The Systems Biology Markup Language (SBML) Level 3 Package: Flux Balance Constraints

Brett G. Olivier¹* and Frank T. Bergmann²

¹Systems Bioinformatics, VU University Amsterdam, The Netherlands
²Modelling of Biol. Processes, Heidelberg University, Germany

Summary

Constraint-based modeling is a well established modelling methodology used to analyze and study biological networks on both a medium and genome scale. Due to their large size, genome scale models are typically analysed using constraint-based optimization techniques. One widely used method is Flux Balance Analysis (FBA) which, for example, requires a modelling description to include: the definition of a stoichiometric matrix, an objective function and bounds on the values that fluxes can obtain at steady state. The Flux Balance Constraints (FBC) Package extends SBML Level 3 and provides a standardized format for the encoding, exchange and annotation of constraint-based models. It includes support for modelling concepts such as objective functions, flux bounds and model component annotation that facilitates reaction balancing. The FBC package establishes a base level for the unambiguous exchange of genome-scale, constraint-based models, that can be built upon by the community to meet future needs (e.g. by extending it to cover dynamic FBC models).

*To whom correspondence should be addressed. Email: sbml-editors@sbml.org
SBML Level 3 Package Specification

SBML Level 3 Package: Flux Balance Constraints (‘fbc’)

Brett G. Olivier  
b.g.olivier@vu.nl  
Systems Bioinformatics  
VU University Amsterdam  
Amsterdam, NH, The Netherlands

Frank T. Bergmann  
fbergmann@caltech.edu  
Computing and Mathematical Sciences  
California Institute of Technology  
Pasadena, CA, US

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The latest release, past releases, and other materials related to this specification are available at 

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Contents

1 Introduction and motivation 3
   1.1 Proposal corresponding to this package specification 3
   1.2 Tracking number 4
   1.3 Package dependencies 4
   1.4 Document conventions 4

2 Background 5
   2.1 Problems with current SBML approaches 5
   2.2 Past work on this problem or similar topics 5

3 Proposed syntax and semantics 6
   3.1 Namespace URI and other declarations necessary for using this package 6
   3.2 Primitive data types 6
      3.2.1 Type FbcType 6
      3.2.2 Type FbcOperation 6
   3.3 The extended Model class 7
      3.3.1 The FBC listOfFluxBounds 7
      3.3.2 The FBC listOfObjectives 7
      3.3.3 A note on units 8
   3.4 The extended Species class 8
   3.5 The FBC FluxBound class 9
   3.6 The FBC Objective class 11
   3.7 The FBC FluxObjective class 12

4 Illustrative examples of the FBC syntax 13
   4.1 Example one: the basic FBC syntax 13
      4.1.1 Kinetic model description 13
      4.1.2 Capacity constraints 14
      4.1.3 Objective function 14
      4.1.4 Complete worked example 15

5 Best practices 17
   5.1 Examples contrasting the current SBML L2 encoding with L3 and FBC 17

6 Future development 20
   6.1 Model annotation 20
      6.1.1 Gene association annotation 20
      6.1.2 A generic annotation mechanism 20

A Validation of SBML documents 23
   A.1 Validation and consistency rules 23

Acknowledgments 28

References 29
1 Introduction and motivation

Constraint based modeling is a widely accepted methodology used to analyze and study biological networks on both a small and whole organism (genome) scale. Typically these models are underdetermined and constraint based methods (e.g. linear, quadratic optimization) are used to optimize specific model properties. This is assumed to occur under a defined set of constraints (e.g. stoichiometric, metabolic) and bounds (e.g. thermodynamic, experimental and environmental) on the values that the solution fluxes can obtain.

Perhaps the most well known (and widely used) analysis method is Flux Balance Analysis (FBA; Orth et al., 2010) which is performed on Genome Scale Reconstructions (GSR's; Oberhardt et al., 2009). Using FBA a target flux is optimized (e.g. maximizing a flux to biomass or minimizing ATP production) while other fluxes can be bounded to simulate a selected growth environment or specific metabolic state.

As constraint based models are generally underdetermined, i.e. few or none of the kinetic rate equations and related parameters are known, it is crucial that a model definition includes the ability to define optimization parameters such as objective functions, flux bounds and constraints. Currently this is not possible in the Systems Biology Markup Language (SBML) Level 2 or Level 3 core specification (Hucka et al., 2011, 2003).

The question of how to encode constraint based (also referred to as steady state or FBA) models in SBML is not new. However, advances in the methods used to construct genome scale constraint based models and the wider adoption of constraint based modeling in biotechnological/medical applications have led to a rapid increase in both the number of models being constructed and the tools used to analyze them.

Faced with such growth, both in number and diversity, the need for a standardized data format for the definition, exchange and annotation of constraint based models has become critical. As the core model components (e.g. species, reactions, stoichiometry) can already be efficiently described in SBML (with its associated active community, software and tool support) the Flux Balance Constraints package aims to extend SBML Level 3 core by adding the elements necessary to encode current and future constraint based models.

1.1 Proposal corresponding to this package specification

This specification for Flux Balance Constraints in SBML Level 3 Version 1 is based on the proposal, by this documents authors, located at the following URL:


The tracking number in the SBML issue tracking system (SBML Team, 2010) for Flux Balance Constraints package activities is 3154219. The version of the proposal used as the starting point for this specification is the version of March 2012. Previous versions of the current proposal are:

Version 3 (March 2012)


Version 2 (March 2011)

http://sbml.org/Community/Wiki/SBML_Level_3_Proposals/Flux_Constraints_Proposal

Version 1 (February 2010)

http://precedings.nature.com/documents/4236/version/1

Details of earlier independent proposals are provided in Section 2.
1.2 Tracking number

As initially listed in the SBML issue tracking system under:

1.3 Package dependencies

The Flux Balance Constraints package adds additional classes to SBML Level 3 Version 1 Core and has no dependency on any other SBML Level 3 package.

1.4 Document conventions

Following the precedent set by the SBML Level 3 Core specification document, we use UML 1.0 (Unified Modeling Language; Eriksson and Penker 1998; Oestereich 1999) class diagram notation to define the constructs provided by this package. We also use color in the diagrams to carry additional information for the benefit of those viewing the document on media that can display color. The following are the colors we use and what they represent:

- **Black**: Items colored black in the UML diagrams are components taken unchanged from their definition in the SBML Level 3 Core specification document.

- **Green**: Items colored green are components that exist in SBML Level 3 Core, but are extended by this package. Class boxes are also drawn with dashed lines to further distinguish them.

- **Blue**: Items colored blue are new components introduced in this package specification. They have no equivalent in the SBML Level 3 Core specification.

