XML format for RBA models

S. Fischer, V. Fromion, A. Goelzer August 17, 2017

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1 Introduction

In this document we present the XML structures used to define a RBA model. A complete RBA model is composed of the following files:

- metabolism.xml (definition of compartments, metabolic species and metabolic reactions).
- parameters.xml (definition of density constraints and user-defined functions).
- proteins.xml (definition of proteins).
- rnas.xml (definition of RNAs).
- dna.xml (definition of DNA).
- enzymes.xml (definition of enzymatic machineries catalyzing metabolic reactions).
- processes.xml (definition of cell processes necessary to growth and maintenance).

For every file, we present the nodes that compose the XML structure. For every node, we show a class diagram that shows the node's attributes and the children node that it may/must contain. We provide a brief description about the relevance of the node in the RBA model.

2 Conventions

2.1 Naming conventions

In XML, classes follow ThisConvention, instances thisConvention. In the Python package, classes follow ThisConvention, instances this_convention. In the Python package, instance listOfThings simply becomes things. Throughout this document, XML conventions are used. Keep in mind that the convention for instances is different when using the Python package.

2.2 Boolean attributes

A boolean attribute evaluates to true if it is "1" or "true" (case does not matter). In all other cases it evaluates to false.

2.3 Variable for transport functions

A metabolite identifier is expected to look like M_met_c, where M is a generic prefix (identical for all metabolites), met is the name of the metabolite and c its compartment. The variable for a transport function (see EnzymaticActivity) must be the prefix of a metabolite, in this case M_met. Every time a transport function based on met is

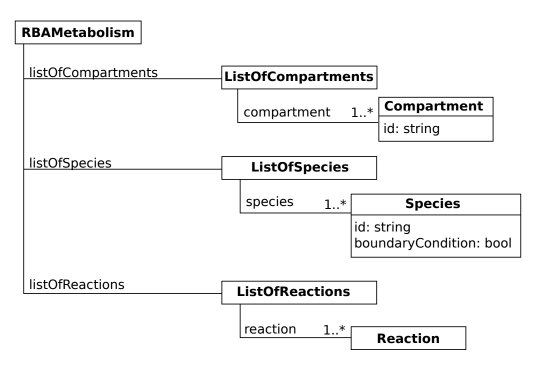


Figure 1: XML structure of metabolism document.

evaluated, the external concentration is used no matter where the transport takes place. Typically, glucose import rates from the periplasm to the cytosol will be based on the concentration of extracellular glucose.

3 metabolism.xml

The metabolism file is strongly inspired by SBML. More precisely, it is a subpart of an SBML file.

3.1 RBAMetabolism

The outermost portion of the metabolism file is an instance of class **RBAMetabolism**, shown in Figure 1.

Currently, **RBAMetabolism** has no simple attributes. It includes exactly one instance of **ListOf** container classes. All **ListOf** classes do not have own attributes, they are merely used to organize a list of instances from another class. This organization was inspired by SBML.

3.2 Compartment

The Compartment class is used to list existing cell compartments.

The *id* attribute The *id* attribute is a string defining the identifier of a compartment.

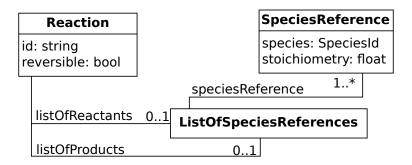


Figure 2: Class storing metabolic reactions.

3.3 Species

The **Species** class is used to define *metabolic* species.

The *id* attribute is a string defining the identifier of a metabolite.

The boundaryCondition attribute The boundaryCondition attribute is a boolean. If the attribute is set to true, the metabolite is considered to be at a constant concentration. In other words, it is not affected by reactions. This is typical for metabolites in the external medium.

3.4 Reaction

The **Reaction** class is used to define metabolic reactions (Fig. 2). Reactants and products are defined using a **ListOfSpeciesReferences**.

The *id* attribute The *id* attribute is a string defining the identifier of a reaction.

The reversible attribute The reversible attribute is a boolean. If the attribute is set to true, the reaction can occur in both directions. If the attribute is set to false, only the forward reaction can occur.

3.5 SpeciesReference

The **SpeciesReference** class is used to refer to a metabolic **Species** and associate with it a stoichiometry (Fig. 2).

The species attribute The species attribute must match the identifier of a Species.

