentration="0" compartment="V1"/>
            <species id="X4" initialConcentration="0" compartment="V1"</pre>
                boundaryCondition="true" constant="true"/>
        </listOfSpecies>
        <listOfParameters>
            <parameter id="K0" value="0.1"/>
            <parameter id="K1" value="0.5"/>
            <parameter id="K2" value="0.1"/>
            <parameter id="K3" value="0.5"/>
            <parameter id="Kv" value="0.5"/>
            <parameter id="Kin" value="0.1"/>
        </listOfParameters>
        stOfRules>
            <rateRule variable="X0">
                 <math xmlns="http://www.w3.org/1998/Math/MathML">
                     <apply>
                         <ci> Kin </ci>
                     </apply>
                 </rateRule>
            <rateRule variable="V1">
                 <math xmlns="http://www.w3.org/1998/Math/MathML">
                     <apply>
                         <ci> Kv </ci>
                     </apply>
                 </rateRule>
        </listOfRules>
```

```
<listOfReactions>
   <reaction id="reaction_1" reversible="false">
       <listOfReactants>
           <speciesReference species="X0"/>
       <listOfProducts>
           <speciesReference species="S1"/>
       </listOfProducts>
       <kineticLaw>
           <math xmlns="http://www.w3.org/1998/Math/MathML">
               <apply>
                   <times/>
                   <ci> KO </ci>
                   <ci> X0 </ci>
                </apply>
           </kineticLaw>
   </reaction>
   <reaction id="reaction_2" reversible="false">
       <listOfReactants>
           <speciesReference species="S1"/>
       </listOfReactants>
       <listOfProducts>
           <speciesReference species="S2"/>
       </listOfProducts>
       <kineticLaw>
           <math xmlns="http://www.w3.org/1998/Math/MathML">
               <apply>
                   <times/>
                   <ci> K1 </ci>
                   <ci> S1 </ci>
                </apply>
           </kineticLaw>
   </reaction>
   <reaction id="reaction_3" reversible="false">
       <listOfReactants>
           <speciesReference species="S2"/>
       </listOfReactants>
       <listOfProducts>
           <speciesReference species="S3"/>
       </listOfProducts>
       <kineticLaw>
           <math xmlns="http://www.w3.org/1998/Math/MathML">
               <apply>
                   <times/>
                   <ci> K2 </ci>
                   <ci> S2 </ci>
                </apply>
           </kineticLaw>
   </reaction>
   <reaction id="reaction_4" reversible="false">
       <listOfReactants>
           <speciesReference species="S3"/>
       </listOfReactants>
       <listOfProducts>
           <speciesReference species="X4"/>
       <kineticLaw>
           <math xmlns="http://www.w3.org/1998/Math/MathML">
               <apply>
                   <times/>
                   <ci> K3 </ci> <ci> S3 </ci>
                </apply>
           </kineticLaw>
   </reaction>
```

The ODE translation of this model is as follows assuming all species variables contain the species concentration:

Parameters and Constants:

$$X_4 = 0$$

 $V_0 = 10$
 $K_0 = 0.1$
 $K_1 = 0.5$
 $K_2 = 0.1$
 $K_3 = 0.5$
 $K_{in} = 0.1$
 $K_v = 0.5$

Initial Conditions of Variables:

$$V_1 = 1$$

 $X_0 = S_1 = S_2 = S_3 = 0$

Differential Equations:

$$\begin{array}{ll} \frac{dV_1}{dt} = K_v & \text{rule} \\ \\ \frac{dX_0}{dt} = K_{in} & \text{rule} \\ \\ \frac{dS_1}{dt} = \frac{K_0 X_0 - K_1 S_1}{V_0} & \text{reactions 1 and 2} \\ \\ \frac{dS_2}{dt} = \frac{K_1 S_1 - K_2 S_2}{V_0} & \text{reactions 2 and 3} \\ \\ \frac{dS_3}{dt} = \frac{K_2 S_2 - K_3 S_3}{V_1} & \text{reactions 3 and 4} \end{array}$$

5.7 Example Involving Function Definitions

This section contains a model which uses the function definition feature of SBML. Consider the following hypothetical system:

$$S_1 \xrightarrow{f(S_1)} S_2$$

where

$$f(x) = x * 2$$

The following is the XML document that encodes the model shown above:

```
<listOfFunctionDefinitions>
           <functionDefinition id="f">
                <math xmlns="http://www.w3.org/1998/Math/MathML">
                   <lambda>
                        <br/>
<br/>
dvar><ci> x </ci></bvar>
                        <apply>
                            <times/>
                           <ci> x </ci>
                           <cn> 2 </cn>
                        </apply>
                    </lambda>
               </functionDefinition>
        </listOfFunctionDefinitions>
        <listOfCompartments>
            <compartment id="compartmentOne" size="1"/>
        </listOfCompartments>
        <listOfSpecies>
           <species id="S1" initialConcentration="0" compartment="compartmentOne"/>
           <species id="S2" initialConcentration="0" compartment="compartmentOne"/>
        </listOfSpecies>
        <listOfReactions>
            <reaction id="reaction_1" reversible="false">
                <listOfReactants>
                    <speciesReference species="S1"/>
                <listOfProducts>
                    <speciesReference species="S2"/>
               </listOfProducts>
               <kineticLaw>
                    <math xmlns="http://www.w3.org/1998/Math/MathML">
                        <apply>
                           <ci> f </ci>
                           <ci> S1 </ci>
                         </apply>
                   </kineticLaw>
           </reaction>
        </listOfReactions>
   </model>
</sbml>
```

5.8 Example Involving delay Functions

The following is a simple model illustrating the use of *delay* to represent a gene that suppresses its own expression. The model can be expressed in a single rule:

$$\frac{dP}{dt} = \frac{\frac{1}{1 + m(P_{delayed})^q} - P}{\tau}$$

where

 $P_{delayed}$ is $delay(P, \Delta_t)$ or P at $t - \Delta_t$ P is protein concentration τ is the response time m is a multiplier or equilibrium constant q is the Hill coefficient

The SBML form of this model is as follows:

```
</or>
        Species>
           <species id="P" compartment="cell" initialConcentration="0"/>
        </listOfSpecies>
        <listOfParameters>
            <parameter id="tau" value="1"/>
           <parameter id="m" value="0.5"/>
           <parameter id="q" value="1"/>
           <parameter id="delta_t" value="1"/>
        </listOfParameters>
        tOfRules>
           <rateRule variable="P">
                <math xmlns="http://www.w3.org/1998/Math/MathML">
                <apply>
                 <divide/>
                  <apply>
                  <minus/>
                   <apply>
                   <divide/>
                   <cn> 1 </cn>
                    <apply>
                     <plus/>
                     <cn> 1 </cn>
                     <apply>
                      <times/>
                      <ci> m </ci>
                      <apply>
                      <power/>
                      <apply>
                       <csymbol encoding="text"</pre>
                                definitionURL="http://www.sbml.org/symbols/delay">
                           delay
                        </csymbol>
                        <ci> P </ci>
                       <ci> delta_t </ci>
                      </apply>
                      <ci>q </ci>
                      </apply>
                    </apply>
                   </apply>
                   </apply>
                  <ci>P </ci>
                  </apply>
                 <ci> tau </ci>
                </apply>
                </rateRule>
       </listOfRules>
   </model>
</sbml>
```

5.9 Example Involving Events

This section presents a simple model system that demonstrates the use of events in SBML. Consider a system with two genes, k_1 and k_2 . k_1 is initially on and k_2 is initially off. When turned on, the two genes lead to the production of two products, P_1 and P_2 , respectively, at a fixed rate. When P_1 reaches a given concentration, k_2 switches off. This system can be represented mathematically as follows:

$$\frac{dP_1}{dt} = k_1 - P_1$$

$$\frac{dP_2}{dt} = k_2 - P_2$$

$$k_2 = \begin{cases} 0 & \text{when } P_1 \le \tau, \\ 1 & \text{when } P_1 > \tau. \end{cases}$$