We also use the following typographical conventions to distinguish the names of objects and data types from other entities; these conventions are identical to the conventions used in the SBML Level 3 Core specification document:

**AbstractClass**: Abstract classes are classes that are never instantiated directly, but rather serve as parents of other object classes. Their names begin with a capital letter and they are printed in a slanted, bold, sans-serif typeface. In electronic document formats, the class names defined within this document are also hyperlinked to their definitions; clicking on these items will, given appropriate software, switch the view to the section in this document containing the definition of that class. (However, for classes that are unchanged from their definitions in SBML Level 3 Core, the class names are not hyperlinked because they are not defined within this document.)

**Class**: Names of ordinary (concrete) classes begin with a capital letter and are printed in an upright, bold, sans-serif typeface. In electronic document formats, the class names are also hyperlinked to their definitions in this specification document. (However, as in the previous case, class names are not hyperlinked if they are for classes that are unchanged from their definitions in the SBML Level 3 Core specification.)

**SomeThing, otherThing**: Attributes of classes, data type names, literal XML, and generally all tokens other than SBML UML class names, are printed in an upright typewriter typeface. Primitive types defined by SBML begin with a capital letter; SBML also makes use of primitive types defined by XML Schema 1.0 (Biron and Malhotra, 2000; Fallside, 2000; Thompson et al., 2000), but unfortunately, XML Schema does not follow any capitalization convention and primitive types drawn from the XML Schema language may or may not start with a capital letter.

For other matters involving the use of UML and XML, we follow the conventions used in the SBML Level 3 Core specification document.
2 Background

2.1 Problems with current SBML approaches

While there is currently no official way of encoding constraint based models in SBML L2 there have been pragmatic approaches used by a variety of groups and applications. Arguably the most comprehensive and widely used format is that used by the COBRA toolbox (Becker et al., 2007) where the metabolic reaction network is well defined using SBML’s Reaction and Species classes. However, other FBA specific model components such as flux bounds and the reactions that take part in the objective function are less well defined. In this case LocalParameter elements are used which (implicitly) rely on e.g. all tools knowing and using the same naming convention for the parameter ID’s. Furthermore, reaction annotations are generally stored as tool specific HTML key-value pairs in a Notes element which has routinely led to different research groups and software using in-house and/or tool specific ways to describe the same information. While a step in the right direction this format is not suitable for implementation in SBML Level 3.

It is worth noting that while SBML Level 2 does have a construct know as Constraint its function is limited to measuring and reporting the model variables behavior in time. In contrast the FluxBound enforces the bounds on a steady-state flux and they can therefore be considered to be independent of one another. In addition, certain attributes that were widely used by the constraint based modelling community such as the Species attribute charge were removed from SBML. This has had the effect that a significant number of tools and models are still limited to using SBML Level 2 Version 1.

2.2 Past work on this problem or similar topics

The problem of describing and annotating FBA models in SBML has been raised at various times in the past few years. In this regard there are two known putative proposals one by Karthik Raman and the other by the Church Laboratory. As far as we are aware these proposals never developed beyond their initial presentation at SBML forums/hackathons. In 2009 the discussion was reopened at the SBML Forum held in Stanford and has subsequently developed into the current active package proposal and this document (see Section 1). In reverse chronological order these are:

Brett Olivier (2009) SBML Level 3 FBA package discussion

Karthik Raman (2005) Flux annotations in SBML

Church laboratory (pre 2005) Metabolic flux model annotations
http://sbml.org/Community/Wiki/Old_known_SBML_annotations_list
3 Proposed syntax and semantics

In this section, we define the syntax and semantics of the Flux Balance Constraints package for SBML Level 3 Version 1. We expound on the various data types and constructs defined in this package, then in Section 4 on page 13, we provide complete examples of using the constructs in an example SBML model.

3.1 Namespace URI and other declarations necessary for using this package

Every SBML Level 3 package is identified uniquely by an XML namespace URI. For an SBML document to be able to use a given SBML Level 3 package, it must declare the use of that package by referencing its URI. The following is the namespace URI for this version of the Flux Balance Constraints package for SBML Level 3 Version 1:

"http://www.sbml.org/sbml/level3/version1/fbc/version1"

In addition, SBML documents using a given package must indicate whether understanding the package is required for complete mathematical interpretation of a model, or whether the package is optional. This is done using the attribute **required** on the `<sbml>` element in the SBML document. For the Flux Balance Constraints package, the value of this attribute must be set to “false”.

The following fragment illustrates the beginning of a typical SBML model using SBML Level 3 Version 1 and this version of the Flux Balance Constraints package:

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

3.2 Primitive data types

Section 3.1 of the SBML Level 3 Version 1 Core specification defines a number of primitive data types and also uses a number of XML Schema 1.0 data types (Biron and Malhotra, 2000). More specifically we make use of **integer**, **double**, **string**, **SId** and **SIdRef**. In addition we make use of two new primitives: the enumerations **FbcType** and **FbcOperation**, see Figure 1 for the interrelation between these entities.

The **SId** type is used as the data type for the identifiers of **FluxBound** (Section 3.5), **FluxObjective** (Section 3.7) and **Objective** (Section 3.6) classes. In the FBC package the **ListOfObjectives** has an attribute of type **SIdRef** that is used to refer to an ‘active’ **Objective**.

3.2.1 Type FbcType

The Flux Balance Constraints package defines a new enumerated type **FbcType** which represents the optimization sense of the objective function. It can have one of the following two values "maximize" or "minimize".

3.2.2 Type FbcOperation

The Flux Balance Constraints package defines a new enumerated type **FbcOperation** which represents a boolean operator. It can take only one of the following values: “lessEqual”, “greaterEqual” or “equal”.

---

3.3 The extended Model class

The SBML Model class is extended with the addition of two children, i.e. a `listOfFluxBounds` and a `listOfObjectives` and a Model may contain at most one of these lists.

3.3.1 The FBC listOfFluxBounds

As shown in Figure 1 the `ListOfFluxBounds` is derived from `SBase` and inherits the attributes `metaid` and `sboTerm`, as well as the subcomponents for Annotation and Notes. `ListOfFluxBounds` must contain at least one `FluxBound` (defined in Section 3.5).

3.3.2 The FBC listOfObjectives

As shown in Figure 1 the `ListOfObjectives` is derived from `SBase` and inherits the attributes `metaid` and `sboTerm`, as well as the subcomponents for Annotation and Notes. Unlike most other SBML `ListOf__` classes, `ListOfObjectives` introduces an additional required attribute `activeObjective`. The `ListOfObjectives` must contain at least one `Objective` (defined in Section 3.6).

Figure 1: A UML representation of the Flux Balance Constraints package. Derived from `SBase`, the FBC classes inherit support for constructs such as SBML Notes and Annotation’s. See Section 1.4 for conventions related to this figure. The individual classes are further discussed in the text.
The activeObjective attribute

This attribute is of type SIdRef and can only refer to the id of an existing Objective. This required attribute exists so that when multiple Objective’s are included in a single model, the model will always be well described i.e. there is a single, primary objective function which defines a single optimum and its associated solution space.