The *stoichiometry* attribute The *stoichiometry* is a positive real number. It repensents the stoichiometry of a **Species** in a given context (typically a **Reaction**).

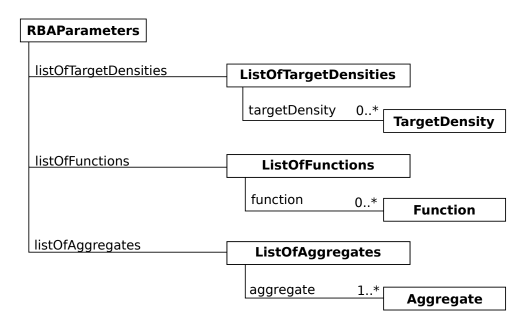


Figure 3: XML structure of parameter document.

4 parameters.xml

The parameter file contains density constraints for the RBA model, user-defined parameters and functions.

4.1 RBAParameters

The outermost portion of the parameter file is an instance of class **RBAParameters**, shown in Figure 3.

RBAParameters has no simple attributes. It includes exactly one instance of **ListOf** container classes. All **ListOf** classes do not have own attributes, they are merely used to organize a list of instances from another class.

4.2 TargetDensity

The **TargetDensity** class is used to define density constraints (Fig. 4). In a RBA model, a density constraint defines how many molecules a given compartment can contain. It inherits **TargetValue** for the constraint definition part.

The *compartment* attribute The *compartment* attribute must match the identifier of a Compartment.

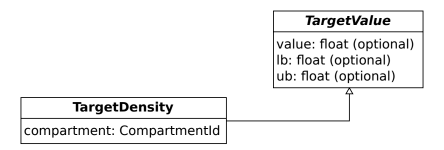


Figure 4: Class used to store density constraints.

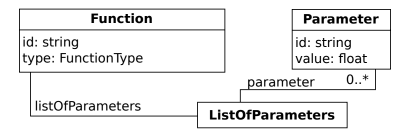


Figure 5: Class used to store user-defined functions.

4.3 TargetValue

The **TargetValue** class is used to define the sign of an additional RBA constraint and the value of its second member (Fig. 4). It is designed to be inherited. The child class usually holds information about the first member of the constraint (e.g. compartment for a density constraint, metabolite for a production constraint).

The value, *lb* and *ub* attributes Every attribute can be left undefined, set to a real number or contain the identifier of a Function or an Aggregate. In the latter case, its value will be growth-rate dependent.

If **value** is defined, the constraint is an equally constraint. **lb** and **ub** are ignored. If **value** is undefined, **lb** (resp. **ub**) defines a lower bound (resp. upper bound) inequality constraint. Note that **lb** and **ub** may both be defined, yielding two separate inequality constraints.

4.4 Function

The **Function** class is used for user-defined functions and parameters (Fig. 5). Implictly, all functions defined here are functions of *growth rate*. Every function holds a **ListOfParameters**, where **Parameter** are defined according to each type of function.

The *id* attribute The *id* attribute is a string defining the identifier of the function.

The type attribute The **type** attribute is a string that must match a known function type. Currently, the supported types are:

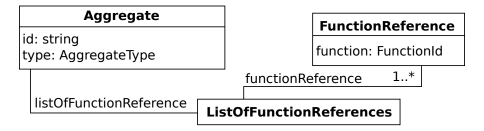


Figure 6: Class used to store user-defined aggregates.

- constant. Constant function with parameter CONSTANT.
- linear. Linear function with parameters LINEAR_CONSTANT, LINEAR_COEF, X_MIN, X_MAX, Y_MIN, Y_MAX. The 4 last parameters are used to saturate the function. If the variable (growth rate) is outside of the [X_MIN, X_MAX] range, it is set to the closest value in that range. The same applies for the return value and the [Y_MIN, Y_MAX] range.
- exponential. Exponential function with parameter *RATE*.
- indicator. Indicator function with parameters X_MIN and X_MAX . This function returns one if the variable (growth rate) is in the [X_MIN, X_MAX], zero otherwise.
- michaelisMenten. Michaelis Menten function with parameters kmax, Km and Y_MIN (optional). If Y_MIN is defined, any return value lower than Y_MIN will be set to Y_MIN.

4.5 Parameter

The **Parameter** class is used to store the values of function parameters (Fig. 5).

The *id* attribute The *id* attribute is a string that should match a valid parameter identifier. The list of valid parameters for each type of **Function** is listed above.

