# MyBacteria design Version 1.0

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# 1. Introduction

The aim of this document is go present the general design of MyBacteria's implementation (the code is documented using Doxygen, technical details are therefore best found in the Doxygen-generated manual).

Before we started working on MyBacteria, we set up a list of requirements that the simulator should fulfill. We wanted to be able to integrate MyBacteria in a whole-cell framework applicable to several organisms. Such a simulator should meet the following requirements.

- Simulate as many bacterial processes as possible.
- Generic formalism, applicable to any bacteria. Once a model has been built for some species, it should be easy to adapt it for another species by modifying input files only.
- Easy to use, inputs should be close to classical biochemical descriptions.
- Integrate various levels of description. The user should be able to focus on a process of interest with a very low level of description while keeping the remaining processes at a higher level.
- Efficient (not more than a couple of hours for one cell cycle).
- Easy to extend and reuse (either by extending the core or coupling with other modules).

We start by listing the central design choices of MyBacteria. Then we present the base components of the simulator, organized around reactants and reactions. The following section describes the same element again, but goes further into hypotheses and critical design elements. Finally, appendices are added to describe elements that have been important in the simulator development but did not fit naturally in the main document (testing strategies, utility classes, etc.).

# 2. Design principles

**Stochastic simulation** In order to meet the requirements listed above, we opted for a Gillespie-based simulator. The Gillespie algorithm has two important features in our context:

- It is a stochastic algorithm, so it naturally enables to simulate low-level stochastic processes.
- It offers a framework where an arbitrary number of reactions can be added.

Using the Gillespie algorithm, we can both simulate events at the molecular level and aggregated processes. The description level of a process simply depends on the number of reactions that the user has chosen to represent the process. MyBacteria starts with an empty system of reactions. The user controls what reactions to add. Processes can easily be tuned to match a specific bacterial species.

**Sequence-based reactions** Standard Gillespie simulators only implement chemical reactions. This does not meet our requirement of simulating a wide variety of processes. For example, it is extremely tedious (nearly impossible) to simulate translation accurately using only chemical reactions. Take a simple molecule of species A translocating along a sequence S of length 100. Here are three possible models for these translocation events:

Subscripts show the position of A along the sequence. Model A and B use only chemical reactions, both involve duplicating translocation events. In practice, this implies creating thousands of reactions and chemical species. In model A, the sequence S can only bind one chemical at a time, which is not realistic for biological systems (there is a high number of ribosomes per mRNA for example). Model B enables several molecules to "bind" the sequence at a given time. However, bound molecules do not interact with each other, e.g. it is impossible to handle sequestration of binding motifs. In MyBacteria, we define new types of reactants and reactions for sequence-based events. A single reaction and a single chemical species are used to represent all translocation events.

We created a variant of the Gillespie algorithm where new types of reactants and reactions can be plugged in. We defined a minimal set of reactants and reactions that handles sequence-based reactions (e.g. binding, translation elongation). All reactions remain low-level, enabling flexible descriptions of complex processes using a limited number of reactions. A lot of information that is provided as an input for these reactions comes from standard sequence annotation. When switching from an organism to another, the key reactions that define the process remain the same. Only sequence information (DNA, position of genes, promoters, etc.) and rates need to be adapted.

**Efficiency** We evaluated performance by simulating protein production. Protein production (from gene to protein) is responsible for a large number of reactions in a bacterial cell (metabolism aside). Simulation is completed within hours even for detailed descriptions of all processes involved (stochastic base-by-base elongation with all cofactors).

This objective was reached by using the latest implementations of the (exact) Gillespie algorithm. We also tuned all new types of reactions to be nearly as efficient as chemical reactions.

**Modularity** We created clear modules in MyBacteria's structure. This allows for core changes and facilitates communication with external modules. Typical core changes involve:

- Plugging in new solver variants (e.g. new implementation of the exact Gillespie algorithm, implementation of approximations such as  $\tau$ -leaping).
- Plugging in new reactants and reactions.

Interfaces of the modules were designed for these operations to be pure plug-in operations (no need to change the code in existing modules). A similar design applies for external programs. Reactant concentrations can be modified during the course of a simulation. This enables to plug-in arbitrary external solvers. For example, we intend to plug-in a deterministic solver for metabolism on MyBacteria.

# 3. Global presentation of the components of the simulator

# 3.1. Components of the simulator

The simulator can be decomposed into several large modules that handle specific tasks during simulation (Fig. 1). First of all, there is an **input/output** module that creates everything that is needed for the simulation from an input file. **Reactants** and **reactions** are user-specified and need to be created on demand, as well as **events** happening throughout the simulations and more technical aspects about which algorithm to use to perform the integration. Once everything is set up, the **solver** follows a simple loop that can be decomposed in three steps. Integration occurs reaction by reaction, at each loop, we go forward one reaction, update the simulation time, concentrations and reaction rates.

- 1. At the beginning of the loop, the **input/output** process checks whether **events** should occur at the current simulation time and whether it needs to write some concentrations to an output file.
- 2. It then hands control over to the **solver**, which is based on Gillespie's approach to integrate a network of chemical reactions. The Gillespie algorithm needs the current reaction rates of all **reactions** and draws a random reaction with a probability proportional to its rate. This task is delegated to a **rate manager**, which uses state-of-the-art methods to maintain the rate list updated and perform the drawing efficiently.

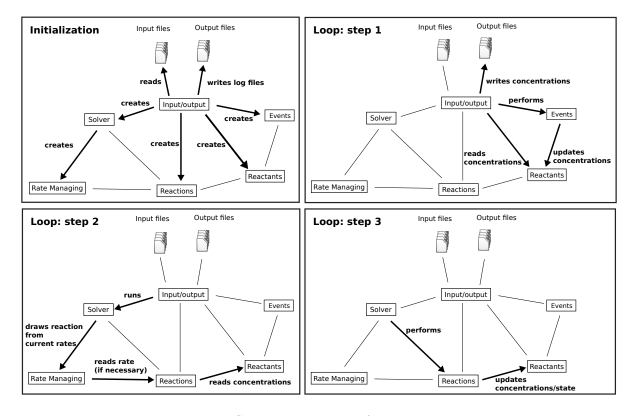


Figure 1: Schematical view of the simulator.

3. Once a **reaction** is drawn, it is performed *i.e.* the concentrations (and the state, see below) of its **reactants** is modified.