```
k_1 = 1 k_2 = 0 \tau = 0.25 P_1 = 0 P_2 = 0
```

The SBML Level 2 representation of this as follows:

```
<?xml version="1.0" encoding="UTF-8"?>
<sbml xmlns="http://www.sbml.org/sbml/level2" level="2" version="1"</pre>
     xmlns:math="http://www.w3.org/1998/Math/MathML">
   <model>
       <listOfCompartments>
            <compartment id="cell"/>
       <listOfSpecies>
            <species id="P1" compartment="cell" initialConcentration="0"/>
            <species id="P2" compartment="cell" initialConcentration="0"/>
        </listOfSpecies>
        <listOfParameters>
            <parameter id="tau" value="0.25"/>
        </listOfParameters>
        tOfRules>
            <rateRule variable="P1">
                <math:math>
                    <math:apply>
                        <math:minus/>
                        <math:ci> k1 </math:ci> <math:ci> P1 </math:ci>
                    </math:apply>
                </math:math>
            </rateRule>
            <rateRule variable="P2">
                <math:math>
                    <math:apply>
                        <math:minus/>
                        <math:ci> k2 </math:ci>
<math:ci> P2 </math:ci>
                    </math:apply>
                </math:math>
            </rateRule>
        </listOfRules>
        <listOfEvents>
            <event>
                <trigger>
                    <math:math>
                        <math:apply>
                            <math:gt/>
                            <math:ci> P1 </math:ci>
                            <math:ci> tau </math:ci>
                        </math:apply>
                    </math:math>
                </trigger>
                <listOfEventAssignments>
                    <eventAssignment variable="k2">
                        <math:math>
                            <math:cn> 1 </math:cn>
                        </math:math>
                    </eventAssignment>
                </listOfEventAssignments>
            </event>
            <event>
                <trigger>
                    <math:math>
                        <math:apply>
                            <math:leq/>
                            <math:ci> P1 </math:ci>
                            <math:ci> tau </math:ci>
                        </math:apply>
```

5.10 Example Involving Two-Dimensional Compartments

The following example is a model that uses a two-dimensional compartment. It is a fragment of a larger model of calcium regulation across the plasma membrane of a cell. The model includes a calcium influx channel, Ca_channel, and a calcium-extruding PMCA pump, Ca_Pump. The model also includes two cytosolic proteins that buffer calcium via the CalciumCalbindin_gt_BoundCytosol and CalciumBuffer_gt_BoundCytosol reactions.

```
<?xml version="1.0" encoding="UTF-8"?>
<sbml xmlns="http://www.sbml.org/sbml/level2" level="2" version="1">
    <model id="facilitated_ca_diffusion">
        <listOfUnitDefinitions>
            <unitDefinition id="substance">
                <listOfUnits>
                     <unit kind="mole" scale="-6"/>
                </listOfUnits>
            </unitDefinition>
            <unitDefinition id="area">
                <listOfUnits>
                     <unit kind="metre" scale="-6" exponent="2" />
                </listOfUnits>
            </unitDefinition>
        <listOfCompartments>
            <compartment id="Extracellular" spatialDimensions="3"/>
<compartment id="PlasmaMembrane" outside="Extracellular" spatialDimensions="2"/>
            <compartment id="Cytosol" outside="PlasmaMembrane" spatialDimensions="3"/>

</listOfCompartments>
        Species>
            <species
              id="CaBPB_C"
              compartment="Cytosol"
              initialConcentration="47.17"/>
            <species
              id="B_C"
              compartment="Cytosol"
              initialConcentration="396.04"/>
            <species
              id="CaB_C"
              compartment="Cytosol"
              initialConcentration="3.96"/>
            <species
              id="Ca_EC"
              compartment="Extracellular"
              initialConcentration="1000"/>
            <species
              id="Ca_C"
              compartment="Cytosol"
              initialConcentration="0.1"/>
            <species</pre>
              id="CaCh_PM"
              compartment="PlasmaMembrane"
              initialConcentration="1"/>
```

```
<species
     id="CaPump_PM"
     compartment="PlasmaMembrane"
     initialConcentration="1"/>
   <species</pre>
     id="CaBP_C"
     compartment="Cytosol"
     initialConcentration="202.83"/>
</listOfSpecies>
<listOfReactions>
   <reaction id="CalciumCalbindin_gt_BoundCytosol" fast="true">
       <listOfReactants>
           <speciesReference species="CaBP_C"/>
           <speciesReference species="Ca_C"/>
       <listOfProducts>
           <speciesReference species="CaBPB_C"/>
       </listOfProducts>
       <kineticLaw>
                  <notes>
                  (((Kf_CalciumCalbindin_BoundCytosol * CaBP_C) * Ca_C) -
                        (Kr_CalciumCalbindin_BoundCytosol * CaBPB_C))
                 </notes>
           <math xmlns="http://www.w3.org/1998/Math/MathML">
               <apply>
                   <minus/>
                   <apply>
                      <times/>
                      <ci> Kf_CalciumCalbindin_BoundCytosol </ci>
                      <ci> CaBP_C </ci>
                      <ci> Ca_C </ci>
                  </apply>
                   <apply>
                      <times/>
                      <ci> Kr_CalciumCalbindin_BoundCytosol </ci>
                      <ci> CaBPB_C </ci>
                   </apply>
               </apply>
           <listOfParameters>
                   <parameter id="Kf_CalciumCalbindin_BoundCytosol" value="20.0"/>
                   <parameter id="Kr_CalciumCalbindin_BoundCytosol" value="8.6"/>
                </listOfParameters>
       </kineticLaw>
   </reaction>
   <reaction id="CalciumBuffer_gt_BoundCytosol" fast="true">
       <listOfReactants>
           <speciesReference species="Ca_C"/>
           <speciesReference species="B_C"/>
       </listOfReactants>
       t0fProducts>
           <speciesReference species="CaB_C"/>
       </listOfProducts>
       <kineticLaw>
             (((Kf_CalciumBuffer_BoundCytosol * Ca_C) * B_C) -
                   (Kr_CalciumBuffer_BoundCytosol * CaB_C))
             </notes>
           <math xmlns="http://www.w3.org/1998/Math/MathML">
               <apply>
                  <minus/>
                   <apply>
                      <times/>
                      <ci> Kf_CalciumBuffer_BoundCytosol </ci>
                      <ci> Ca_C </ci>
```

```
<ci> B_C </ci>
               </apply>
               <apply>
                   <times/>
                   <ci> Kr_CalciumBuffer_BoundCytosol </ci>
                   <ci> CaB_C </ci>
               </apply>
           </apply>
       <listOfParameters>
           <parameter id="Kf_CalciumBuffer_BoundCytosol" value="0.1"/>
<parameter id="Kr_CalciumBuffer_BoundCytosol" value="1.0"/>
       </listOfParameters>
   </kineticLaw>
</reaction>
<reaction id="Ca_Pump">
   <listOfReactants>
       <speciesReference species="Ca_C"/>
    </listOfReactants>
   <listOfProducts>
       <speciesReference species="Ca_EC"/>
    </listOfProducts>
   <listOfModifiers>
       <modifierSpeciesReference species="CaPump_PM"/>
   </listOfModifiers>
    <kineticLaw>
       <notes>
         CaPump_PM)
         </notes>
       <math xmlns="http://www.w3.org/1998/Math/MathML">
           <apply>
               <divide/>
               <apply>
                   <times/>
                   <ci> Vmax </ci>
                   <ci> kP </ci>
                   <ci> CaPump_PM </ci>
                   <apply>
                       <minus/>
                       <ci> Ca_C </ci>
                       <ci> Ca_Rest </ci>
                   </apply>
               </apply>
               <apply>
                   <times/>
                   <apply>
                       <plus/>
                       <ci>Ca_C </ci>
                       <ci> kP </ci>
                   </apply>
                   <apply>
                       <plus/>
                       <ci>Ca_Rest </ci>
                       <ci> kP </ci>
                   </apply>
               </apply>
           </apply>
       <listOfParameters>
           <parameter id="Vmax" value="-4000"/>
           <parameter id="kP" value="0.25"/>
           <parameter id="Ca_Rest" value="0.1"/>
       </listOfParameters>
   </kineticLaw>
</reaction>
<reaction id="Ca_channel">
```