3.3.3 A note on units

The main unit definitions that should be considered when using the Flux Balance Constraints package are the global model definitions of “extent” and “time” as all FBC flux related classes (i.e. FluxBound and FluxObjective implicitly attain the same unit as the Reaction that they reference). More details on units can be found in their respective class definitions.

3.4 The extended Species class

The Flux Balance Constraints package extends the SBML Level 3 Version 1 Core Species class with the addition of two attributes charge and chemicalFormula.

![Species (extended)](image)

Figure 2: A UML representation of the extended SBML Species class used in the Flux Balance Constraints package. See Section 1.4 for conventions related to this figure.

The charge attribute

The optional attribute charge which contains a signed integer referring to the Species object’s charge and is defined as it was in the SBML Level 2 Version 1 specification: “The optional field charge takes an integer indicating the charge on the species (in terms of electrons, not the SI unit coulombs).”

The chemicalFormula attribute

The optional attribute chemicalFormula containing a string that represents the Species objects elemental composition.

While there are many ways of referring to an elemental composition the purpose of the chemicalFormula attribute is to allow reaction balancing and validation which is particularly important in constraint based models.

The format of chemicalFormula must consist only of atomic names (as in the Periodic Table) or user defined compounds either of which take the form of a single capital letter followed by zero or more lowercase letters. Where there is more than a single atom present, this is indicated with an integer. With regards to order (and enhance inter-operability) it is recommended to use the Hill system order Hill (1900, 2012).

<table>
<thead>
<tr>
<th>H2O4S</th>
<th>C2H5Br</th>
<th>BrH</th>
</tr>
</thead>
<tbody>
<tr>
<td>C10H12N5O13P3</td>
<td>CH3I</td>
<td></td>
</tr>
</tbody>
</table>

Table 1: Examples of chemical formulas written using the Hill System. As described in Section 3.4
Using this notation the number of carbon atoms in a molecule is indicated first, followed by the number of hydrogen atoms and then the number of all other chemical elements in alphabetical order. When the formula contains no carbon; all elements, including hydrogen, are listed alphabetically.

### 3.5 The FBC FluxBound class

**FluxBound** is a new FBC class derived from SBML `SBase` that inherits `metaid` and `sboTerm`, as well as the subcomponents for `Annotation` and `Notes`. The purpose of this class is to hold a single (in)equality that provides the maximum or minimum value that a reaction flux can obtain at steady state. It implements four attributes.

**The id and name attributes**

A FluxBound has two optional attributes: `id` an attribute of type `SId` and `name` an attribute of type `string`.

**The reaction attribute**

The required `reaction` attribute of type `SIdRef`. This attribute must refer to a `Reaction` element defined within the enclosing model.

**The operation attribute**

The `operation` attribute contains a value of type `FbcOperation` that can take a limited set of boolean operators as defined in Section 3.2.2. The `operation` attribute represents a mathematical (in)equality of the form `reaction < operator > value` e.g. $R_3 \geq 0$, $R_5 \leq \infty$ and $R_7 = 1.0$. The mapping between traditional mathematical symbols and `FbcOperation` values is as follows:

- $\leq \leftrightarrow \text{"lessEqual"}$
- $\geq \leftrightarrow \text{"greaterEqual"}$
- $= \leftrightarrow \text{"equal"}$

**The value attribute**

The `value` attribute holds a `double` value representing the numerical value of the flux bound. This may include an explicitly defined $\pm \infty$ encoded as a value, e.g. "INF".

**Encoding the FluxBound**

As described in Section 3.5 the flux bound represents a mathematical (in)equality of the form `<reaction> <operator> <value>`. In SBML Level 3 Version 1 with FBC this is encoded as:

```xml
<fbc:listOfFluxBounds>
  <fbc:fluxBound fbc:id="R1b" fbc:reaction="R1" fbc:operation="greaterEqual" fbc:value="1.2"/>
  <fbc:fluxBound fbc:id="R2b" fbc:reaction="R2" fbc:operation="lessEqual" fbc:value="-1.2"/>
  <fbc:fluxBound fbc:id="R3b" fbc:reaction="R3" fbc:operation="equal" fbc:value="1"/>
</fbc:listOfFluxBounds>
```

For an example of how the FluxBound relates to the description of the underlying mathematical model please see Section 4.1.2.

**Units**

The `value` defined by the FluxBound has the units of the `reaction` that it refers to i.e. the globally defined unit of “extent per time.”

Consistency of flux bounds

It is possible, and in some cases necessary, to declare more than one FluxBound relating to a particular reaction. To allow as much flexibility to the user as possible, there is no restriction on the number of flux bounds and the operation specified by each that may be declared for a given reaction. However, in the case where multiple flux bounds are declared for one reaction the combined set must not produce inconsistent values for the upper and lower bounds of the constraint.

For example:

```xml
<fluxBound id="b1" reaction="R" operation="greaterEqual" value="3"/>
<fluxBound id="b2" reaction="R" operation="equal" value="2"/>
```

is obviously inconsistent since it defines \( R \geq 3 \) and \( R = 2 \) - which cannot be correct. However

```xml
<fluxBound id="b1" reaction="R" operation="greaterEqual" value="3"/>
<fluxBound id="b2" reaction="R" operation="equal" value="3"/>
```

would be fine since, whilst the information is essentially repeated, \( R \geq 3 \) and \( R = 3 \) produce the consistent result that \( R = 3 \).

Reactions with undefined flux bounds

In the spirit of SBML Level 3 Version 1 Core the Flux Balance Constraints package does not define any default values for any element. However, in the case of a reaction with no defined flux bounds it is possible to infer this information from the reaction reversibility. In this case: irreversible reactions should be considered to be positive, \( 0 \leq J \leq \infty \) and reversible ones free/unbound, \( -\infty \leq J \leq \infty \).

Similarly, there is also the potential for a bound to “seemingly” conflict with the reaction that it bounds’ reversibility, e.g. a reaction is irreversible but has bounds \( -\infty \leq J \leq \infty \). In the context of this package, flux bounds should be considered authoritative. This follows from the fact that a FluxBound can enforce an irreversible reaction, by restricting the flux \( 0 \leq J \leq \infty \), as well as a reversible reaction \( -\infty \leq J \leq \infty \). It is left to the software implementation to deal with any obvious inconsistencies.
3.6 The FBC Objective class

The FBC Objective class is derived from SBML SBase and inherits metaid and sboTerm, as well as the subcomponents for Annotation and Notes. An integral component in a complete description of a steady-state model is the so-called ‘objective function’ which generally consist of a linear combination of model variables (fluxes) and a sense (direction). In the FBC package this concept is succinctly captured in the Objective class.

The id and name attributes

An Objective has a required attribute id of type SId and an optional attribute name of type string.