# 3.2. Reactant hierarchy

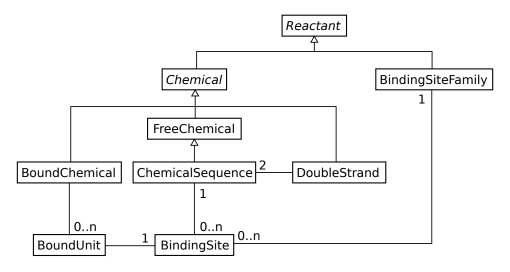


Figure 2: UML diagram of Reactant hierarchy

This section gives a quick overview of the contents of the Reactant hierarchy (Fig. 2). More details about how reactants are implemented can be found later.

# 3.2.1. Reactant

Reactant is a global abstract interface. All entities that can participate in a reaction must inherit from it.

# 3.2.2. Chemical

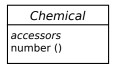


Figure 3: Chemical class

Chemical is an abstract class (Fig. 3). It defines all standard chemical entities. Chemical represents a *pool* of a given chemical species, meaning that one may access its current number at any time.

# 3.2.3. FreeChemical

# **Input format**

FreeChemical <name> [<initial quantity>]



Figure 4: FreeChemical class

FreeChemical (Fig. 4) is a subclass of Chemical that represents free chemical (e.g. molecules diffusing in the cytosol or extracellular medium).

# 3.2.4. BoundChemical

# Input format

BoundChemical <name>

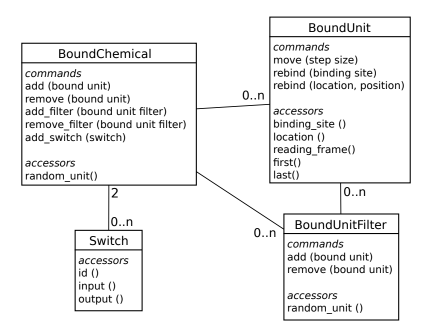


Figure 5: BoundChemical class

BoundChemical (Fig. 5) is a subclass of Chemical that represents chemicals that are bound to a sequence. Even though BoundChemical represents a pool of molecules, single elements are not interchangeable, they are defined by their position on a sequence. BoundChemical uses class BoundUnit to represent molecules individually. It uses BoundUnitFilter to organize bound units according to outside criteria needed for reactions (classify according to binding sites, motifs read, etc.). It also uses Switches on specific switch sites that are sequence dependent (this will be explained in detail later).

For example, a RNA polymerase (RNAP) bound to DNA is a BoundChemical. A typical instance would be a RNAP bound to DNA on position 1000, another could be bound to position 2000. A BoundUnitFilter can be used to sort RNAPs according to the base they are trying to load (A, C, G or T). Finally, a Switch would be used to indicate termination sites.

# 3.2.5. Chemical Sequence

# Input format

ChemicalSequence (Fig. 6) is a subclass of FreeChemical. It is defined by a sequence and the ability to bind elements. However, instances of a sequence are *not* treated individually, it is impossible to tell to which instance a given chemical bound. An

```
ChemicalSequence
commands
add (number)
remove (number)
bind_unit (first, last)
unbind_unit (first, last)
add_switch_site (position, switch_id)
watch_site (binding site)
set_appariated_sequence (chemical sequence)
start_strand (position)
extend_strand (strand_id, position)
accessors
number_sites (first, last)
number_available_sites (first, last)
partial strands ()
is_out_of_bounds (first, last)
is_switch_site (position, switch_id)
length ()
sequence ()
sequence (first, last)
relative (absolute position)
appariated_sequence ()
complementary (position)
```

Figure 6: ChemicalSequence class

object called SequenceOccupation maintains occupation levels at sites of interest. For example, suppose the sequence is an mRNA carrying a ribosome binding site for the protein DnaA. The number of available sites is obtained by removing the number of bound chemicals occupying the site from the number of instances of the mRNA currently in the cell. A ChemicalSequence can be appariated to another ChemicalSequence. A ChemicalSequence can be created from a sequence or as a product of another sequence, in which case a TransformationTable is needed to generate the product's sequence from the parent's, and a ProductTable stores the parent/product relationship.

### 3.2.6. DoubleStrand

# Input format

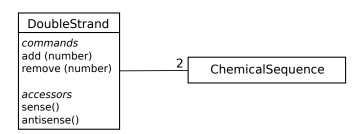


Figure 7: DoubleStrand class

DoubleStrand (Fig. 7) links two ChemicalSequence together that are biochemically linked (e.g. DNA), one sequence being complementary to the other. It enables segment extension on the appariated strand and free end binding (see interface of ChemicalSequence). A DoubleStrand is created from a sense sequence that is specified similarly to a ChemicalSequence. However, the complementary sequence is created from a TransformationTable that specifies how to transform the sense sequence into antisense sequence (e.g. for DNA,  $A \to T$ ,  $T \to A$ ,  $C \to G$ ,  $G \to C$ ).

# 3.2.7. BindingSiteFamily

# Input format

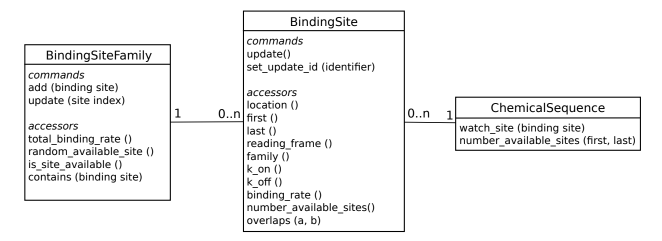


Figure 8: BindingSiteFamily class

BindingSiteFamily (Fig. 8) is a subclass of Reactant. Contrary to Chemical, it does not represent a countable pool of molecules. Each family contains a number of related instances of BindingSite (e.g. ribosome binding sites). BindingSiteFamily, BindingSite and ChemicalSequence use a notification pattern (via update methods) to dynamically maintain the number of available sites for each binding site as well as binding rates up to date. If a binding site is used to load polymerases, a reading frame should be provided to specify where a polymerase will start reading the sequence after binding.

# 3.3. Reaction hierarchy

This section gives a quick overview of the reaction hierarchy (Fig. 9). More details about how reactions are implemented can be found later.