The value attribute is a real number representing the value of the attribute.

4.6 Aggregate

The **Aggregate** class is used to assemble user-defined functions (Fig. 6). Every aggregate holds a **ListOfFunctionReferences**, where each **FunctionReference** refers to a previously defined function.

The *id* attribute The *id* attribute is a string defining the identifier of the aggregate.

The *type* attribute The *type* attribute is a string that must match a known aggregate type. Currently, the supported types are:

• multiplication. The result is the multiplication of the values returned by the function listed in the aggregate at current growth rate.

4.7 FunctionReference

The **FunctionReference** class is used to refer to a user-defined **Function** (Fig. 6).

The function attribute The function attribute is a string that must match the identifier of a user-defined Function.

5 proteins.xml, rnas.xml and dna.xml

All these files use instances of the class **RBAMacromolecules**.

5.1 RBAMacromolecules

The outermost portion of the metabolism file is an instance of class **RBAMacro-molecules**, shown in Figure 7.

RBAMacromolecules has no simple attributes. It contains exactly one instance of ListOfComponents and ListOfMacromolecules.

5.2 Component

The **Component** class is used to define the components of a **Macromolecule** (Fig. 7). For example, these are expected to be amino acids, vitamins and ions for proteins. Note that they use separate identifiers as metabolic **Species**. **ComponentMaps** define how they are assembled from metabolites.

The *id* attribute The *id* attribute is a string defining the identifier of a component.

The weight attribute The weight attribute is a real number defining the weight of a component. This information is essential for the density constraints. The weight of a macromolecule is defined as the sum of the weight of its components. Note that units used for the weight constraint must be consistent across files.

The name and type attributes The name and type attributes are string that provide additional information about the component. The name is a standard name of the component (e.g. full amino acid name). The type can be used to distinguish components if necessary (e.g. in amino acids, vitamins, ions).

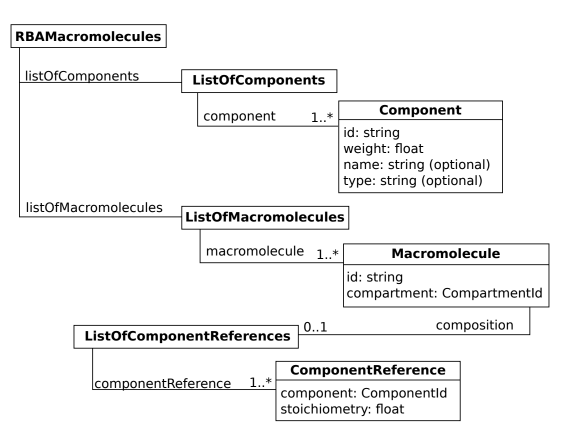


Figure 7: XML structure of macromolecule document.

5.3 Macromolecule

The **Macromolecule** class is used to define macromolecular species (Fig. 7). Its composition is given by a **ListOfComponentReferences**.

The *id* attribute The *id* attribute is a string defining the identifier of the macromolecule.

The compartment attribute The compartment attribute must match the identifier of a Compartment. It represents the compartment where the molecule is thought to be active.

5.4 ComponentReference

The **ComponentReference** class is used to refer to a **Component** and associate with it a stoichiometry (Fig. 7).

The *component* attribute The *component* attribute must match the identifier of a Component defined in the same RBAMacromolecules instance.

The stoichiometry attribute The stoichiometry is a positive real number. It repensents the stoichiometry of a Component in a given context (typically the composition of a Macromolecule).

6 processes.xml

The process file is used to define constraints related to complex cellular processes.

6.1 RBAProcesses

The outermost portion of the process file is an instance of class **RBAProcesses**, shown in Figure 8.

RBAMacromolecules has no simple attributes. It contains exactly one instance of **ListOfProcesses** and **ListOfComponentMaps**.

6.2 Process

The **Process** class is used to define cellular processes (Fig. 9). Processes cover a wide variety of non-metabolic reactions.

A **Process** revolves around 3 optional substructures. We encourage you to study existing examples to understand how the different substructures are used.

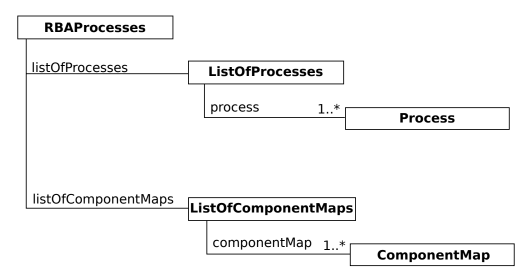


Figure 8: XML structure of process document.