The type attribute

The required type attribute contains an FbcType type which represents the sense of the optimality constraint and can take one of two values:

- maximize → “maximize”
- minimize → “minimize”

The listOfFluxObjectives element

The element listOfFluxObjectives which contains a ListOfFluxObjectives is derived from and functions like a typical SBML ListOf class with the restriction that it must contain one or more elements of type FluxObjective (see Section 3.7). This implies that if an Objective is defined there should be at least one FluxObjective contained in a ListOfFluxObjectives.

![Objective Class Diagram](image)

Figure 4: A UML representation of the Flux Balance Constraints package Objective class. See Section 1.4 for conventions related to this figure.

Encoding the Objective

The Flux Balance Constraints package allows for the definition of multiple model objectives with one being designated as active (see Section 3.6) as illustrated in this example:

```xml
<fbc:listOfObjectives fbc:activeObjective="obj1">
  <fbc:objective fbc:id="obj1" fbc:type="maximize">
    <fbc:listOfFluxObjectives>
      <fbc:fluxObjective fbc:reaction="R101" fbc:coefficient="1"/>
    </fbc:listOfFluxObjectives>
  </fbc:objective>
  <fbc:objective fbc:id="obj2" fbc:type="minimize">
    <fbc:listOfFluxObjectives>
      <fbc:fluxObjective fbc:reaction="R102" fbc:coefficient="-2.5"/>
      <fbc:fluxObjective fbc:reaction="R103" fbc:coefficient="1"/>
    </fbc:listOfFluxObjectives>
  </fbc:objective>
</fbc:listOfObjectives>
```

Note how both Objective instances differ in type and each contains different set of FluxObjectives (see Section 3.7). For an example of how the Objective relates to the description of the underlying mathematical model please see Section 4.1.3.
3.7 The FBC FluxObjective class

The FBC FluxObjective class is derived from SBML SBase and inherits metaid and sboTerm, as well as the sub-components for Annotation and Notes.

The FluxObjective class is a relatively simple container for a model variable weighted by a signed linear coefficient.

**The id and name attributes**

A FluxObjective has two optional attributes: id an attribute of type SId and name an attribute of type string.

**The reaction and coefficient attributes**

The required reaction is of type SIdRef and is restricted to refer only to a Reaction while the coefficient attribute holds a double referring to the coefficient that this FluxObjective takes in the enclosing Objective. For example the objective Maximize: $1 \cdot R_1 + 2 \cdot R_2$ would be encoded as

```xml
<objective fbc:id="obj1" fbc:type="maximize">
  <listOfFluxObjectives>
    <fluxObjective fbc:reaction="R1" fbc:coefficient="1"/>
    <fluxObjective fbc:reaction="R2" fbc:coefficient="2"/>
  </listOfFluxObjectives>
</objective>
```

**Figure 5:** A UML representation of the Flux Balance Constraints package FluxObjective class. See Section 1.4 for conventions related to this figure.

**Units**

As described above the FluxObjective defined here as $n \cdot J$ where the coefficient $(n)$ is dimensionless and the value $(J)$ takes the units of the reaction flux i.e. "extent per time". Therefore, the FluxObjective $(n \cdot J)$ has the unit "extent per time" where the units of reaction "extent" and "time" are defined globally.
4 Illustrative examples of the FBC syntax

This section contains a worked example showing the encoding of a model suitable for Flux Balance Analysis using the Flux Balance Constraints package.

4.1 Example one: the basic FBC syntax

4.1.1 Kinetic model description

Figure 6: FBC syntax example: a simple four reaction pathway. The reactions are R1, R2, X1, X2 with fixed species IN, OUT, ATP, NADH and variable species A, B.

As shown in Figure 6 this example is a simple four reaction pathway that transforms metabolite IN to OUT. The model was created and analyzed using the SBW Flux Balance FBC implementation Bergmann (2012); Bergmann and Sauro (2006). In SBML each reaction is represented as a chemical process transforming reactants to products, e.g. reaction R1 is encoded in XML as (see also the complete example provided at the end of this section):

```xml
<reaction id="R1" reversible="false" fast="false">
  <listOfReactants>
    <speciesReference species="IN" stoichiometry="1" constant="true"/>
  </listOfReactants>
  <listOfProducts>
    <speciesReference species="A" stoichiometry="1" constant="true"/>
  </listOfProducts>
</reaction>
```

Using the reagent identity and stoichiometry it is possible to compactly describe this network in terms of its reaction stoichiometry as shown in Table 2 where each reaction is represented as a column.

```
<table>
<thead>
<tr>
<th></th>
<th>R1</th>
<th>R2</th>
<th>X1</th>
<th>X2</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>1</td>
<td>0</td>
<td>-1</td>
<td>-1</td>
</tr>
<tr>
<td>B</td>
<td>0</td>
<td>-1</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>
```

Table 2: Example one: stoichiometric matrix, N

While the stoichiometry contains the structural properties of the reaction network the full description of a biological model can be described as a set of ordinary differential equations (ODEs). Of course other formalisms do exist,
but here we will concentrate exclusively on kinetic models where the change in concentration of each variable component in the system ($\frac{ds}{dt}$) is a non-linear function of the rates of the reactions which either create or consume it (the product of the stoichiometric matrix, $N$ and the vector of reaction rates, $v$).

$$\frac{ds}{dt} = Nv$$  \hspace{1cm} (1)

The formulation of the kinetic model, as shown in Equation 1 is typical of the kind that can already be described using SBML Level 3 Version 1 Core where the vector $v$ would contain rate equations as a function of parameters and variable species. In a steady-state, constraint based model these rates are considered unknowns and the system of equations can be rewritten as a set of linear constraints (see Equation 2):

$$NJ = 0$$  \hspace{1cm} (2)

Note that the rate vector $v$ is now represented as the steady-state flux vector $J$. However, in order to perform a typical steady-state analysis such as flux balance analysis (FBA) we need to include more information into the model description. SBML Level 3 Version 1 Core does not have an unambiguous way of encoding either a capacity constraint or an objective target and for this we need to use the additional constructs provided by the Flux Balance Constraints package. In the following sections the same model data is shown encoded as XML and as a Linear Program (LP) in the common format used by IBM CPLEX.

### 4.1.2 Capacity constraints

A capacity constraint: in this example the maximum limit (upper bound) of the flux through reaction $R1$ is set to be one (with an arbitrary unit of flux). In LP format this can be written as:

<table>
<thead>
<tr>
<th>Bounds</th>
</tr>
</thead>
<tbody>
<tr>
<td>$R1 \leq 1.0$</td>
</tr>
</tbody>
</table>

the same information encoded as XML:

```
<fbcl:fluxBounds>
  <fbcl:fluxBound fbc:reaction="R1" fbc:operation="lessEqual" fbc:value="1"/>
</fbcl:fluxBounds>
```

### 4.1.3 Objective function

This described a target which can be maximized or minimized: in this example the flux through reaction $R2$ will be maximized.