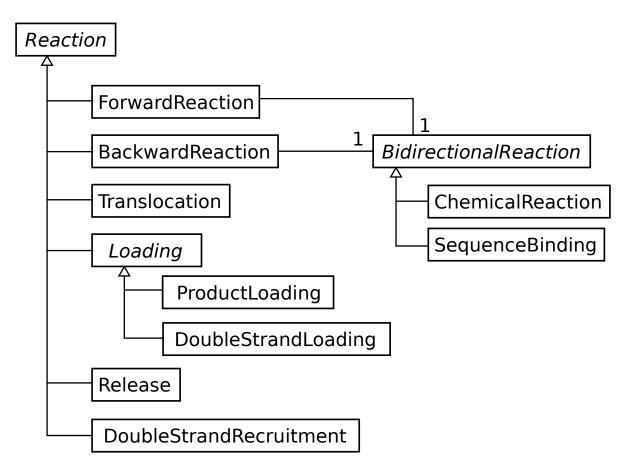


Figure 9: UML diagram of Reaction hierarchy.

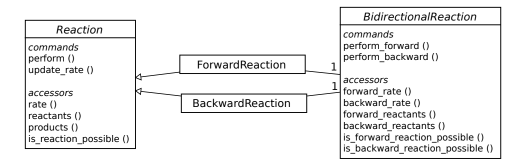


Figure 10: Reaction and BidirectionalReaction classes.

# 3.3.1. Reaction

There are two abstract classes used to define reactions: Reaction for one-way reactions and BidirectionalReaction for reversible reactions. Two adapter classes ForwardReaction and BackwardReaction split reversible reactions in two one-way reactions (Fig. 10). In the end, the solver only handles one-way reactions. A reaction can necessarily be performed, its rate updated and accessed and is composed of reactants and products.

# 3.3.2. ChemicalReaction

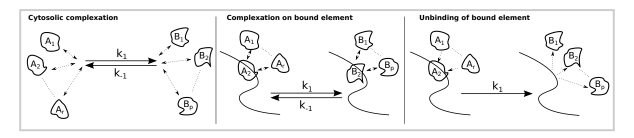


Figure 11: Schematic view of a ChemicalReaction.

# Input format

ChemicalReaction [<chemical> <stoichiometry>]^ $\{1..n\}$  rates <k\_1> <k\_-1> <

**Formula** A ChemicalReaction represents association/dissociation of an arbitrary number of elements (Fig. 11). It is defined by

$$a_1A_1 + a_2A_2 + \dots + a_rA_r \xrightarrow[k_{-1}]{k_1} b_1B_1 + \dots + b_pB_p$$

where

- $A_i$  and  $B_i$  are of type FreeChemical. They can be of type BoundChemical in two cases: (i) a reaction containing a BoundChemical on each side, (ii) an *irreversible* reaction where a *reactant* is a BoundChemical and where there are no bound product. In both cases, the associated stoichiometric coefficient must be 1.
- $a_i$  and  $b_i$  are stoichiometric coefficients.
- $k_1$  and  $k_{-1}$  are rate constants.

**Action** When the reaction is performed, the number of chemicals involved is changed according to their stoichiometric coefficient. If BoundChemical are involved on each side, the simulator will assume that the bound chemical that is consumed is replaced by the bound chemical on the other side of the equation (*i.e.* it will be bound at the location previously occupied by the precursor). If there is a BoundChemical on the reactant side of an irreversible reaction, the simulator will assume that the reaction describes the unbinding of this bound unit into the cytosol.

**Rate** The rates are given by

$$\lambda_{forward} = k_1 \prod_{i=1}^{r} [A_i]^{a_i}$$

$$\lambda_{backward} = k_{-1} \prod_{i=1}^{p} [B_i]^{b_i}$$

# 3.3.3. SequenceBinding

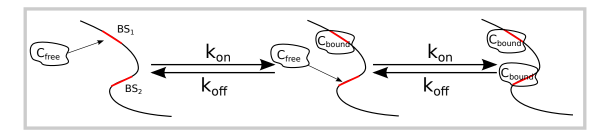


Figure 12: Schematic view of a SequenceBinding.

# Input format

SequenceBinding <chemical> <bound form> <binding site family>

**Formula** A SequenceBinding represents binding of a free element on a binding site of a sequence (Fig. 12). It is defined by

$$C_{free} + BSF \Longrightarrow C_{bound}$$

where

- $C_{free}$  is of type FreeChemical.
- $\bullet$  BSF is of type BindingSiteFamily.
- $C_{bound}$  is of type BoundChemical.

**Action** When the forward reaction is performed, a random available binding site is drawn from the binding site family (drawing is weighted by affinity). A  $C_{free}$  molecule is removed from the pool and a  $C_{bound}$  added to the ChemicalSequence bearing the binding site. When the backward reaction is performed, a random molecule of  $C_{bound}$  is removed from the pool (and from its sequence) and a  $C_{free}$  molecule is added.

**Rate** The rates are given by

$$\lambda_{forward} = \frac{[C_{free}]}{V_c} \sum_{\text{sites } s \in BSF} (k_{on})_s \times \text{Number of sites } s \text{ available}$$

$$\lambda_{backward} = \frac{1}{V_c} \sum_{\text{molecules } m \in C_{bound}} (k_{off})_{\text{site on which } m \text{ is bound}}$$

- $(k_{on})_s$  is the association constant of  $C_{free}$  with binding site s.
- $(k_{off})_s$  is the dissociation constant of  $C_{bound}$  with binding site s.
- $V_c$  is the volume of the cell.

# 3.3.4. Translocation

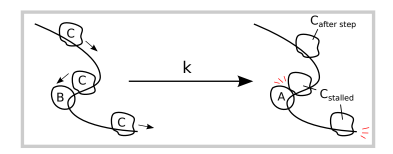


Figure 13: Schematic view of a Translocation.

# Input format

**Formula** A Translocation represents movement of a bound element along a sequence (Fig. 13). It is defined by

$$C \xrightarrow{k} C_{\text{after step}}$$

or

$$C \xrightarrow{k} C_{\text{stalled form}}$$

where

- $\bullet$  C is of type BoundChemical.
- $C_{\text{after step}}$  is of type BoundChemical.
- ullet  $C_{
  m stalled\ form}$  is of type BoundChemical.
- $\bullet$  k is a rate constant.

**Action** When the reaction is performed, a random C is chosen. Generally, it is replaced by a  $C_{\rm after\ step}$ , moved by a step of a given size along the sequence the original C is bound to. If the chemical cannot move because it reached the end of the sequence, it is replaced by  $C_{\rm stalled\ form}$ .

**Rate** The rate is given by

$$\lambda = k[C]$$

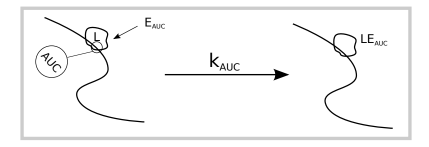


Figure 14: Schematic view of a Loading.