The Machinery is the molecular entity enabling the process (e.g. ribosome for translation.) Each machinery unit has a limited production/degradation capacity. Every target of a process has an intrinsic metabolite cost (metabolites needed to produce/degrade it and byproducts). However, if a machinery is defined, there is an additional metabolic cost to produce the machinery that will enable the production/degradation of the target. This is similar to the production of Enzymes in order to catalyze metabolic Reactions.

Operations define the sets of macromolecules that a process produces or degrades. A Macromolecule can only be produced if it was related to a Process. For example, a protein can only be produced if there is a translation process where proteins are explicitly listed as a production in Operations. Operations break down Macromolecules in metabolic Species and Machinery costs. Operations does not define how many molecules of a Macromolecule must be produced.

Targets define the fluxes that a process must maintain for the cell to work properly. Some fluxes, such as enzyme production, are already enclosed in the base model. They should not appear in Targets. On the other hand, fluxes such as production of housekeeping proteins need to be explicitly added to Targets.

The *id* attribute The *id* attribute is a string defining the identifier of a process.

The name attribute The name attribute is a string that can be used to give the process a more human understandable name.

6.3 Machinery

The Machinery class defines the machinery used by a process (Fig. 9). Machinery has no simple attributes. If a Machinery is defined, it defines a capacity constraint. Every Machinery unit has a metabolic cost defined by a Machinery Composition.

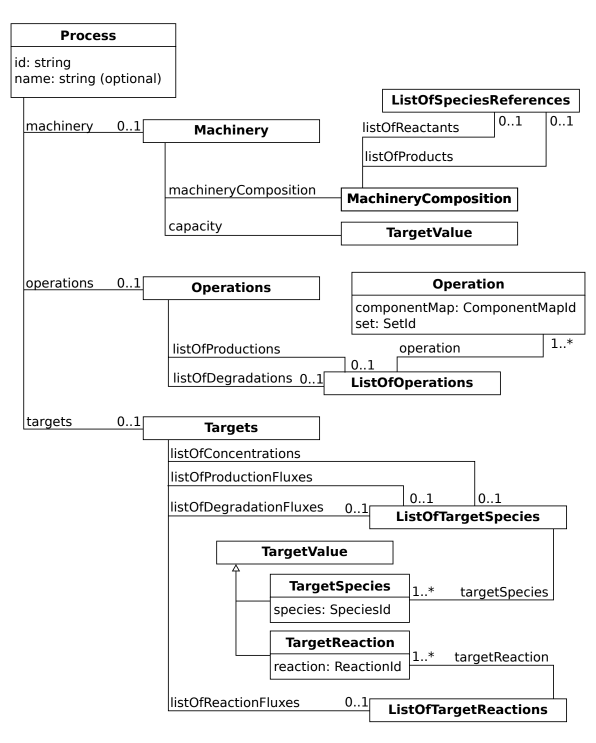


Figure 9: Class used to store processes.

Every unit also has a capacity defined by a **TargetValue**. The capacity defines how many targets a **Machinery** can process in 1 unit of time. Total capacity (base capacity multiplied by number of **Machinery** units) must always exceed the number of targets produced.

6.4 MachineryComposition

The MachineryComposition class defines the assembly costs of a complex molecular machinery (Fig. 9). MachineryComposition has no simple attributes. It contains two ListOfSpeciesReferences. One is for reactants, the other for byproducts of the assembly reaction. Note that in this case, SpeciesReferences can refer to both metabolic Species and Macromolecules. The assembly reaction should contain obvious components of the machinery, but also metabolic costs related to assembly (such as ATP/GTP costs) unless these costs are already covered by a process.

6.5 Operations

The Operations relates Macromolecules production/degradation to some given Process (Fig. 9). Operations has no simple attributes. It may contain two ListOfOperations, one for production and one for degradation.

6.6 Operation

The **Operations** class defines how **Macromolecules** are produced/degraded (Fig. 9). **Operation** is used to break down **Macromolecules** into metabolites by linking them to a **ComponentMap**.

The componentMap attribute The componentMap attribute must match the identifier of a ComponentMap. This ComponentMap will be used to compute the metabolic costs.