<table>
<thead>
<tr>
<th>Maximize</th>
</tr>
</thead>
<tbody>
<tr>
<td>objective1_objf: + 1.0 $R2$</td>
</tr>
</tbody>
</table>

the same information encoded as XML:

```
<fbcl:listOfObjectives fbc:activeObjective="objective1">
  <fbcl:objective fbc:id="objective1" fbc:type="maximize">
    <fbcl:listOfFluxObjectives>
      <fbcl:fluxObjective fbc:reaction="R2" fbc:coefficient="1"/>
    </fbcl:listOfFluxObjectives>
  </fbcl:objective>
</fbcl:listOfObjectives>
4.1.4 Complete worked example

To conclude we show how the complete model described in Figure 6 encoded as both an LP and as XML. Formulated as an LP the problem can be written as:

```
\% Example one LP format
Maximize
  objective1_objf: + 1.0 R2
Subject To
  A: + R1 - X1 - X2 = 0.0
  B: - R2 + X1 + X2 = 0.0
Bounds
  R1 <= 1.0
  0.0 <= R2 <= +inf
  0.0 <= X1 <= +inf
  0.0 <= X2 <= +inf
END
```

Solving this we find that maximization of flux through $R_2$ gives an optimal solution $R_2 = 1$, shown in Equation 3, with one possible solution for $J$.

\[
\begin{pmatrix}
  1 & 0 & -1 & -1 \\
  0 & -1 & 1 & 1 \\
\end{pmatrix}
\begin{pmatrix}
  1.0 \\
  1.0 \\
  0.0 \\
  1.0 \\
\end{pmatrix} = 0
\tag{3}
\]

Finally we provide the complete model, described above, encoded using the Flux Balance Constraints package:
Section 4.1 Example one: the basic FBC syntax

```xml
<model>
  <species id="B" compartment="compartment" initialConcentration="0" substanceUnits="substance" hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false"/>
  <species id="ATP" compartment="compartment" initialConcentration="0" substanceUnits="substance" hasOnlySubstanceUnits="false" boundaryCondition="false" constant="false"/>
  <species id="NADH" compartment="compartment" initialConcentration="0" substanceUnits="substance" hasOnlySubstanceUnits="false" boundaryCondition="true" constant="false"/>
</listOfSpecies>

<listOfReactions>
  <reaction id="R1" reversible="false" fast="false">
    <listOfReactants>
      <speciesReference species="IN" stoichiometry="1" constant="true"/>
    </listOfReactants>
    <listOfProducts>
      <speciesReference species="A" stoichiometry="1" constant="true"/>
    </listOfProducts>
  </reaction>
  <reaction id="R2" reversible="false" fast="false">
    <listOfReactants>
      <speciesReference species="B" stoichiometry="1" constant="true"/>
    </listOfReactants>
    <listOfProducts>
      <speciesReference species="OUT" stoichiometry="1" constant="true"/>
    </listOfProducts>
  </reaction>
  <reaction id="X1" reversible="false" fast="false">
    <listOfReactants>
      <speciesReference species="A" stoichiometry="1" constant="true"/>
    </listOfReactants>
    <listOfProducts>
      <speciesReference species="ATP" stoichiometry="1" constant="true"/>
      <speciesReference species="B" stoichiometry="1" constant="true"/>
    </listOfProducts>
  </reaction>
  <reaction id="X2" reversible="false" fast="false">
    <listOfReactants>
      <speciesReference species="A" stoichiometry="1" constant="true"/>
    </listOfReactants>
    <listOfProducts>
      <speciesReference species="B" stoichiometry="1" constant="true"/>
      <speciesReference species="NADH" stoichiometry="1" constant="true"/>
    </listOfProducts>
  </reaction>
</listOfReactions>

<fbc:listOfFluxBounds>
  <fbc:fluxBound fbc:reaction="R1" fbc:operation="greaterEqual" fbc:value="0"/>
  <fbc:fluxBound fbc:reaction="R1" fbc:operation="lessEqual" fbc:value="1"/>
  <fbc:fluxBound fbc:reaction="R2" fbc:operation="greaterEqual" fbc:value="0"/>
  <fbc:fluxBound fbc:reaction="R2" fbc:operation="lessEqual" fbc:value="INF"/>
  <fbc:fluxBound fbc:reaction="X1" fbc:operation="greaterEqual" fbc:value="0"/>
  <fbc:fluxBound fbc:reaction="X1" fbc:operation="lessEqual" fbc:value="INF"/>
  <fbc:fluxBound fbc:reaction="X2" fbc:operation="greaterEqual" fbc:value="0"/>
  <fbc:fluxBound fbc:reaction="X2" fbc:operation="lessEqual" fbc:value="INF"/>
</fbc:listOfFluxBounds>

<fbc:listOfObjectives fbc:activeObjective="objective1">
  <fbc:objective fbc:id="objective1" fbc:type="maximize">
    <fbc:fluxObjective fbc:reaction="R1" fbc:coefficient="1"/>
  </fbc:objective>
  <fbc:objective fbc:reaction="R2" fbc:coefficient="1"/>
</fbc:listOfObjectives>
</model>
```
5 Best practices

In this section, we recommend a number of practices for using and interpreting various constructs in the Flux Balance Constraints package. These recommendations are non-normative, but we advocate them strongly; ignoring them will not render a model invalid, but may reduce inter-operability between software and models.

5.1 Examples contrasting the current SBML L2 encoding with L3 and FBC

These examples contrast some elements of an existing model, iJR904 from the BiGG Database encoded in the COBRA format Becker et al. (2007); Reed et al. (2003); Schellenberger et al. (2010) that have been translated into SBML Level 3 Version 1 using the CBMPy implementation of the FBC package Olivier (2012); Olivier et al. (2005) and libSBML experimental ver. 5.6.0 Bornstein et al. (2008).

Objective function definition

SBML Level 2 objective function

```xml
<reaction id="R_BiomassEcoli" name="BiomassEcoli" reversible="false">
  <kineticLaw>
    <math xmlns="http://www.w3.org/1998/Math/MathML">
      <ci>FLUX_VALUE</ci>
    </math>
    <listOfParameters>
      <parameter id="LOWER_BOUND" value="0" units="mmol_per_gDW_per_hr"/>
      <parameter id="UPPER_BOUND" value="999999" units="mmol_per_gDW_per_hr"/>
      <parameter id="OBJECTIVE_COEFFICIENT" value="1"/>
      <parameter id="FLUX_VALUE" value="0" units="mmol_per_gDW_per_hr"/>
    </listOfParameters>
  </kineticLaw>
</reaction>
```

The SBML Level 3 objective function

```xml
<fbc:listOfObjectives fbc:activeObjective="obj1">
  <fbc:objective fbc:id="obj1" fbc:type="maximize">
    <fbc:listOfFluxObjectives>
      <fbc:fluxObjective fbc:reaction="R_BiomassEcoli" fbc:coefficient="1"/>
    </fbc:listOfFluxObjectives>
  </fbc:objective>
</fbc:listOfObjectives>
```

Species definition

It is particularly useful to contrast the differences in the Species definition as it is used in genome scale models.

SBML Level 2 Species annotation version 1

To begin with we let's examine the SBML Level 2 Version 1 species definition used by the BiGG database and COBRA Becker et al. (2007); Schellenberger et al. (2010). Note how the name attribute is overloaded with the chemical formula.