# **3.3.5.** Loading

# Input format

LoadingTable <name> \

[<template> <element\_to\_load> <occupied\_polymerase> <rate>,]^{1..n} ProductLoading <bound chemical> <loading table> DoubleStrandLoading <bound chemical> <loading table> <stalled form>

**Formula** A Loading typically represents loading of elements by a polymerase onto a template sequence (Fig. 14). It is defined by

$$L + E \longrightarrow LE$$

where

- L is of type BoundChemical.
- E is an element to load, of type FreeChemical. It is defined in a LoadingTable associated with the reaction.
- *LE* is the occupied form of the loader, of type BoundChemical. It is defined in a LoadingTable associated with the reaction.

**Action** Each instance of L reads a specific template. Using its LoadingTable, we know which E it tries to load, which LE is yielded if loading occurs and the loading rate associated with the template. When the reaction is performed, a random L is chosen according to loading rates. An element to load E is removed from the pool and L is replaced with LE. A ProductLoading assembles loaded elements into a product that will eventually be release in the cytosol (e.g. RNA synthesis), while DoubleStrandLoading extends segments along a DoubleStrand (e.g. DNA replication). In DoubleStrandLoading, loading may fail because the loader met a previously synthesized segment. In the latter case, it is replaced by a BoundChemical representing its stalled form.

**Rate** The rate is given by

$$\lambda = \sum_{t \in templates} k_t[L_t][E_t]$$

where

- $k_t$  is the loading rate associated with template t.
- $L_t$  corresponds to loaders L reading template t.
- $E_t$  is the chemical to load onto template t.

# 3.3.6. DoubleStrandRecruitment

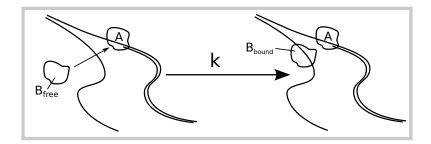


Figure 15: Schematic view of a DoubleStrandRecruitment.

# Input format

DoubleStrandRecruitment <BoundChemical> <FreeChemical> <bound form> <rate>

**Formula** A DoubleStrandRecruitment typically represents recruitment of a DNA polymerase by the replication fork on the opposite strand (Fig. 15). It is defined by

$$A + B_{free} \xrightarrow{k} A + B_{bound}$$

where

- A is of type BoundChemical, bound to a DoubleStrand.
- $B_{free}$  is of type FreeChemical.
- $B_{bound}$  is a BoundChemical representing the bound form of  $B_{free}$ .
- $\bullet$  k is a rate constant.

**Action** When the reaction is performed, a random A is chosen. If A is not bound to a DoubleStrand, the reaction is ignored. If the position opposite to a on the DoubleStrand is already occupied, the reaction is ignored. Else, a  $B_{free}$  is bound on the complementary ChemicalSequence, opposite to A as a  $B_{bound}$ .

**Rate** The rate is given by

$$\lambda = k[A][B_{free}]$$

# **3.3.7.** Release

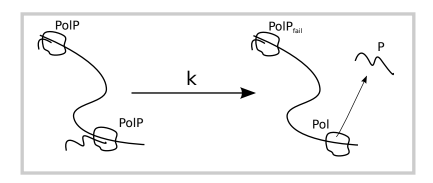


Figure 16: Schematic view of a Release.

# Input format

**Formula** A Release represents release of a product from a polymerase (Fig. 16).

$$PolP \xrightarrow{k} Pol + P$$

or

$$PolP \xrightarrow{k} PolP_{fail}$$

where

- PolP is a BoundChemical representing a polymerase-product complex.
- P is of type ChemicalSequence. It is a product that is released by PolP defined in a ProductTable associated with reaction.
- Pol is a BoundChemical representing an empty polymerase.
- $PolP_{fail}$  is a BoundChemical representing the polymerase-product complex in case release failed because P was not a valid product defined in the ProductTable associated with reaction.
- $\bullet$  k is a rate constant.

**Action** When the reaction is performed, a random PolP is chosen. A ProductTable uses its binding and current position to determine what product P it has synthesized. If P is defined in the product table, it is released in the cytosol and PolP is replaced by an empty version of the polymerase Pol. If there is no P corresponding to current PolP position, the simulator assumes that PolP has not reached its actual terminator and it is replaced by  $PolP_{fail}$  to enable other treatments (e.g. abnormal termination or continuing synthesis).

**Rate** The rate is given by

$$\lambda = k[PolP]$$

# 3.3.8. Degradation

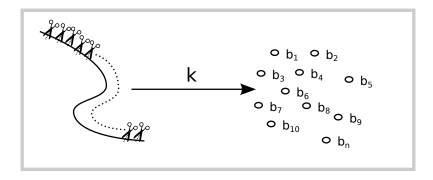


Figure 17: Schematic view of degradation reaction.

# Input format

CompositionTable <name> [<letter> [<chemical composing letter>]^{1..m}]^{1..n}
Degradation <chemical sequence> <composition table> <rate>

**Formula** A Degradation represents decomposition of a sequence into base components (Fig. 17). It is defined by

$$CS \xrightarrow{k} b_1 + b_2 + \dots + b_N$$

where

- ullet CS is of type ChemicalSequence.
- $b_i$  are of type FreeChemical. They are found in a CompositionTable specified in the reaction.
- $\bullet$  k is the degradation constant.

**Action** When the reaction is performed, a CS is removed from the pool. A CompositionTable is specified along the reaction. It allows base-by-base conversion of the sequence of CS into components yielded by degradation. The pools of base components is updated accordingly. In the simulator, a degradation reaction is effectively implemented as a ChemicalReaction.

Rate The rate is given by

$$\lambda = k[CS]$$

# 3.4. Switches

# Input format

Switch <name> <input\_bound\_chemical> <output\_bound\_chemical>
SwitchSite <chemical\_sequence> <position> <switch\_name>

Switches are intrinsically linked to BoundChemicals but apply to specific BoundUnits through SwitchSites located on ChemicalSequences. Every time an instance of input\_bound\_chemica steps on a switch site, it *immediately* becomes an output\_bound\_chemical.

For example, during transcription, an RNA polymerase (RNAP) goes through an initiation state, then loops through several elongation states (loading of a nucleotide and translocation). Once it reaches a termination site represented by a SwitchSite, the RNAP leaves its current elongation state and enters termination state. It stops performing polymerization reactions and typically releases the polymerization product and unbinds from DNA.