```
<listOfReactants>
                  <speciesReference species="Ca_EC"/>
               </listOfReactants>
               t0fProducts>
                  <speciesReference species="Ca_C"/>
               </listOfProducts>
               <listOfModifiers>
                  <modifierSpeciesReference species="CaCh_PM"/>
               </listOfModifiers>
               <kineticLaw>
                  <notes>
                     (J0 * Kc * (Ca_EC - Ca_C) / (Kc + Ca_C) * CaCh_PM)
                    </notes>
                  <math xmlns="http://www.w3.org/1998/Math/MathML">
                      <apply>
                          <divide/>
                          <apply>
                              <times/>
                              <ci> CaCh_PM </ci>
                              <ci> JO </ci>
                              <ci> Kc </ci>
                              <apply>
                                  <minus/>
                                  <ci> Ca_EC </ci>
                                  <ci> Ca_C </ci>
                              </apply>
                          </apply>
                          <apply>
                              <plus/>
                              <ci> Kc </ci>
                              <ci> Ca_C </ci>
                          </apply>
                      </apply>
                  <listOfParameters>
                      <parameter id="J0" value="0.014"/>
                      <parameter id="Kc" value="0.5"/>
                  </listOfParameters>
               </kineticLaw>
           </reaction>
       </model>
</sbml>
```

6 Discussion

The volume of data now emerging from molecular biotechnology leave little doubt that extensive computer-based modeling, simulation and analysis will be critical to understanding and interpreting the data (Abbott, 1999; Gilman, 2000; Popel and Winslow, 1998; Smaglik, 2000). This has lead to an explosion in the development of computer tools by many research groups across the world. The explosive rate of progress is exciting, but the rapid growth of the field is accompanied by problems and pressing needs.

One problem is that simulation models and results often cannot be directly compared, shared or re-used, because the tools developed by different groups often are not compatible with each other. As the field of systems biology matures, researchers increasingly need to communicate their results as computational models rather than box-and-arrow diagrams. They also need to reuse published and curated models as library components in order to succeed with large-scale efforts (e.g., the Alliance for Cellular Signaling; Gilman, 2000; Smaglik, 2000). These needs require that models implemented in one software package be portable to other software packages, to maximize public understanding and to allow building up libraries of curated computational models.

We offer SBML to the systems biology community as a suggested format for exchanging models between simulation/analysis tools. SBML is an open model representation language oriented specifically towards representing systems of biochemical reactions.

Our vision for SBML is to create an open standard that will enable different software tools to exchange computational models. SBML is not static; we continue to develop and experiment with it, and we interact with other groups who seek to develop similar markup languages. We plan on continuing to evolve SBML with the help of the systems biology community to make SBML increasingly more powerful, flexible and useful.

6.1 Future Enhancements: SBML Level 3 and Beyond

As mentioned above, SBML Level 2 is intended to provide a foundation for modeling biochemical networks. Many people have expressed a desire to see additional capabilities added to SBML. The following summarizes additional features that are under consideration to be included in SBML Level 3:

- Arrays. This will enable the creation of arrays of components (species, reactions, compartments and submodels).
- Connections. This will be a mechanism for describing the connections between items in an array. For example, it should be possible to create a 2-D array of compartments and then a 4-D array of reactions which transport species between the compartments, where the third dimension is the connections between the compartments. Two possible ways of describing a connection scheme are: (1) sparse/explicit, simply listing the relative co-coordinates of connected objects for patterns of points; (2) algebraic, where a conditional equation describes whether two objects are connected.
- Geometry. This will enable the encoding of the spatial characteristics of models including the geometry of compartments, the diffusion properties of species and the specification of different species concentrations across different regions of a cell.
- *Model Composition*. This will enable a large model to be built up out of instances of other models. It will also allow the reuse of model components and the creation of several instances of the same model.
- Multi-state and Complex Species. This will allow the straight-forward construction of models involving species with a large number of states or species composed of subcomponents. The representation scheme would be designed to contain the combinatorial explosion of objects that often results from these types of models.
- Controlled Vocabularies. This will enable models and model components to be labelled with instances of controlled vocabularies. The model vocabulary will describe the SBML features used by the model.
- *Diagrams*. This feature will allow components to be annotated with data to enable the display of the model in a diagram.
- Dynamic Structure. This will enable model structure to vary during simulation. One aspect of aspect of this allowing rules and reactions to have their effect conditional on the state of the model system. For example in SBML Level 2 it is possible to create a rule with the effect:

$$\frac{ds}{dt} = \begin{cases} 0 & \text{if } s > 0\\ y & \text{otherwise} \end{cases}$$

Dynamic restructuring would enable the expression of the following example:

if
$$s > 0$$
 $\frac{ds}{dt} = y$

where s is not determined by the rule when $s \leq 0$.

- Rules for Initial Conditions. This will enable the encoding of rules that are only executed at the start of a simulation i.e. set initial conditions. These rules will be similar to assignment rules and are evaluated in the same sequence as assignment rules but only in the very first instance of simulation.
- *Tie-breaking algorithm*. This will include a controlled vocabulary and associated attributes on models to indicate the simultaneous event tie-breaking algorithm required to correctly simulate the model.

6.2 Relationships to Other Efforts

There are a number of ongoing efforts with similar goals as those of SBML. Many of them are oriented more specifically toward describing protein sequences, genes and related entities for database storage and search. These are generally not intended to be computational models, in the sense that they do not describe entities and behavioral rules in such a way that a simulation package could "run" the models.

The effort perhaps closest in spirit to SBML is CellMLTM (Hedley et al., 2001). CellML is an XML-based markup language designed for storing and exchanging computer-based biological models. It includes facilities for representing model structure, mathematics and additional information for database storage and search. Models are described in terms of networks of connections between discrete components, where a component is a functional unit that may correspond to a physical compartment or simply a convenient modeling abstraction. Components contain variables and connections contain mappings between the variables of connected components. CellML provides facilities for grouping components and specifying the kinds of relationships that may exist between components. It also uses MathML (W3C, 2000b) for expressing mathematical relationships between components and provides the ability to use ECMAScript (formerly known as JavaScript) to define functions.

The constructs in CellML tend to be at a more abstract and general level than those in SBML Level 2, and describe the structure and underlying mathematics of cellular models in a very general way. By contrast, SBML is closer to the internal object model used in model analysis software. Because SBML Level 2 is being developed in the context of interacting with a number of existing simulation packages, it is a more concrete language than CellML and may be better suited to its purpose of enabling interoperability with existing simulation tools.

The development of SBML Level 2 has benefited from discussions with the developers of CellML. The developers of SBML and CellML are actively engaged in ensuring that the two representations can be translated between each other.

Acknowledgments

SBML was developed with funding and support from the ERATO Kitano Symbiotic Systems project, a project funded by the Japan Science and Technology Corporation and hosted in part at the California Institute of Technology.

SBML was first conceived at the JST/ERATO-sponsored First Workshop on Software Platforms for Systems Biology, held in April, 2000, at the California Institute of Technology in Pasadena, California, USA. The participants collectively decided to begin developing a common XML-based declarative language for representing models. A draft version of the Systems Biology Markup Language was developed by the Caltech ERATO team and delivered to all collaborators in August, 2000. This draft version underwent extensive discussion over mailing lists and then again during the Second Workshop on Software Platforms for Systems Biology held in Tokyo, Japan, November 2000. A revised version of SBML was issued by the Caltech ERATO team in December, 2000, and after further discussions over mailing lists and in meetings, we produced a description of SBML Level 1 (Hucka et al., 2001).

SBML Level 2 was conceived at the 5th Workshop on Software Platforms for Systems Biology, held in July 2002, at the University of Hertfordshire, UK. The participants collectively decided to revise the form of SBML in Level 2. The first draft of this document was released in August 2002. The final set of features in SBML Level 2 described in this document was finalized in May 2003 at the 7th Workshop on Software Platforms for Systems Biology in Ft. Lauderdale, Florida.

SBML Level 2 was developed with the help of many people, especially the authors of BioSketchPad, BioSpice, CellDesigner, Cellerator, CellML, COPASI, DBSolve, E-Cell, ESS, Gepasi, Jarnac, JDesigner, JigCell, MCell, NetBuilder, Promot/DIVA, StochSim, and Virtual Cell, and members of the sysbio and sbml-discuss mailing lists. We are particularly grateful to the following people for discussions, advice and comments: Nicolas Allen, Adam Arkin, Hamid Bolouri, Ben Bornstein, Dennis Bray, Roger Brent, Steve Burbeck, Claudine Chaouiya, Kwang Cho, Athel Cornish-Bowden, Manuel Corpas, Autumn Cuellar, John Doyle, Serge Dronov, Drew Endy, David Fell, Carl Firth, Ed Frank, Akira Funahashi, Ralph Gauges, Martin Ginkel, Victoria Gor, Igor Goryanin, Warren Hedley, Charles Hodgman, Stefan Hoops, Nick Juty, Jay Kaserger, Sarah Keating, Hiroaki Kitano, Ben Kovitz, Andreas Kremling, Nicolas Le Novère, Fred Livingston, Les Loew, Daniel Lucio, Joanne Matthews, Mike McCollum, Pedro Mendes, Eric Minch, Eric Mjolsness, David Morley, Mineo Morohashi, Poul Nielsen, Greg Peterson, Mark Poolman, Carole Proctor, Wayne Rindone, Sven Sahle, Takeshi Sakurada, Vijay Saraswat, Herbert Sauro, James Schaff, Maria Schilstra, John Schwacke, Cliff Shaffer, Bruce Shapiro, Tom Shimizu, Herbert Sauro, Hugh Spence, Jörg Stelling, Kouichi Takahashi, Masaru Tomita, Marc Vass, John Wagner, Jonathan Webb, Jörg Weimar, Darren Wilkinson, Olaf Wolkenhauer and Tau-Mu Yi.

Appendix

A Summary of Notation

The definitive explanation for the notation used in this document can be found in the companion notation document (Hucka, 2000). Here we briefly summarize some of the main components of the notations used in describing SBML.

Within the definitions of the various object classes introduced in this document, the following types of expressions are used many times:

The symbols field1, field2, etc., represents fields in a data structure. The colon immediately after the name separates the name of the attribute from the type of data that it stores.

More complex specifications use square brackets ([]) just after a type name. This is used to indicate that the field contains a list of elements. Specifically, the notation [0..*] signifies a list containing zero or more elements; the notation [1..*] signifies a list containing at least one element; and so on. The approach used here to translate from a list form into XML is, first, create a subelement named listOf_____s, where the blank indicates the capitalized name of the field, and then put a list of elements named after the field as the content of the listOf_____s element. listOf_____s elements representing list of the form [0..*] are optional: a missing listOf_____ element indicates that the list is empty. listOf_____ elements should always have content.

Expressions in curly braces ({}) shown after an attribute type indicate additional constraints placed on the field. We express constraints using XML Schema language. In the examples above, the expression {use="optional" default="0.0"} indicates that the field field3 is optional and that it has a default value of 0.0. A constraint of the form {namespace="X"} indicates that the field is not in the SBML Level 2 XML namespace but resides in the given XML namespace X. If a field is in a different namespace then the type of the field will not be defined by the SBML UML. In the examples above, the math field and its content is defined in the MathML namespace.

A field definition of the form X:(A:B) defines an element X that contains a field A with type B. If A is the string ANY then the element X contains an arbitrary sequence of elements. A field definition of the form X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content is constrained by constraint X:(A:B) is similar except that the field X and its content X:(A:B) is similar except that the field X and its content X is similar except that X is similar except that

B Differences between Level 1 Version 2 and Level 2

Compared to SBML Level 1 Version 2, SBML Level 2 introduces the following changes:

- SBML Level 2 supports the inclusion of metadata using the same approach as CellML (Cuellar et al., 2002). All structures in SBML can be annotated with optional content in RDF (Resource Description Format; Lassila and Swick, 1999) following the guidelines put forward by Cuellar et al.. (Section 3.1.)
- All data structures, including Sbml and listOf______ elements, are now derived from the type SBase. (Section 3.1.) This means all major structures in SBML can have separate annotations and metadata associated with them.
- A new field, id, replaces the name field previously defined for most SBML structures to identify each part of a model. (See Section 3.3.) The id field has a type of SId, whose definition is similar to SName

- in Level 1. In SBML Level 2, the name field is optional and is defined to allow any Unicode characters allowed by the string type of XML Schema (Biron and Malhotra, 2000).
- Formulas in Level 2 are expressed using MathML (W3C, 2000b) 2.0. The field named formula previously available on the KineticLaw and Rule structures has been replaced by a MathML element named math containing MathML content. In addition, stoichiometry numbers are now also expressed using MathML, allowing for more flexibility in defining reactions. (Sections 3.6, 4.8 and 4.9.)
- The namespace for identifiers in a model does not contain any built-in symbols; gone, for example, are the predefined rate laws of SBML Level 1. The approach taken in SBML Level 2 is that each model must itself define whatever functions it needs using the new FunctionDefinition mechanism. Although SBML Level 2 does define two built-in entities (a symbol representing time and another symbol representing delay functions), these are referenced using a feature of MathML and are not in the same namespace as identifiers defined by a model. (Section 3.6.4.)
- SBML Level 2 makes explicit a previously unstated assumption, that the XML encoding of a model uses UTF-8. SBML documents must refer to the UTF-8 encoding in their XML declaration. (Section 4.1.)
- The top-level Model structure can contain an optional list of global user-defined functions expressed in MathML and organized in new structures of type FunctionDefinition. (Sections 4.2 and 4.3.)
- The top-level Model structure can contain an optional list of event definitions organized in structures of type Event. Events define discrete changes in model behavior at specific times during a time simulation of the model. (Section 4.2 and 4.10.)
- Unlike in SBML Level 1, unit identifiers in Level 2 are in a separate namespace from the namespace used for models, functions, species, compartments, reactions and parameters. Also, the unit names "meter" and "liter" are not defined in Level 2 because the SBML user community deemed them unnecessary. Finally, Unit structures now have the additional fields multiplier and offset to enable the definition of non-SI units. (Section 4.4.)
- The Compartment structure has a new field, spatialDimensions, whose value is a positive integer specifying the number of dimensions in space the compartment possesses. This enables the definition of such things as two-dimensional membranes. As a side-effect, the units of species concentration in SBML Level 2 depend on the spatial dimensions of the compartment where the species is located. To support these new capabilities, Compartment now uses a field named size instead of volume, and there are two new built-in units for area and length. (Sections 4.4 and 4.5.)
- All attributes representing initial conditions or parameter values, including compartment sizes and species concentrations, are optional in Level 2. A missing value for one of these fields implies that the value is either unknown, not required for analysis, or should be obtained from an external source. (Sections 4.5, 4.6 and 4.7.)
- The Compartment, Species and Parameter structures each have a new boolean field named constant. This field specifies whether the variables represented by these structures can be changed by rules and reactions. (Sections 4.5, 4.6 and 4.7.)
- The Species structure has a new field, initialConcentration, for setting the initial value of a species in terms of its concentration. This is in addition to the ability, carried over from Level 1, to set the values in terms of amounts. (Section 4.6.)
- The rule structures are simpler compared to SBML Level 1. There is no longer a type attribute on Rule, and new structures AssignmentRule and RateRule replace SBML Level 1's ParameterRule, SpeciesConcentrationRule and CompartmentVolumeRule. (Section 4.8.)
- The Reaction structure has a new list of *modifiers* in addition to the list of reactants and products. The listOfModifiers enumerates species that affect a reaction but are neither created nor destroyed by the reaction. (Section 4.9.)

C XML Schema for SBML

The following is an XML Schema definition for the Systems Biology Markup Language, using the W3C Recommendation for XML Schema version 1.0 of 2 May 2001 (Biron and Malhotra, 2000; Fallside, 2000; Thompson et al., 2000).