```xml
<species id="M_atp_c" name="ATP_C10H12N5O13P3" compartment="Cytosol" charge="-4"/>
```
**Section 5.1 Examples contrasting the current SBML L2 encoding with L3 and FBC**

**SBML Level 2 Species annotation version 2**

A newer variation of the above, probably necessitated by the discontinuation of the `charge` attribute in SBML and libSBML.

```xml
<species id="M_atp_c" name="ATP" compartment="c">
  <notes>
    <body xmlns="http://www.w3.org/1999/xhtml">
      <p>FORMULA: C10H12N5O13P3</p>
      <p>CHARGE: -4</p>
    </body>
  </notes>
</species>
```

**The SBML Level 3 Species attributes**

Hopefully, with the adoption of SBML FBC these species properties can be unified into a common format.

```xml
<species metaid="meta_M_atp_c" id="M_atp_c" name="ATP" compartment="Cytosol" boundaryCondition="false" initialConcentration="0" hasOnlySubstanceUnits="false" fbc:charge="-4" fbc:chemicalFormula="C10H12N5O13P3"/>
```

**Reaction definition and flux bounds**

**SBML Level 2 Reaction**

```xml
<reaction id="R_GTHS" name="glutathione synthetase" reversible="false">
  <notes>
    <html:p>Abbreviation: R_GTHS</html:p>
    <html:p>EC Number: 6.3.2.3</html:p>
    <html:p>SUBSYSTEM: Cofactor and Prosthetic Group Biosynthesis</html:p>
    <html:p>Equation: [c] : atp + glucys + gly --&gt; adp + gthrd + h + pi</html:p>
    <html:p>Confidence Level: 0</html:p>
    <html:p>LOCUS:b2947#ABBREVIATION:gshB#ECNUMBERS:6.3.2.3#</html:p>
    <html:p>NAME: glutathione synthase#ABBREVIATION:GshB#</html:p>
    <html:p>GENE ASSOCIATION: (b2947)</html:p>
  </notes>
  <listOfReactants>
    <speciesReference species="M_atp_c" stoichiometry="1"/>
    <speciesReference species="M_glucys_c" stoichiometry="1"/>
    <speciesReference species="M_gly_c" stoichiometry="1"/>
  </listOfReactants>
  <listOfProducts>
    <speciesReference species="M_adp_c" stoichiometry="1"/>
    <speciesReference species="M_gthrd_c" stoichiometry="1"/>
    <speciesReference species="M_h_c" stoichiometry="1"/>
    <speciesReference species="M_pi_c" stoichiometry="1"/>
  </listOfProducts>
  <kineticLaw>
    <math xmlns="http://www.w3.org/1998/Math/MathML">
      <ci>FLUX_VALUE</ci>
    </math>
    <listOfParameters>
      <parameter id="LOWER_BOUND" value="0" units="mmol_per_gDW_per_hr"/>
      <parameter id="UPPER_BOUND" value="999999" units="mmol_per_gDW_per_hr"/>
      <parameter id="OBJECTIVE_COEFFICIENT" value="0"/>
      <parameter id="FLUX_VALUE" value="0" units="mmol_per_gDW_per_hr"/>
    </listOfParameters>
  </kineticLaw>
</reaction>
```
The SBML Level 3 Reaction and FluxBound

As an example and where (unambiguously) possible the SBML Level 2 annotation has been converted into MIRIAM compliant RDF; in this case the EC number. In addition this example highlights an open issue, namely, how to deal with the information currently encoded in the Notes element. Currently this information is not completely encodable using the current SBML annotation mechanism, however, for an example of an alternative mechanism, implemented as a tool specific annotation please see Section 6.

```xml
<reaction metaid="meta_R_GTHS" id="R_GTHS" name="glutathione synthetase" reversible="false">
  <annotation>
      <rdf:Description rdf:about="#meta_R_GTHS">
        <bqbiol:is>
          <rdf:Bag>
            <rdf:li rdf:resource="http://identifiers.org/ec-code/6.3.2.3"/>
          </rdf:Bag>
        </bqbiol:is>
      </rdf:Description>
    </rdf:RDF>
  </annotation>

  <listOfReactants>
    <speciesReference species="M_atp_c" stoichiometry="1"/>
    <speciesReference species="M_glucys_c" stoichiometry="1"/>
    <speciesReference species="M_gly_c" stoichiometry="1"/>
  </listOfReactants>

  <listOfProducts>
    <speciesReference species="M_adp_c" stoichiometry="1"/>
    <speciesReference species="M_gthrd_c" stoichiometry="1"/>
    <speciesReference species="M_h_c" stoichiometry="1"/>
    <speciesReference species="M_pi_c" stoichiometry="1"/>
  </listOfProducts>
</reaction>

<fbc:listOfFluxBounds>
  <fbc:fluxBound fbc:id="R_GTHS_lower_bnd" fbc:reaction="R_GTHS" fbc:operation="greaterEqual" fbc:value="0"/>
  <fbc:fluxBound fbc:id="R_GTHS_upper_bnd" fbc:reaction="R_GTHS" fbc:operation="lessEqual" fbc:value="999999"/>
</fbc:listOfFluxBounds>
```
6 Future development

In this section we highlight some open issues not addressed in this version of the FBC specification.

6.1 Model annotation

While the Flux Balance Constraints package addresses the basic constraint based model structure the effective annotation of individual model components is critical to its efficient (re)use. Here two examples of custom annotations are provided as an example of how this could be achieved.

6.1.1 Gene association annotation

An example of an annotation currently encoded in the Notes element that is not encodable using the SBML Annotation mechanism is the so called “Gene Association”.

In order to capture this information we propose the following mechanism which can be used to represent a logical expression containing “genes” in this case the final (transcribed/translated) protein product of a gene which forms a sub-component of a protein.