A Switch is not considered a reaction because there is no rate associated with it (switches are performed automatically before the solver chooses the next reaction). We dedicate a section to these elements because they play a central role in the simulator's philosophy. The user can use generic reactions that apply in general (e.g. transcription of any gene based on its sequence) and use switches every time something more specific is needed. As seen before, termination sites for transcription are expected to be Switch-Sites. Similarly, important regulation sites can be implemented using SwitchSites.

# 3.5. Solver loop

Once Reactions and Reactants are defined, they must be integrated properly. We use variants of the Gillespie algorithm to provide a framework where reactions are performed according to their current reaction rate. Roughly speaking, the main hypothesis of this framework is that reaction timings are distributed according to exponential distributions. This allows for many mathematical simplifications and harmonious integration of an arbitrary number of reactions. The central point of the algorithm is that the probability that a reaction will be the next reaction in the system is proportional to its rate (mathematically speaking, the reaction is obtained by multinomial drawing according to rates).

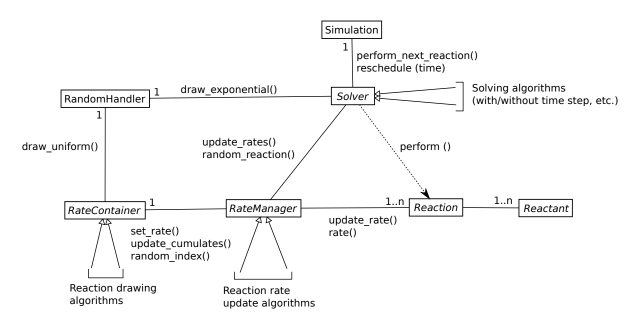


Figure 18: Solver loop. The loop is driven by the Solver class that defines how and when rates should be updated. The update task is performed by a RateManager. Once rates are known, multinomial drawing is delegated to a RateContainer. A central RandomHandler is used so that the solver only uses one seed, enabling simulation reproducibility.

The solving loop is depicted in Figure 18. The Gillespie algorithm has many variants. We decided to implement it using three *abstract* classes. By using inheritance, variants can be combined for each step of the algorithm (how to update reactions, how to select a reaction). The three central classes are:

- Solver: Children of this class decide how and when rates should be updated, e.g. update rates after every reaction, only after a given time step, etc. Note that they do not perform any of these computations, they just organize how the algorithm should work.
- RateManager: Children of this class are responsible for updating reaction rates when prompted to by a Solver class. Recomputing all rates is generally inefficient, so various implementations of this task can be used to improve the global loop speed.
- RateContainer: Childern of this class are responsible for storing reaction rates in a specific structure *adapted* to multinomial drawing. Again many implementations exist, their efficiency depends on the system that is integrated.

The implementations of these three classes will be described later in the document.

# 3.6. Events

Events enable users to change molecule numbers outside of the solver loop at specific times (Fig. 19). A Simulation instance handles both a Solver instance and an EventHandler instance. Every time an event timing is reached, the solver loop is stopped, the event(s) is (are) performed, the solver is reinitialized and the simulation resumes. Different Event implementations are offered to modify molecule numbers in a convenient way.

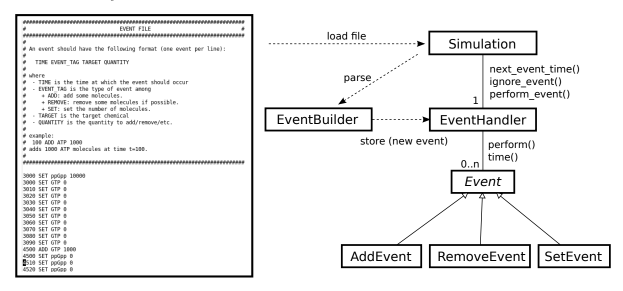


Figure 19: Events: another way to modify chemical concentrations aside from reactions, e.g. to simulate the injection of a chemical inside a cell.

# 3.7. Input/Output handling

# 3.7.1. Simulator Input

The simulator needs the following to work:

- A general input file defining simulation parameters. A sample file is provided were all options are described (e.g. length of simulation, what to output, algorithm variants). One important parameter is the location of the files the simulator should open to read reactants, reactions and events.
- An arbitrary number of files where reactants, reactions and events are declared. The simulator solves dependencies across files, it is not necessary to declare reactants in the same file as or before reactions using them.

# Caution:

- All reactants must be declared in some file with their appropriate type (e.g. FreeChemical or BoundChemical).
- Multiple declarations are forbidden, a name cannot be reused.

# 3.7.2. Simulator Output

Outputs provided by the simulator are:

- A general output file logging parametes used for simulation (input files used, random seed, algorithms used, etc.).
- A concentration file with the number of molecules over time (for the chemicals and at a time step defined in the parameter file).
- If a DoubleStrand was added in the chemicals to ouput, a replication file describing replication advancement of that DoubleStrand.

# 4. Detailed design

# 4.1. Reactants

# 4.1.1. FreeChemical

FreeChemical simply represents a pool of interchangeable molecules distributed uniformly in the cell. Computationnally, only the number of molecules in the pool is relevant.

### 4.1.2. BoundChemical

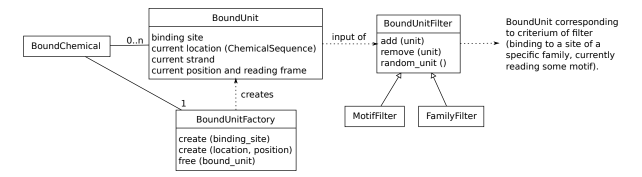


Figure 20: BoundChemical are in fact a pool of individual BoundUnit created using a BoundUnitFactory. A BoundUnit is characterized by the ChemicalSequence it bound to and its current position. Reaction then use BoundUnitFilter to sort BoundUnit according to some criterium of reference (e.g. Loading reactions sort BoundUnit according to the motif they read).

BoundChemical represents molecules of the same chemical species, but there are specifities for each unit of a BoundChemical, as all units are bound at different locations of different ChemicalSequence (Fig. 20). A BoundUnitFactory is used to recycle BoundUnits, avoiding memory reallocation throughout simulation. BoundUnitFilters are used to sort BoundUnits according to criteria useful for reactions (Fig. 20).

BoundUnits are passed from one BoundChemical species to another through reactions, their attributes are updated if needed. They are only destroyed once they are unbound from their ChemicalSequence.