```
<?xml version="1.0" encoding="UTF-8"?>
< xsd:schema
        targetNamespace="http://www.sbml.org/sbml/level2"
        xmlns:xsd="http://www.w3.org/2001/XMLSchema"
        xmlns:xsi="http://www.w3.org/2001/XMLSchema-instance"
        xmlns:xlink="http://www.w3.org/1999/xlink"
        xmlns:mml="http://www.w3.org/1998/Math/MathML"
        xmlns="http://www.sbml.org/sbml/level2"
        elementFormDefault="qualified"
        attributeFormDefault="unqualified"
        version="1">
    <xsd:import</pre>
        namespace="http://www.w3.org/1998/Math/MathML"
        schemaLocation="http://www.w3.org/Math/XMLSchema/mathml2/mathml2.xsd"/>
    <xsd:annotation>
        <xsd:documentation>
      File name : sbml.xsd
      Author : M. Hucka, A. Finney, D. Lucio
     Description: XML Schema for the Systems Biology Markup Language Level 2.
                    This is designed for XML Schema version 1.0.
      Version: 1
      Copyright 2003 California Institute of Technology and Japan Science and
     Technology Corporation.
      This library is free software; you can redistribute it and/or modify it
      under the terms of the GNU Lesser General Public License as published
      by the Free Software Foundation; either version 2.1 of the License, or
      any later version.
      This file is distributed in the hope that it will be useful, but
     WITHOUT ANY WARRANTY, WITHOUT EVEN THE IMPLIED WARRANTY OF
     MERCHANTABILITY OR FITNESS FOR A PARTICULAR PURPOSE. The software
      and documentation provided hereunder is on an "as is" basis, and the
      California Institute of Technology and Japan Science and Technology
      Corporation have no obligations to provide maintenance, support,
     updates, enhancements or modifications. In no event shall the
     California Institute of Technology or the Japan Science and Technology
      Corporation be liable to any party for direct, indirect, special,
      incidental or consequential damages, including lost profits, arising
      out of the use of this software and its documentation, even if the
      California Institute of Technology and/or Japan Science and Technology
      Corporation have been advised of the possibility of such damage. See
     the GNU Lesser General Public License for more details.
      You should have received a copy of the GNU Lesser General Public License
      along with this library; if not, write to the Free Software Foundation,
      Inc., 59 Temple Place, Suite 330, Boston, MA 02111-1307 USA.
</xsd:documentation>
    </xsd:annotation>
    <!--The definition of SId follows.-->
    <xsd:simpleType name="SId">
        <xsd:annotation>
            <xsd:documentation>
                The type SId is used throughout SBML as the type of the 'id'
                attributes on model elements.
            </xsd:documentation>
        </xsd:annotation>
        <xsd:restriction base="xsd:string">
            <xsd:pattern value="(_|[a-z]|[A-Z])(_|[a-z]|[A-Z]|[0-9])*"/>
        </xsd:restriction>
    </xsd:simpleType>
    <!--The definition of SBase follows.-->
```