The set attribute The set attribute must refer to a Macromolecule set. Currently, the only acceptable values are protein, rna and dna. The set contains the Macromolecules that are being produced/degraded.

6.7 ComponentMap

The ComponentMap class is used to convert Macromolecules in metabolic and machinery costs (Fig.10).

There are two types of costs. The **ConstantCost** is a purely metabolic cost. It contains metabolites that are consumed/produced independent of the **Macromolecule**'s length (e.g. translation initiation). The **ListOfCosts** container details **Costs** depending on the individual **Components** of the **Macromolecule**. They cover metabolic costs used to assemble the **Component** onto the nascent **Macromolecule**. They also

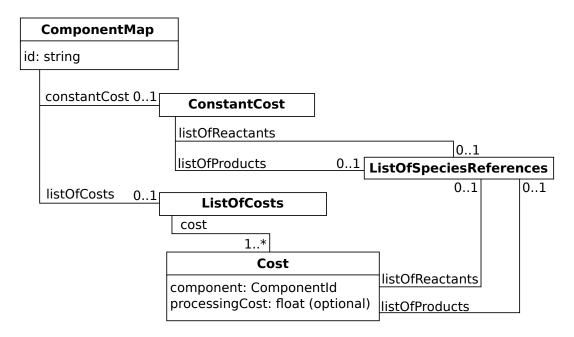


Figure 10: Class used to store production/degradation of macromolecules.

cover processing costs, *i.e.* how many **Machinery** units are needed to assemble the **Component**.

The *id* attribute The *id* attribute is a string defining the identifier of a component map.

6.8 ConstantCost

The **ConstantCost** class defines metabolites consumed and byproducts generated by an assembly process (Fig.10). It contains two **ListOfSpeciesReferences**, one for metabolites consumed and one for metabolites produced. Note that in this context, a **Species-Reference** must refer to a metabolic **Species**.

69 Cost

The **Cost** class defines metabolites consumed and byproducts generated when assembling a specific **Component** (Fig.10). It contains two **ListOfSpeciesReferences**, one for metabolites consumed and one for metabolites produced. Note that in this context, a **SpeciesReference** must refer to a metabolic **Species**. Additionally, it defines a processing cost used in a **Machinery**'s capacity constraint.

The *component* attribute The component attribute is a string that must match the identifier of a Component.

The processingCost attribute The processingCost attribute is a real value that is used to compute how many Machinery units are needed to assemble the Component. For example, let the processing cost of an amino acid be 1. The capacity of the Machinery (the ribosome) is the number of amino acids it can assemble per unit of time. The processing cost allows to compute how many ribosomes are needed to produce the Component and, in the end, the Macromolecule (in this example the number of amino acids divided by the ribosome's capacity).

6.10 Targets

The **Targets** class defines what molecules a **Process** must produce for the cell to work properly (Fig. 9). **Targets** has no simple attributes.

It contains 3 **ListOfTargetSpecies**. These targets allow to define metabolic **Species** or **Macromolecule** fluxes. One is for production fluxes, another for degradation fluxes. The last list is for maintaining a target at a given concentration. The difference with a simple production flux is that keeping a target at a concentration depends on growth rate. More precisely, the flux needed to keep the concentration is the growth rate multiplied by the target concentration. Note that all fluxes must be positive. If the target is a **Macromolecule**, production/degradation can only occur if an **Operations** section of some **Process** defines how the **Macromolecule** is actually produced/degraded.

It contains a **ListOfTargetReactions**. It is also possible to define target fluxes as reaction fluxes. These targets add constraints on the flux of a specific metabolic **Reaction**. In this case, fluxes may be positive or negative.

6.11 TargetSpecies

The **TargetSpecies** class defines constraints for a species flux (Fig. 9). It inherits **TargetValue** for the constraint definition part, allowing for equality or inequality constraints.

The species attribute The species attribute is a string that must match the identifier of a metabolic Species or a Macromolecule. Note that the Macromolecule must be broken down into metabolite costs through the Operations section of some Process. Otherwise no cost will be applied.

6.12 TargetReaction

The **TargetReaction** class defines constraints for a species flux (Fig. 9). It inherits **TargetValue** for the constraint definition part, allowing for equality or inequality constraints.

The reaction attribute The reaction attribute is a string that must match the identifier of a metabolic Reaction.