# **4.1.3.** Chemical Sequence

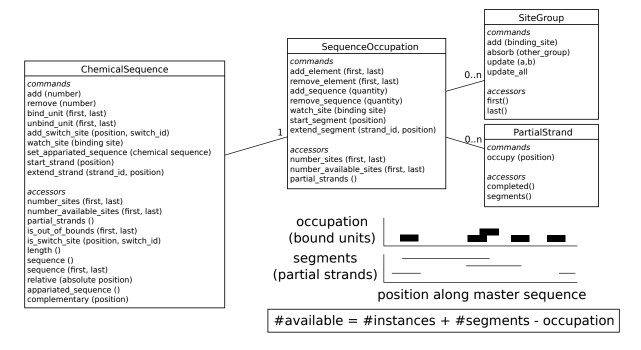


Figure 21: ChemicalSequence represents a pool of polymeres that can be elongated and on which BoundUnits bind through BindingSites. For binding to occur, availability of BindingSite is assessed using a utility class SequenceOccupation that records the number of instances of the polymer, the position of BoundUnits and elongation of PartialStrands. SiteGroup is used to notify sites of availability changes more efficiently.

ChemicalSequence handles a pool of polymers. A pool is defined by a master sequence describing what a typical polymer looks like (e.g. the sequence of DnaA protein) and the number of instances of the master sequence in the pool. For efficiency reason, we do the following assumptions.

# **Simplifying assumptions**

- No deviation from master sequence, all instances are identical.
- BoundUnits are not assigned to a specific instance of the sequence, they are positioned on the master sequence.

# Consequences

- No direct inference of collisions is possible.
- A chemical can bind on a partial strand, yet move along the whole sequence freely.
- Degradation of an instance does not cause unbinding.

**Site availability** Despite our simplifying assumptions it is still possible to provide an accurate description of site availability. Availability depends of the number of sequences, number and position of bound elements, number and position of newly polymerized sequence segments (Fig. 21).

# 4.1.4. DoubleStrand

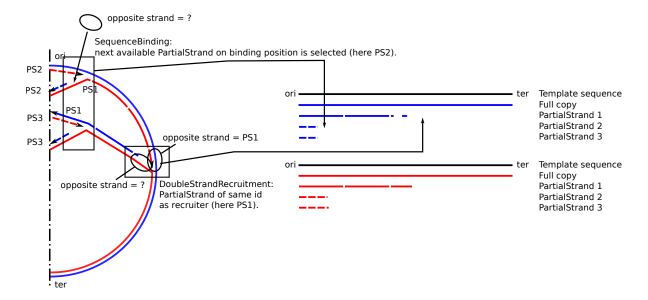


Figure 22: Strands of a DoubleStrand are identified according to creation order. Every time a new segment is polymerized, it is necessary to determine which PartialStrand is elongated. If a polymerase has been recruited on the complementary strand by DoubleStrandRecruitment, it is automatically assigned the same partial strand as the recruiter.

**Strand identification** Because DoubleStrand typically represents DNA, we expect that the DoubleStrand will contain a lot of PartialStrands. For replication, it is important to know exactly which strand are opposite to one another for DoubleStrandRecruitment to work properly. We use strand identification as shown in Figure 22.

# 4.1.5. BindingSiteFamily

The task of a BindingSiteFamily is to regroup all the binding sites that can participate in a same SequenceBinding reaction. To simplify the reaction, it stores the subrate associated with each binding site. In order to update the rate properly when availability of sites changes, an *observer pattern* is used (Fig. 23).

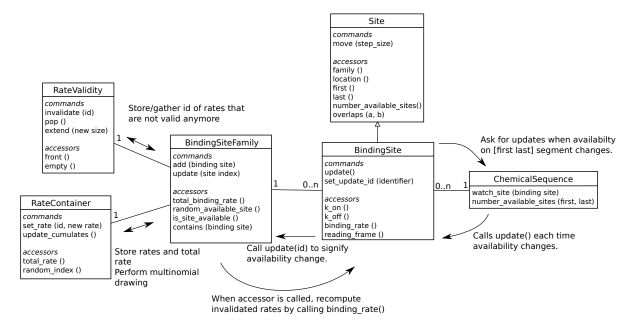


Figure 23: Schematical view of the Observer pattern used to keep availability of binding sites up to date for SequenceBinding reactions.

Every BindingSite is viewed as an observer by the ChemicalSequence it belongs to. Every time a change occurs on the site, the BindingSite is notified. The latter binding site notifies its BindingSiteFamily using a specific identifier, letting the family know which binding rate is out of date. This information is stored in a RateValidity class. It is only when it is really needed (i.e. when a SequenceBinding wants to access total rate or a random site) that rates are recomputed. This avoids useless computations e.g. in the case of a translocation, where a bound unit is first unbound from its ChemicalSequence then rebound. If the bound unit does not move away from the site, two updates will be sent, but the rate will only be recomputed once at the end.

# 4.2. Reactions

# 4.2.1. ChemicalReaction

Nothing particular.

# **4.2.2.** SequenceBinding

**Binding** Because of the way BindingSiteFamily is implemented, the reaction can easily and efficiently access the binding rate at all times, no matter what reactions have occured previously and how site availability changed in the meantime.

**Unbinding** SequenceBinding uses a FamilyFilter (see detailed description of BoundChemical) to filter out all BoundUnits that are bound to a binding site of the BindingSiteFamily associated with the reaction. BoundUnits that have bound to sites of a different family or that have moved away from the binding site through Translocation are *not* candidates for unbiding.

### 4.2.3. Translocation

**Collisions** For now, Translocation ignores collisions, making its implementation straightforward.

**Stalled form** Translocation enters stalled form if a BoundUnit reached the end of a sequence.

# **4.2.4.** Loading

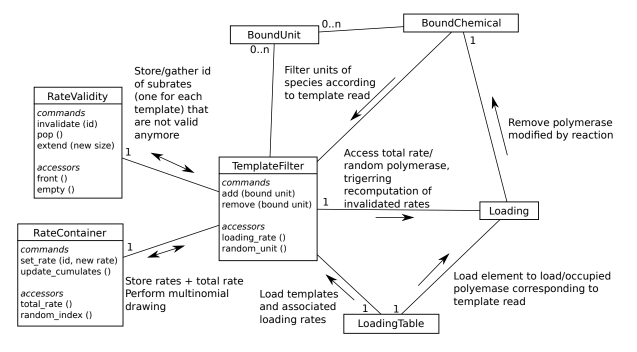


Figure 24: Schematical view of the pattern used to keep subrates associated with each template up to date in a Loading reaction.