```
<xsd:complexType name="SBase" abstract="true">
    <xsd:annotation>
        <xsd:documentation>
            The SBase type is the base type of all main components in SBML.
            It supports attaching metadata, notes and annotations to components.
        </xsd:documentation>
    </xsd:annotation>
    <xsd:sequence>
        <xsd:element name="notes" min0ccurs="0">
            <xsd:complexType>
                <xsd:sequence>
                     < xsd: anv
                         namespace="http://www.w3.org/1999/xhtml"
                         processContents="skip"
                         maxOccurs="unbounded"/>
                </xsd:sequence>
            </xsd:complexType>
        </xsd:element>
        <xsd:element name="annotation" minOccurs="0">
            <xsd:complexType>
                <xsd:sequence>
                     <xsd:any processContents="skip" maxOccurs="unbounded"/>
                </xsd:sequence>
            </xsd:complexType>
        </xsd:element>
    </xsd:sequence>
    <xsd:attribute name="metaid" type="xsd:ID" use="optional"/>
</xsd:complexType>
<!--The definition of FunctionDefinition follows.-->
<xsd:complexType name="FunctionDefinition">
    <xsd:complexContent>
        <xsd:extension base="SBase">
            <xsd:sequence>
                <xsd:element ref="mml:math"/>
            </xsd:sequence>
            <xsd:attribute name="id" type="SId" use="required"/>
<xsd:attribute name="name" type="xsd:string" use="optional"/>
        </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<!--The definition of UnitKind follows.-->
<xsd:simpleType name="UnitKind">
    <xsd:restriction base="xsd:string">
        <xsd:enumeration value="ampere"/>
        <xsd:enumeration value="becquerel"/>
        <xsd:enumeration value="candela"/>
        <xsd:enumeration value="Celsius"/>
        <xsd:enumeration value="coulomb"/>
        <xsd:enumeration value="dimensionless"/>
        <xsd:enumeration value="farad"/>
        <xsd:enumeration value="gram"/>
        <xsd:enumeration value="gray"/>
        <xsd:enumeration value="henry"/>
        <xsd:enumeration value="hertz"/>
        <xsd:enumeration value="item"/>
        <xsd:enumeration value="joule"/>
        <xsd:enumeration value="katal"/>
        <xsd:enumeration value="kelvin"/>
        <xsd:enumeration value="kilogram"/>
        <xsd:enumeration value="litre"/>
        <xsd:enumeration value="lumen"/>
        <xsd:enumeration value="lux"/>
        <xsd:enumeration value="metre"/>
        <xsd:enumeration value="mole"/>
        <xsd:enumeration value="newton"/>
        <xsd:enumeration value="ohm"/>
        <xsd:enumeration value="pascal"/>
<xsd:enumeration value="radian"/>
        <xsd:enumeration value="second"/>
```