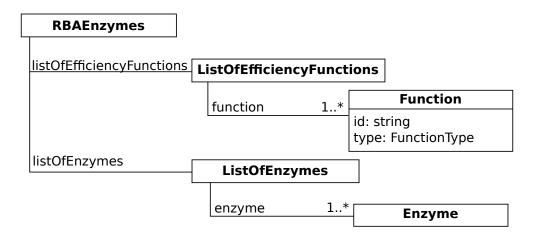


Figure 11: XML structure of enzyme document.

7 enzymes.xml

The enzyme file is used to define enzyme composition and efficiency. It defines efficiency constraints in the RBA model. These constraints ensure that a reaction flux is smaller than the product of efficiency and concentration of the enzyme catalyzing the reaction.

7.1 RBAEnzymes

The outermost portion of the metabolism file is an instance of class **RBAenzymes**, shown in Figure 11.

RBAenzymes has no simple attributes.

It contains exactly one instance of **ListOfEfficiencyFunctions**. This list contains **Function** definitions without a **ListOfParameter**. The variable of the **Functions** is the growth rate. These functions can be used to define different sets of efficiency functions. A classical application is to define one function per external medium. Suppose the function identifiers are **medium_1**, ..., **medium_N**. Every **Enzyme** must redefine these functions and associate enzyme-specific parameters. For example, for enzyme A, parameters for function **medium_1** will reflect the efficiency of A in the first medium. This structure allows to solve RBA for different sets of enzyme efficiencies.

It contains excatly one instance of **ListOfEnzymes** that is used to store **Enzyme** information.

7.2 Enzyme

The **Enzyme** class is used to define enzymes (Fig. 12).

It contains a Machinery Composition that refers to metabolic Species and Macromolecules composing the Enzyme. Note that the composition can be left unspecified. In this case, the reaction associated with the enzyme is considered spontaneous.

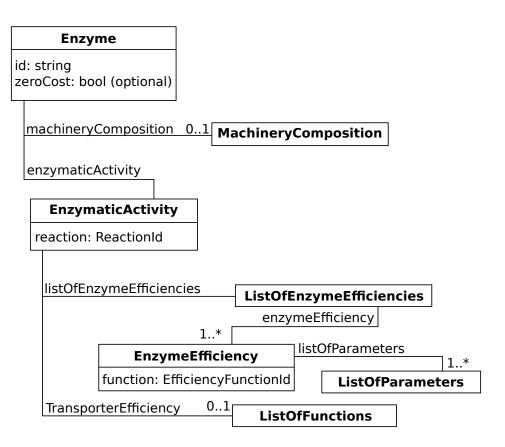


Figure 12: Class storing enzyme information.

It contains an **EnzymaticActivity** instance that is used to specify the various **EnzymeEfficiency** functions. It also contains transporter activity information (if applicable).

The *id* attribute The *id* attribute is a string defining the identifier of the enzyme.

The zeroCost attribute The zeroCost attribute is a boolean value. If set to true, the reaction associated may occur without having to produce the enzyme. If set to false or unspecified, an efficiency constraint is created where the flux through the reaction has to be smaller than the product of enzyme efficiency and enzyme concentration.

7.3 EnzymaticActivity

The **EnzymaticActivity** class is used to store information about enzyme efficiency and transporter efficiency (Fig. 12).

It contains one instance of **ListOfEnzymeEfficiencies**. This list redefines the efficiency functions in **RBAenzymes** with parameters specific to an enzyme.

It contains one instance of **ListOfFunctions** representing transporter efficiency. Each **Function** in the list must specify a **variable** attribute. The **variable** attribute is the metabolite whose external concentration affects the transport rate (usually the chemical imported or a cofactor). The transport efficiency is computed by *multiplying* the results of all the functions listed. Note that the transport efficiency can depend on an arbitrary number of chemicals.

The reaction attribute The reaction attribute must match the identifier of a metabolic Reaction. This is a one-to-one mapping. A Reaction can only have one associated Enzyme. If several Enzymes catalyze the same Reaction, the Reaction must be duplicated.

7.4 EnzymeEfficiency

The **EnzymeEfficiency** class is used to define parameters for an efficiency function (Fig. 12). It contains one instance of **ListOfParameters**. The **Parameters** must match the type of the efficiency function (as specified in the **RBAenzymes** instance) being redefined.

The function attribute The function attribute must match the identifier of an efficiency function defined in the same **RBAenzymes** instance.