Handling each polymerase individually The main challenge with Loading is to maintain the subrates associated with each motif up to date. It needs to maintain a list of all BoundUnits reading a specifing motif. To this end it uses a TemplateFilter (see detailed implementation of BoundChemical). Every time a BoundUnit becomes of the type of the BoundChemical associated with the reaction, the filter looks what motif defined in the LoadingTable it is currently reading. If the motif could not be found, an UNKNOWN TEMPLATE error message is displayed, the BoundUnit is not recorded in the filter and will not participate in the Loading reaction. The implementation is very similar to that used for BindingSiteFamily (Fig. 24).

ProductLoading vs DoubleStrandLoading There difference between the two processes is rather small. We just added a failure condition in the case of DoubleStrandLoading for convenience. Depending on what reactions are used to synthesize a DoubleStrand it might be possible that a polymerase arrives upon a position that has already been synthesized. In this case, the DoubleStrandLoading fails and the polymerase is replaced by the polymerase in its stalled form.

### **4.2.5.** Release

Fail polymerase (unknown product) When a release is triggered, a BoundUnit from the BoundChemical associated with the Release reaction is randomly chosen. Because the BoundUnit knows its current position and its binding site, it will assume that product it has synthesized starts the reading frame of the binding site and ends at the position directly preceding its current reading frame (we assume that the polymerase translocates onto a terminating sequence which does not contribute to product synthesis). If the product is found in the ProductTable, everything works normally.

If the product is not found, we display a Unknown Product error message but keep the simulation alive. The fail polymerase in the reaction enables the user to define a rescue pathway. If the release competes with some other reaction for the original polymerase, the fail polymerase can be the original polymerase itself. If products overlap and the polymerase was stalled due to a termination site of another product, fail polymerase can be a polymerase in a sythesizing step (e.g. ProductLoading) so synthesis will resume until the next termination site is reached.

# 4.3. Solver loop

Here we describe the implementations provided for each step of the algorithm. Most of the details are explained in a side document (Dinh et al., 2016). We only give a quick overview here.

### 4.3.1. RateContainer classes

We start with the lowest level classes, which perform one of the central tasks of the Gillespie algorithm: drawing a reaction from reaction rates. For efficiency reasons, we

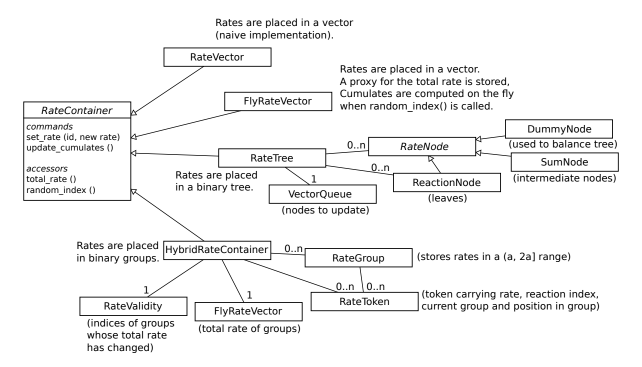


Figure 25: Implementations provided to store rates and perform a multinomial drawing. Implicitly, all theses classes use RandomHandler to perform their random drawings.

proposes several implementation of the algorithm (Fig. 25). Comparison and description of theses classes are given in Dinh et al. (2016).

Note that multinomial drawing occurs within the solver loop, but also within some reactions such as Loading or SequenceBinding, so these classes are used quite extensively throughout the simulation.

# **4.3.2.** RateManager classes

The second layer of the solver loop ensures that the rates are updated when needed to. Two implementations are proposed for this task (Fig. 26). The NaiveRateManager updates every rate. While it is inefficient, it can be used as a reference to test other managers. The DependencyRateManager uses an observer pattern to update only reactions for which a reactant concentration has changed (see Dinh et al. (2016) for further details).

# 4.3.3. Solver classes

For the moment, only one solver class is fully available to the user, NaiveSolver, which implements the exact Gillespie algorithm. Another variant called ManualDispatchSolver is implemented, were the user can assign a time step to each reaction at which its rate will be updated (Fig. 27). However, when the rate of a reaction is a constant, there

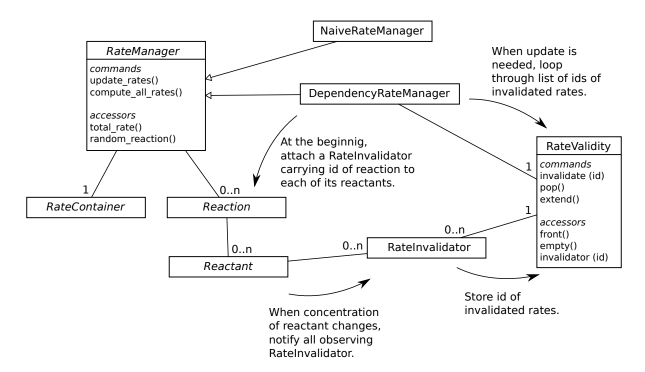


Figure 26: Implementations provided to update reaction rates. Note that the drawing part of the algorithm is always delegated to a RateContainer. DependencyRateManager uses an Observer pattern to monitor which rates have changed.

is a risk that its reactants will run out and the reaction will be impossible to realize or reactant number will become negative. In the simulator, the latter case is forbidden, so ManualDispatchSolver ignores reactions impossible to perform due to reactant inavailability.

# 4.4. Input/Output handling

# 4.4.1. Parsing system

The parsing system used by the simulator is pretty simple (Fig. 28). A SimulationParams class is used to store simulation parameters (which will be used to create and drive the Solver), CellState stores reactions to integrate and EventHandler stores events. At the moment, an *ad hoc* input format is used.

# 4.4.2. Builder and interpreter

The parsing system is designed to cut each line into words, then the words are interpreted by Builders that try to create instances of each of the class of the simulator. The Parser loops through the Builders until an instance was successfully created. Line format is checked token-wise by an Interpreter (Fig. 29). If no Builder is able to match the line, a FormatException is raised. If some dependency could not be solved,

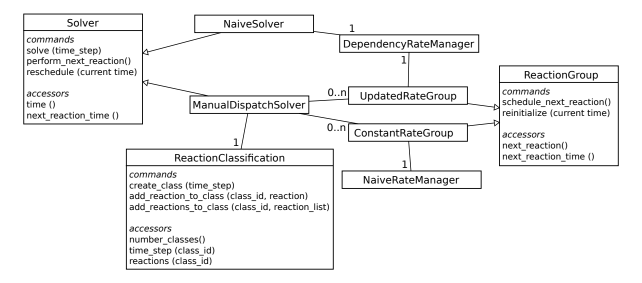


Figure 27: Two implementations of the Solver class organizing rate updating. NaiveSolver forces recomputation of rates at each time step. ManualDispatchSolver puts reactions into groups: reactions in UpdatedRateGroup are updated after every reaction while those in ConstantRateGroup only at user-defined steps defined in ReactionClassification. Note that all Solver classes use at leaste a variant of RateManager at some point to delegate storing and updating of rates.

a DependencyException is raised. In the latter case, the Parser will postpone the line until dependency can be successfully solved.