```
<xsd:enumeration value="siemens"/>
         <xsd:enumeration value="sievert"/>
         <xsd:enumeration value="steradian"/>
         <xsd:enumeration value="tesla"/>
         <xsd:enumeration value="volt"/>
         <xsd:enumeration value="watt"/>
         <xsd:enumeration value="weber"/>
    </xsd:restriction>
</xsd:simpleType>
<!--The definition of Unit follows.-->
<xsd:complexType name="Unit">
    <xsd:complexContent>
         <xsd:extension base="SBase">
             <xsd:attribute name="kind" type="UnitKind" use="required"/>
<xsd:attribute name="exponent" type="xsd:integer" default="1"/>
             <xsd:attribute name="scale" type="xsd:integer" default="0"/>
             <xsd:attribute name="multiplier" type="xsd:double" default="1"/>
<xsd:attribute name="offset" type="xsd:double" default="0"/>
         </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<!--The definition of UnitDefinition follows.-->
<xsd:complexType name="ListOfUnits">
    <xsd:complexContent>
         <xsd:extension base="SBase">
             <xsd:sequence>
                  <xsd:element name="unit" type="Unit" maxOccurs="unbounded"/>
             </xsd:sequence>
         </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="UnitDefinition">
    <xsd:complexContent>
         <xsd:extension base="SBase">
             <xsd:sequence>
                  <xsd:element name="listOfUnits" type="ListOfUnits"/>
             </xsd:sequence>
             <xsd:attribute name="id" type="SId" use="required"/>
             <xsd:attribute name="name" type="xsd:string" use="optional"/>
         </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<!--The definition of Compartment follows.-->
<xsd:complexType name="Compartment">
    <xsd:complexContent>
         <xsd:extension base="SBase">
             <xsd:attribute name="id" type="SId" use="required"/>
             <xsd:attribute name="name" type="xsd:string" use="optional"/>
<xsd:attribute name="size" type="xsd:double" use="optional"/>
             <xsd:attribute name="spatialDimensions" use="optional" default="3">
                  <xsd:simpleType>
                       <xsd:restriction base="xsd:integer">
                           <xsd:minInclusive value="0"/>
                           <xsd:maxInclusive value="3"/>
                      </xsd:restriction>
                  </xsd:simpleType>
             </xsd:attribute>
             <xsd:attribute name="units" type="SId" use="optional"/>
             <xsd:attribute name="outside" type="SId" use="optional"/>
<xsd:attribute name="constant" type="xsd:boolean" use="optional" default="true"/>
         </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<!--The definition of Species follows.-->
<xsd:complexType name="Species">
    <xsd:complexContent>
         <xsd:extension base="SBase">
             <xsd:attribute name="id" type="SId" use="required"/>
             <xsd:attribute name="name" type="xsd:string" use="optional"/>
```

```
<xsd:attribute name="compartment" type="SId"/>
              <xsd:attribute name="initialAmount" type="xsd:double" use="optional"/>
              <xsd:attribute name="initialConcentration" type="xsd:double" use="optional"/>
<xsd:attribute name="substanceUnits" type="SId" use="optional"/>
<xsd:attribute name="spatialSizeUnits" type="SId" use="optional"/>
                  name="boundaryCondition" type="xsd:boolean" use="optional" default="false"/>
              <xsd:attribute name="charge" type="xsd:integer" use="optional"/>
<xsd:attribute name="constant" type="xsd:boolean" use="optional" default="false"/>
         </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<!--The definition of Parameter follows.-->
<xsd:complexType name="Parameter">
    <xsd:complexContent>
         <xsd:extension base="SBase">
              <xsd:attribute name="id" type="SId" use="required"/>
<xsd:attribute name="name" type="xsd:string" use="optional"/>
<xsd:attribute name="value" type="xsd:double" use="optional"/>
              <xsd:attribute name="units" type="SId" use="optional"/>
<xsd:attribute name="constant" type="xsd:boolean" use="optional" default="true"/>
         </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="ListOfParameters">
    <xsd:complexContent>
         <xsd:extension base="SBase">
                   <xsd:element name="parameter" type="Parameter" maxOccurs="unbounded"/>
              </xsd:sequence>
         </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<!--The definition of Rule follows. -->
<xsd:complexType name="Rule" abstract="true">
    <xsd:complexContent>
         <xsd:extension base="SBase">
              <xsd:sequence>
                   <xsd:element ref="mml:math"/>
              </xsd:sequence>
         </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="AlgebraicRule">
    <xsd:complexContent>
         <xsd:extension base="Rule"/>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="AssignmentRule">
    <xsd:complexContent>
         <xsd:extension base="Rule">
              <xsd:attribute name="variable" type="SId" use="required"/>
         </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="RateRule">
    <xsd:complexContent>
         <xsd:extension base="Rule">
              <xsd:attribute name="variable" type="SId" use="required"/>
         </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<!--The definition of Reaction follows.-->
<xsd:complexType name="KineticLaw">
    <xsd:complexContent>
         <xsd:extension base="SBase">
              <xsd:sequence>
                   <xsd:element ref="mml:math"/>
                   <xsd:element name="listOfParameters" type="ListOfParameters" minOccurs="0"/>
```

```
</xsd:sequence>
            <xsd:attribute name="timeUnits" type="SId" use="optional"/>
            <xsd:attribute name="substanceUnits" type="SId" use="optional"/>
        </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="SimpleSpeciesReference" abstract="true">
    <xsd:complexContent>
        <xsd:extension base="SBase">
            <xsd:attribute name="species" type="SId" use="required"/>
        </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="ModifierSpeciesReference">
    <xsd:complexContent>
        <xsd:extension base="SimpleSpeciesReference"/>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="ListOfModifierSpeciesReferences">
   <xsd:complexContent>
        <xsd:extension base="SBase">
            <xsd:sequence>
                <xsd:element</pre>
                    name="modifierSpeciesReference"
                    type="ModifierSpeciesReference"
                    maxOccurs="unbounded"/>
            </xsd:sequence>
        </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="StoichiometryMath">
   <xsd:complexContent>
        <xsd:extension base="SBase">
            <xsd:sequence>
                <xsd:element ref="mml:math"/>
            </xsd:sequence>
        </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="SpeciesReference">
    <xsd:complexContent>
        <xsd:extension base="SimpleSpeciesReference">
            <xsd:sequence>
                <xsd:element name="stoichiometryMath" type="StoichiometryMath" minOccurs="0"/>
            </xsd:sequence>
            <xsd:attribute name="stoichiometry" type="xsd:double" use="optional" default="1"/>
        </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="ListOfSpeciesReferences">
   <xsd:complexContent>
        <xsd:extension base="SBase">
            <xsd:sequence>
                <xsd:element</pre>
                    name="speciesReference" type="SpeciesReference" maxOccurs="unbounded"/>
            </xsd:sequence>
        </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="Reaction">
    <xsd:complexContent>
        <xsd:extension base="SBase">
            <xsd:sequence>
                <xsd:element</pre>
                    name="listOfReactants" type="ListOfSpeciesReferences" minOccurs="0"/>
                    name="listOfProducts" type="ListOfSpeciesReferences" minOccurs="0"/>
                <xsd:element</pre>
                    name="listOfModifiers"
```

```
type="ListOfModifierSpeciesReferences"
                    minOccurs="0"/>
                 <xsd:element</pre>
                    name="kineticLaw" type="KineticLaw" minOccurs="0"/>
            </xsd:sequence>
            <xsd:attribute name="id" type="SId" use="required"/>
            <xsd:attribute name="name" type="xsd:string" use="optional"/>
            <xsd:attribute name="reversible" type="xsd:boolean" use="optional" default="true"/>
            <xsd:attribute name="fast" type="xsd:boolean" use="optional"/>
        </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<!--The definition of Event follows.-->
<xsd:complexType name="EventAssignment">
    <xsd:complexContent>
        <xsd:extension base="SBase">
            <xsd:sequence>
                <xsd:element ref="mml:math"/>
            <xsd:attribute name="variable" type="SId" use="required"/>
        </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="ListOfEventAssignments">
    <xsd:complexContent>
        <xsd:extension base="SBase">
            <xsd:sequence>
                 <xsd:element</pre>
                    name="eventAssignment" type="EventAssignment" maxOccurs="unbounded"/>
            </xsd:sequence>
        </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="MathField">
    <xsd:complexContent>
        <xsd:extension base="SBase">
            <xsd:sequence>
                <xsd:element ref="mml:math"/>
            </xsd:sequence>
        </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<xsd:complexType name="Event">
    <xsd:complexContent>
        <xsd:extension base="SBase">
            <xsd:sequence>
                 <xsd:element name="trigger" type="MathField"/>
                <xsd:element name="delay" type="MathField" minOccurs="0"/>
                <xsd:element name="listOfEventAssignments" type="ListOfEventAssignments"/>
            </xsd:sequence>
            <xsd:attribute name="id" type="SId" use="optional"/>
            <xsd:attribute name="name" type="xsd:string" use="optional"/>
<xsd:attribute name="timeUnits" type="SId" use="optional"/>
        </xsd:extension>
    </xsd:complexContent>
</xsd:complexType>
<!-- The definition of Model follows.-->
<xsd:complexType name="Model">
    <xsd:complexContent>
        <xsd:extension base="SBase">
            <xsd:sequence>
                <xsd:element name="listOfFunctionDefinitions" minOccurs="0">
                     <xsd:complexType>
                         <xsd:complexContent>
                             <xsd:extension base="SBase">
                                  <xsd:sequence>
                                      <xsd:element</pre>
                                          name="functionDefinition"
                                          type="FunctionDefinition"
```