```xml
<fbc:listOfGeneAssociations>
  <fbc:geneAssociation fbc:id="gal" fbc:reaction="R_ACGAptspp">
    <fbc:or>
      <fbc:and>
        <fbc:gene>b2415</fbc:gene>
        <fbc:gene>b2417</fbc:gene>
        <fbc:gene>b2416</fbc:gene>
        <fbc:gene>b1101</fbc:gene>
      </fbc:and>
      <fbc:and>
        <fbc:gene>b2415</fbc:gene>
        <fbc:gene>b0679</fbc:gene>
        <fbc:gene>b2416</fbc:gene>
      </fbc:and>
    </fbc:or>
  </fbc:geneAssociation>
</fbc:listOfGeneAssociations>
```

6.1.2 A generic annotation mechanism

An annotation issue previously introduced in the example shown in Section 4 where a SBML Level 2 Reaction is encoded using the Flux Balance Constraints package.

SBML Level 2 Reaction

```xml
<reaction id="R_GTHS" name="glutathione_synthetase" reversible="false">
  <notes>
    <html:p>Abbreviation: R_GTHS</html:p>
    <html:p>EC Number: 6.3.2.3</html:p>
    <html:p>SUBSYSTEM: Cofactor and Prosthetic Group Biosynthesis</html:p>
    <html:p>Equation: [c] : atp + glucys + gly --&gt; adp + gthrd + h + pi</html:p>
    <html:p>Confidence Level: 0</html:p>
    <html:p>LOCUS:b2947#ABBREVIATION:gshB#ECNUMBERS:6.3.2.3#NAME:glutathione synthase#ABBREVIATION:GshB#GENE ASSOCIATION: (b2947)</html:p>
  </notes>
  <listOfReactants>
    <speciesReference species="M_atp_c" stoichiometry="1"/>
    <speciesReference species="M_glucys_c" stoichiometry="1"/>
    <speciesReference species="M_gly_c" stoichiometry="1"/>
  </listOfReactants>
</reaction>
```
SBML Level 3 Reaction

Note how that in order to maintain all the annotations encoded in the SBML Level 2 Reaction notes a custom annotation is introduced i.e. the KeyValueData class.

In general this example highlights the need for a community supported annotation mechanism for genome scale, constraint based models.

More information about this annotation is available at http://pysces.sourceforge.net/KeyValueData
Section 6.1 Model annotation

<speciesReference species="M_gthrd_c" stoichiometry="1"/>
<speciesReference species="M_h_c" stoichiometry="1"/>
<speciesReference species="M_pi_c" stoichiometry="1"/>
</listOfProducts>
</reaction>
A Validation of SBML documents

A.1 Validation and consistency rules

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

- A checked box indicates a requirement for SBML conformance. If a model does not follow this rule, it does not conform to the Flux Balance Constraints specification. (Mnemonic intention behind the choice of symbol: “This must be checked.”)

- A triangle indicates a recommendation for model consistency. If a model does not follow this rule, it is not considered strictly invalid as far as the Flux Balance Constraints specification is concerned; however, it indicates that the model contains a physical or conceptual inconsistency. (Mnemonic intention behind the choice of symbol: “This is a cause for warning.”)

- A star indicates a strong recommendation for good modeling practice. This rule is not strictly a matter of SBML encoding, but the recommendation comes from logical reasoning. As in the previous case, if a model does not follow this rule, it is not strictly considered an invalid SBML encoding. (Mnemonic intention behind the choice of symbol: “You’re a star if you heed this.”)

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

For convenience and brevity, we use the shorthand “fbc:x” to stand for an attribute or element name x in the namespace for the Flux Balance Constraints package, using the namespace prefix fbc. In reality, the prefix string may be different from the literal “fbc” used here (and indeed, it can be any valid XML namespace prefix that the modeler or software chooses). We use “fbc:x” because it is shorter than to write a full explanation everywhere we refer to an attribute or element in the Flux Balance Constraints package namespace.

General rules about this package

fbc-10101 ✓ To conform to the Flux Balance Constraints package specification for SBML Level 3 Version 1, an SBML document must declare the following XML namespace:

```
```

(References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.1 on page 6.)

fbc-10102 ✓ Wherever they appear in an SBML document, elements and attributes from the Flux Balance Constraints package must be declared either implicitly or explicitly to be in the XML namespace “http://www.sbml.org/sbml/level3/version1/fbc/version1”. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.1 on page 6.)

General rules about identifiers

fbc-10301 ✓ (Extends validation rule #10301 in the SBML Level 3 Version 1 Core specification.) Within a Model the values of the attributes id and fbc:id on every instance of the following classes of objects must be unique across the set of all id and fbc:id attribute values of all such objects in a model: the Model itself, plus all contained FunctionDefinition, Compartment, Species, Reaction, SpeciesReference, ModifierSpeciesReference, Event, and Parameter objects, plus...
the FluxBound, Objective and FluxObjective objects defined by the Flux Balance Constraints package. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.2 on page 6.)

The value of attribute fbc:id attribute must always conform to the syntax of the SBML data type SID. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.2 on page 6.)

**Rules for the extended SBML class**

- **fbc-20101** ✓ The value of attribute fbc:required on the SBML object must be of the data type boolean. (References: SBML Level 3 Version 1 Core, Section 4.1.2.)
- **fbc-20102** ✓ The value of attribute fbc:required on the SBML object must be set to “false”. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.1 on page 6.)

**Rules for extended Model object**

- **fbc-20201** ✓ There may be at most one instance of each of the following kinds of objects within a Model object using Flux Balance Constraints: ListOfFluxBounds and ListOfObjectives. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.3 on page 7.)
- **fbc-20202** ✓ The various ListOf__ subobjects with an Model object are optional, but if present, these container object must not be empty. Specifically, if any of the following classes of objects are present on the Model, it must not be empty: ListOfFluxBounds and ListOfObjectives. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.3 on page 7.)
- **fbc-20203** ✓ Apart from the general notes and annotation subobjects permitted on all SBML objects, a ListOfFluxBounds container object may only contain FluxBound objects. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.3 on page 7.)
- **fbc-20204** ✓ Apart from the general notes and annotation subobjects permitted on all SBML objects, a ListOfObjectives container object may only contain Objective objects. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.3 on page 7.)
- **fbc-20205** ✓ A ListOfFluxBounds object may have the optional attributes metaid and sboTerm defined by SBML Level 3 Core. No other attributes from the SBML Level 3 Core namespace or the Flux Balance Constraints namespace are permitted on a ListOfFluxBounds object. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.3 on page 7.)
- **fbc-20206** ✓ A ListOfObjectives object may have the optional attributes metaid and sboTerm defined by SBML Level 3 Core. Additionally the ListOfObjectives must contain the attribute activeObjective. No other attributes from the SBML Level 3 Core namespace or the Flux Balance Constraints namespace are permitted on a ListOfObjectives object. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.3 on page 7.)
- **fbc-20207** ✓ The value of attribute fbc:activeObjective on the ListOfObjectives object must be of the data type SIDRef. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.3.2 on page 8.)
The value of attribute \texttt{fbc:activeObjective} on the ListOfObjectives object must be the identifier of an existing \texttt{Objective}. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.3.2 on page 8.)

\textbf{Rules for extended Species object}

\textbf{fbc-20301} \checkmark A \texttt{SBML Species} object may have the optional attributes \texttt{fbc:charge} and \texttt{fbc:chemicalFormula}. No other attributes from the Flux Balance Constraints namespaces are permitted on a \texttt{Species}. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.4 on page 8)

\textbf{fbc-20302} \checkmark The value of attribute \texttt{fbc:charge} on the \texttt{SBML Species} object must be of the data type \texttt{integer}. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.4 on page 8).

\textbf{fbc-20303} \checkmark The value of attribute \texttt{fbc:chemicalFormula} on the \texttt{SBML Species} object must be set to a \texttt{string} consisting only of atomic names or user defined compounds and their occurrence. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.4 on page 8.)

\textbf{Rules for FluxBound object}

\textbf{fbc-20401} \checkmark A \texttt{FluxBound} object may have the optional SBML Level 3 Core attributes \texttt{metaid} and \texttt{sboTerm}. No other attributes from the SBML Level 3 Core namespace are permitted on a \texttt{FluxBound}. (References: SBML Level 3 Version 1 Core, Section 3.2.)

\textbf{fbc-20402} \checkmark A \texttt{FluxBound} object may have the optional SBML Level 3 Core subobjects for notes and annotations. No other elements from the SBML Level 3 Core namespace are permitted on a \texttt{FluxBound}. (References: SBML Level 3 Version 1 Core, Section 3.2.)

\textbf{fbc-20403} \checkmark A \texttt{FluxBound} object must have the required attributes \texttt{fbc:reaction}, \texttt{fbc:operation} and \texttt{fbc:value}, and may have the optional attributes \texttt{fbc:id} and \texttt{fbc:name}. No other attributes from the SBML Level 3 Flux Balance Constraints namespace are permitted on a \texttt{FluxBound} object. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.5 on page 9.)

\textbf{fbc-20404} \checkmark The attribute \texttt{fbc:reaction} of a \texttt{FluxBound} must be of the data type \texttt{SIdRef}. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.5 on page 9.)

\textbf{fbc-20405} \checkmark The attribute \texttt{fbc:name} of a \texttt{FluxBound} must be of the data type \texttt{string}. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.5 on page 9.)

\textbf{fbc-20406} \checkmark The attribute \texttt{fbc:operation} of a \texttt{FluxBound} must be of the data type \texttt{FbcOperation} and thus its value must be one of “lessEqual”, “greaterEqual” or “equal”. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.5 on page 9.)

\textbf{fbc-20407} \checkmark The attribute \texttt{fbc:value} of a \texttt{FluxBound} must be of the data type \texttt{double}. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.5 on page 9.)

\textbf{fbc-20408} \checkmark The value of the attribute \texttt{fbc:reaction} of a \texttt{FluxBound} object must be the identifier of an existing \texttt{Reaction} object defined in the enclosing \texttt{Model} object. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.5 on page 9.)

\textbf{fbc-20409} \checkmark The combined set of all \texttt{FluxBound}’s with identical values for \texttt{fbc:reaction} must be consistent. That is while it is possible to define a lower and an upper bound for a reaction, it is not possible to define multiple lower or upper bounds. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.5 on page 9.)
**Rules for Objective object**

**fbc-20501** ✓ A **Objective** object may have the optional SBML Level 3 Core attributes **metaid** and **sboTerm**. No other attributes from the SBML Level 3 Core namespace are permitted on a **Objective**. (References: SBML Level 3 Version 1 Core, Section 3.2.)

**fbc-20502** ✓ A **Objective** object may have the optional SBML Level 3 Core subobjects for notes and annotations. No other elements from the SBML Level 3 Core namespace are permitted on a **Objective**. (References: SBML Level 3 Version 1 Core, Section 3.2.)

**fbc-20503** ✓ A **Objective** object must have the required attributes **fbc:id** and **fbc:type** and may have the optional attribute **fbc:name**. No other attributes from the SBML Level 3 Flux Balance Constraints namespace are permitted on a **Objective** object. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.6 on page 11.)

**fbc-20504** ✓ The attribute **fbc:name** on a **Objective** must be of the data type **string**. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.6 on page 11.)

**fbc-20505** ✓ The attribute **fbc:type** on a **Objective** must be of the data type **FbcType** and thus its value must be one of “minimize” or “maximize”. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.6 on page 11.)

**fbc-20506** ✓ A **Objective** object must have one and only one instance of the **ListOfFluxObjectives** object. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.6 on page 11.)

**fbc-20507** ✓ The **ListOfFluxObjectives** subobject within a **Objective** object must not be empty. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.6 on page 11.)

**fbc-20508** ✓ Apart from the general notes and annotation subobjects permitted on all SBML objects, a **ListOfFluxObjectives** container object may only contain **FluxObjective** objects. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.6 on page 11.)

**fbc-20509** ✓ A **ListOfFluxObjectives** object may have the optional **metaid** and **sboTerm** defined by SBML Level 3 Core. No other attributes from the SBML Level 3 Core namespace or the Flux Balance Constraints namespace are permitted on a **ListOfFluxObjectives** object. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.6 on page 11.)

**Rules for FluxObjective object**

**fbc-20601** ✓ A **FluxObjective** object may have the optional SBML Level 3 Core attributes **metaid** and **sboTerm**. No other attributes from the SBML Level 3 Core namespace are permitted on a **FluxObjective**. (References: SBML Level 3 Version 1 Core, Section 3.2.)

**fbc-20602** ✓ A **FluxObjective** object may have the optional SBML Level 3 Core subobjects for notes and annotations. No other elements from the SBML Level 3 Core namespace are permitted on a **FluxObjective**. (References: SBML Level 3 Version 1 Core, Section 3.2.)

**fbc-20603** ✓ A **FluxObjective** object must have the required attributes **fbc:reaction** and **fbc:coefficient**, and may have the optional attributes **fbc:id** and **fbc:name**. No other attributes from the SBML Level 3 Flux Balance Constraints namespace are permitted on a **FluxObjective** object. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.7 on page 12.)
Section A.1 Validation and consistency rules

The attribute `fbc:name` on a `FluxObjective` must be of the data type `string`. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.7 on page 12.)

The value of the attribute `fbc:reaction` of a `FluxObjective` object must conform to the syntax of the SBML data type `SIdRef`. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.7 on page 12.)

The value of the attribute `fbc:reaction` of a `FluxObjective` object must be the identifier of an existing `Reaction` object defined in the enclosing `Model` object. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.7 on page 12.)

The value of the attribute `fbc:coefficient` of a `FluxObjective` object must conform to the syntax of the SBML data type `double`. (References: SBML Level 3 Package Specification for Flux Balance Constraints, Version 1, Section 3.7 on page 12.)
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