```
maxOccurs="unbounded"/>
                 </xsd:sequence>
            </xsd:extension>
        </xsd:complexContent>
    </rsd:complexType>
</xsd:element>
<xsd:element name="listOfUnitDefinitions" minOccurs="0">
    <xsd:complexType>
        <xsd:complexContent>
            <xsd:extension base="SBase">
                 <xsd:sequence>
                     <xsd:element</pre>
                         name="unitDefinition"
                         type="UnitDefinition"
                         maxOccurs="unbounded"/>
                 </xsd:sequence>
            </xsd:extension>
        </xsd:complexContent>
    </xsd:complexType>
</xsd:element>
<xsd:element name="listOfCompartments" minOccurs="0">
    <xsd:complexType>
        <xsd:complexContent>
            <xsd:extension base="SBase">
                 <xsd:sequence>
                     <xsd:element</pre>
                         name="compartment"
                         type="Compartment"
                         maxOccurs="unbounded"/>
                 </xsd:sequence>
            </xsd:extension>
        </xsd:complexContent>
    </xsd:complexType>
</xsd:element>
<xsd:element name="listOfSpecies" minOccurs="0">
    <xsd:complexType>
        <xsd:complexContent>
            <xsd:extension base="SBase">
                <xsd:sequence>
                     <xsd:element</pre>
                         name="species" type="Species" maxOccurs="unbounded"/>
                 </xsd:sequence>
            </xsd:extension>
        </xsd:complexContent>
    </xsd:complexType>
</xsd:element>
<xsd:element name="listOfParameters" minOccurs="0">
    <xsd:complexType>
        <xsd:complexContent>
            <xsd:extension base="SBase">
                <xsd:sequence>
                     <xsd:element</pre>
                         name="parameter"
type="Parameter"
                         maxOccurs="unbounded"/>
                </xsd:sequence>
            </xsd:extension>
        </xsd:complexContent>
    </xsd:complexType>
</xsd:element>
<xsd:element name="listOfRules" minOccurs="0">
    <xsd:complexType>
        <xsd:complexContent>
            <xsd:extension base="SBase">
                 <xsd:choice maxOccurs="unbounded">
                     <xsd:element</pre>
                         name="algebraicRule"
                         type="AlgebraicRule"
                         minOccurs="0"/>
```

```
<xsd:element</pre>
                                                   name="assignmentRule"
                                                   type="AssignmentRule"
                                                   minOccurs="0"/>
                                              <xsd:element</pre>
                                                   name="rateRule"
                                                   type="RateRule"
                                                   minOccurs="0"/>
                                          </xsd:choice>
                                     </xsd:extension>
                                </xsd:complexContent>
                            </xsd:complexType>
                       </xsd:element>
                       <xsd:element name="listOfReactions" minOccurs="0">
                           <xsd:complexType>
                                <xsd:complexContent>
                                     <xsd:extension base="SBase">
                                          <xsd:sequence>
                                              <xsd:element</pre>
                                                  name="reaction" type="Reaction" maxOccurs="unbounded"/>
                                         </xsd:sequence>
                                     </xsd:extension>
                                </xsd:complexContent>
                            </xsd:complexType>
                       </xsd:element>
                       <xsd:element name="listOfEvents" minOccurs="0">
                           <xsd:complexType>
                                <xsd:complexContent>
                                     <xsd:extension base="SBase">
                                          <xsd:sequence>
                                              <xsd:element</pre>
                                                  name="event" type="Event" maxOccurs="unbounded"/>
                                         </xsd:sequence>
                                     </xsd:extension>
                                </xsd:complexContent>
                            </xsd:complexType>
                       </xsd:element>
                  </xsd:sequence>
                  <xsd:attribute name="id" type="SId" use="optional"/>
<xsd:attribute name="name" type="xsd:string" use="optional"/>
              </xsd:extension>
         </xsd:complexContent>
    </xsd:complexType>
    <!-- The following is the type definition for the top-level element in an SBML document.-->
    <xsd:complexType name="Sbml">
         <xsd:complexContent>
              <xsd:extension base="SBase">
                  <xsd:sequence>
                       <xsd:element name="model" type="Model"/>
                  </xsd:sequence>
                  <xsd:attribute name="level" type="xsd:positiveInteger" use="required" fixed="2"/>
<xsd:attribute name="version" type="xsd:positiveInteger" use="required" fixed="1"/>
              </xsd:extension>
         </xsd:complexContent>
    </xsd:complexType>
    <!--The following is the (only) top-level element allowed in an SBML document.--> <xsd:element name="sbml" type="Sbml"/>
     <!--The end.-->
</xsd:schema>
```

References

- Abbott, A. (1999). Alliance of US labs plans to build map of cell signalling pathways. Nature, 402:219–200.
- Arkin, A. P. (2001). Simulac and Deduce. Available via the World Wide Web at http://gobi.lbl.gov/~aparkin/Stuff/Software.html.
- Biron, P. V. and Malhotra, A. (2000). XML Schema part 2: Datatypes (W3C candidate recommendation 24 October 2000). Available via the World Wide Web at http://www.w3.org/TR/xmlschema-2/.
- Bosak, J. and Bray, T. (1999). XML and the second-generation Web. Scientific American, 280(5):89–93.
- Bray, D., Firth, C., Le Novère, N., and Shimizu, T. (2001). *StochSim*. Available via the World Wide Web at http://www.zoo.cam.ac.uk/comp-cell/StochSim.html.
- Bray, T., D. Hollander, D., and Layman, A. (1999). Namespaces in XML. World Wide Web Consortium 14-January-1999. Available via the World Wide Web at http://www.w3.org/TR/1999/REC-xml-names-19990114/.
- Bray, T., Paoli, J., Sperberg-McQueen, C. M., and Maler, E. (2000). Extensible markup language (XML) 1.0 (second edition), W3C recommendation 6-October-2000. Available via the World Wide Web at http://www.w3.org/TR/1998/REC-xml-19980210.
- Bureau International des Poids et Mesures (2000). The International System of Units (SI) supplement 2000: addenda and corrigenda to the 7th edition (1998). Available via the World Wide Web at http://www.bipm.fr/pdf/si-supplement2000.pdf.
- Cuellar, A. A., Nelson, M., and Hedley, W. (2002). The CellML metadata 1.0 specification working draft—16 January 2002. Available via the World Wide Web at http://cellml.org/public/metadata/cellml_metadata_specification.html.
- Eriksson, H.-E. and Penker, M. (1998). UML Toolkit. John Wiley & Sons, New York.
- Fallside, D. C. (2000). XML Schema part 0: Primer (W3C candidate recommendation 24 October 2000). Available via the World Wide Web at http://www.w3.org/TR/xmlschema-0/.
- Gilman, A. (2000). A letter to the signaling community. Alliance for Cellular Signaling, The University of Texas Southwestern Medical Center. Available via the World Wide Web at http://afcs.swmed.edu/afcs/Letter_to_community.htm.
- Goryanin, I. (2001). DBsolve: Software for metabolic, enzymatic and receptor-ligand binding simulation. Available via the World Wide Web at http://homepage.ntlworld.com/igor.goryanin/.
- Goryanin, I., Hodgman, T. C., and Selkov, E. (1999). Mathematical simulation and analysis of cellular metabolism and regulation. *Bioinformatics*, 15(9):749–758.
- Harold, E. R. and Means, E. S. (2001). XML in a Nutshell. O'Reilly & Associates.
- Hedley, W. J., Nelson, M. R., Bullivant, D., Cuellar, A., Ge, Y., Grehlinger, M., Jim, K., Lett, S., Nickerson, D., Nielsen, P., and Yu, H. (2001). CellML specification. Available via the World Wide Web at http://www.cellml.org/public/specification/20010810/cellml_specification.html.
- Hucka, M. (2000). SCHUCS: A notation for describing model representations intended for XML encoding. Available via the World Wide Web at http://www.sbml.org/.
- Hucka, M., Finney, A., Sauro, H. M., and Bolouri, H. (2001). Systems Biology Markup Language (SBML) Level 1: Structures and facilities for basic model definitions. Available via the World Wide Web at http://www.sbml.org.

- Hucka, M., Finney, A., Sauro, H. M., Bolouri, H., Doyle, J. C., Kitano, H., Arkin, A. P., Bornstein, B. J., Bray, D., Cornish-Bowden, A., Cuellar, A. A., Dronov, S., Gilles, E. D., Ginkel, M., Gor, V., Goryanin, I. I., Hedley, W. J., Hodgman, T. C., Hofmeyr, J.-H., Hunter, P. J., Juty, N. S., Kasberger, J. L., Kremling, A., Kummer, U., Le Novère, N., Loew, L. M., Lucio, D., Mendes, P., Minch, E., Mjolsness, E. D., Nakayama, Y., Nelson, M. R., Nielsen, P. F., Sakurada, T., Schaff, J. C., Shapiro, B. E., Shimizu, T. S., Spence, H. D., Stelling, J., Takahashi, K., Tomita, M., Wagner, J., and Wang, J. (2003). The Systems Biology Markup Language (SBML): A medium for representation and exchange of biochemical network models. Bioinformatics, 19(4):524–531.
- Lassila, O. and Swick, R. (1999). Resource description framework (RDF) model and syntax specification. Available via the World Wide Web at http://www.w3.org/TR/REC-rdf-syntax/.
- Mendes, P. (1997). Biochemistry by numbers: Simulation of biochemical pathways with Gepasi 3. *Trends in Biochemical Sciences*, 22:361–363.
- Mendes, P. (2000). New research software to simulate biochemical processes. Available via the World Wide Web at http://www.vbi.vt.edu/pr/press_releases/press_20001218_news_new-software.htm.
- Mendes, P. (2001). Gepasi 3.21. Available via the World Wide Web at http://www.gepasi.org.
- Morton-Firth, C. J. and Bray, D. (1998). Predicting temporal fluctuations in an intracellular signalling pathway. *Journal of Theoretical Biology*, 192:117–128.
- Oestereich, B. (1999). Developing Software with UML: Object-Oriented Analysis and Design in Practice. Addison-Wesley Publishing Company.
- Popel, A. and Winslow, R. L. (1998). A letter from the directors... Center for Computational Medicine & Biology, Johns Hopkins School of Medicine, Johns Hopkins University. Available via the World Wide Web at http://www.bme.jhu.edu/ccmb/ccmbletter.html.
- Sauro, H. M. (2000). Jarnac: A system for interactive metabolic analysis. In Hofmeyr, J.-H. S., Rohwer, J. M., and Snoep, J. L., editors, Animating the Cellular Map: Proceedings of the 9th International Meeting on BioThermoKinetics. Stellenbosch University Press.
- Sauro, H. M. and Fell, D. A. (1991). SCAMP: A metabolic simulator and control analysis program. *Mathl. Comput. Modelling*, 15:15–28.
- Sauro, H. S. (2001). JDesigner: A simple biochemical network designer. Available via the World Wide Web at http://members.tripod.co.uk/sauro/biotech.htm.
- Schaff, J., Slepchenko, B., and Loew, L. M. (2000). Physiological modeling with the Virtual Cell framework. In Johnson, M. and Brand, L., editors, *Methods in Enzymology*, volume 321, pages 1–23. Academic Press, San Diego.
- Schaff, J., Slepchenko, B., Morgan, F., Wagner, J., Resasco, D., Shin, D., Choi, Y. S., Loew, L., Carson, J., Cowan, A., Moraru, I., Watras, J., Teraski, M., and Fink, C. (2001). Virtual Cell. Available via the World Wide Web at http://www.nrcam.uchc.edu.
- Schilstra, M. and Bolouri, H. (2002). Netbuilder. Available via the World Wide Web at http://strc.herts.ac.uk/bio/maria/NetBuilder/index.html.
- Shapiro, B. E., Levchenko, A., and Mjolsness, E. (2001). Automatic model generation for signal transduction with applications to MAP-kinase pathways. In Kitano, H., editor, *Foundations of Systems Biology*, chapter 7, pages 145–161. MIT Press.
- Shapiro, B. E., Levchenko, A., and Mjolsness, E. D. (2003). Cellerator web site. Available via the World Wide Web at http://www-aig.jpl.nasa.gov/public/mls/cellerator.
- Smaglik, P. (2000). For my next trick... Nature, 407:828–829.

- Stelling, J., Ginkel, M., Bettenbrok, K., and Gilles, E. D. (2001). Towards a virtual biological laboratory. In Kitano, H., editor, *Foundations of Systems Biology*, chapter 5, pages 189–212. MIT Press.
- Thompson, H. S., Beech, D., Maloney, M., and Mendelsohn, N. (2000). XML Schema part 1: Structures (W3C candidate recommendation 24 October 2000). Available via the World Wide Web at http://www.w3.org/TR/xmlschema-1/.
- Tomita, M., Hashimoto, K., Takahashi, K., Shimizu, T., Matsuzaki, Y., Miyoshi, F., Saito, K., Tanida, S., Yugi, K., Venter, J. C., and Hutchison, C. (1999). E-Cell: Software environment for whole cell simulation. *Bioinformatics*, 15(1):72–84.
- Tomita, M., Nakayama, Y., Naito, Y., Shimizu, T., Hashimoto, K., Takahashi, K., Matsuzaki, Y., Yugi, K., Miyoshi, F., Saito, Y., Kuroki, A., Ishida, T., Iwata, T., Yoneda, M., Kita, M., Yamada, Y., Wang, E., Seno, S., Okayama, M., Kinoshita, A., Fujita, Y., Matsuo, R., Yanagihara, T., Watari, D., Ishinabe, S., and Miyamoto, S. (2001). E-Cell. Available via the World Wide Web at http://www.e-cell.org/.
- Unicode Consortium (1996). The Unicode Standard, Version 2.0. Addison-Wesley Developers Press, Reading, Massachusetts.
- Vass, M., Shaffer, C. A., Tyson, J. J., Ramakrishnan, N., and Watson, L. T. (2003). The jigcell model builder: A tool for modeling intra-cellular regulatory networks. *Bioinformatics*. In Press.
- W3C (2000a). Naming and addressing: URIs, URLs, ... Available via the World Wide Web at http://www.w3.org/Addressing/.
- W3C (2000b). W3C's math home page. Available via the World Wide Web at http://www.w3.org/Math/.