# 4.4.3. Output

Two classes are used to produce output. ChemicalLogger logs chemical numbers through time. DoubleStrandLogger logs partial strands of a DoubleStrand.

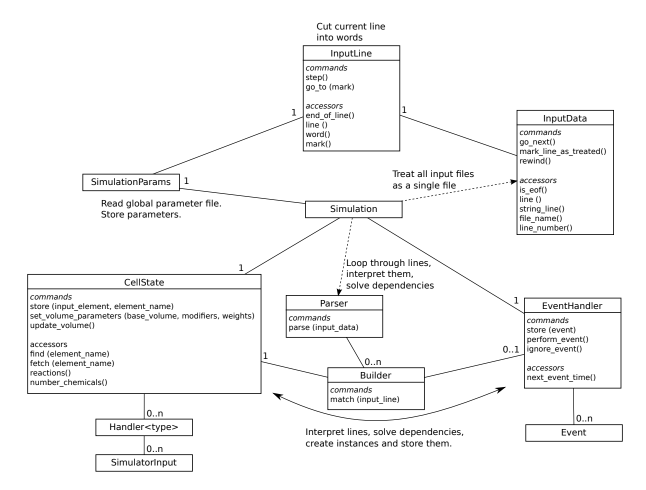
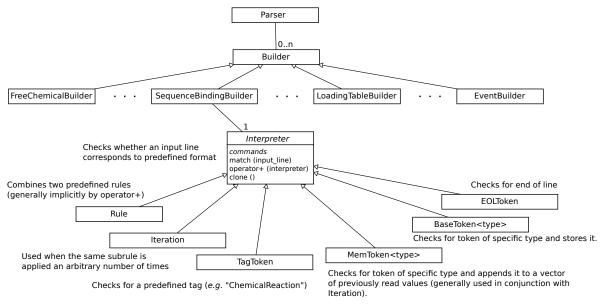


Figure 28: Architecture of the parsing system. SimulationParams reads the global parameter file and stores parameter values. InputData is used to provide a simplified interface to the input files and mark treated lines. A Parser and Builders are used to make sense of individual lines. Everything that is created is stored in EventHandler (for events) and CellState (for the rest).



example:

TagToken("SequenceBinding") + BaseToken<string> (unit\_to\_bind) + BaseToken<string> (bound\_unit) + BaseToken<string> (binding\_site\_family)

Figure 29: Interpreter system used. For each class of the simulator, a Builder is responsible for interpreting current line, solving dependencies, checking validity of parameters. If format is invalid or dependencies could not be resolved, exceptions are raised to warn the user.

# A. Utility classes

# A.1. Exceptions

Base class provided by C++ standard library.

std::exception

DependencyException

ParserException

Exception used to indicate that something went wrong while interpreting. Parent class is generally used when a wrong parameter value was provided.

Exception used to indicate that a dependency could not be solved in a reaction or an event.

FormatException

Exception used to indicate that the format is invalid.

Figure 30: Exceptions used in the simulator.

Programming by contract (see Section B) covers most internal errors that might happen. Exceptions are only used when user input is treated. They are used to signify inconstencies in input files during the parsing step (Fig. 30).

# A.2. Random handler

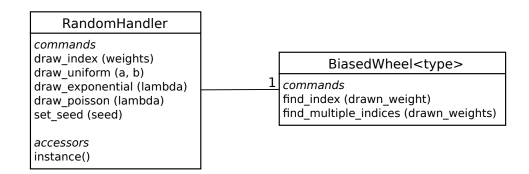


Figure 31: RandomHandler and BiasedWheel used for multinomial drawing. RandomHandler uses the Singleton pattern, meaning exactly one instance of the class is created and used throughout the simulator. It has to be accessed using RandomHandler::instance().

A unique RandomHandler is used throughout the simulation to control the random seed by using a Singleton pattern (Fig. 31).

# A.3. Factories

Factories are used to remember user options and handle memory more efficiently (Fig. 32).

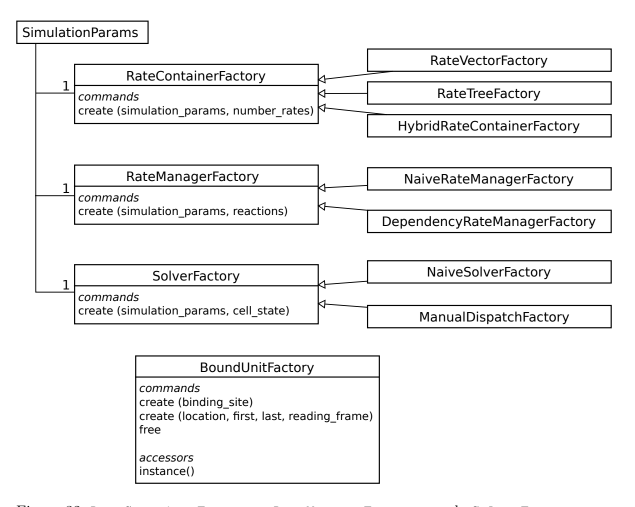


Figure 32: RateContainerFactory, RateManagerFactory and SolverFactory are used by SimulationParams to record what RateContainer, RateManager and Solver the user wishes to use (Abstract Factory Pattern). BoundUnitFactory is used to control memory usage by BoundUnits.It enables recycling of bound units, avoiding memory reallocation. It uses the Singleton pattern to make sure all instances of BoundUnits are stored at the same place.

# A.4. Vector-based containers

The simulator uses two non-standard containers, VectorList and VectorQueue. A typical example is a BoundChemical storing all its BoundUnits. There is typically a high turnover of bound units and units reacting are drawn uniformly within the list of bound units. Naively, we would use a list to perform such a task, but there are huge performance issues. Using standard C++ std::list would imply a lot of memory reallocation each time a BoundUnit is added/removed (because a node of the list is created/deleted). What is more, if, by random drawing, we decide that it is the 10th unit that is going to react, we need to loop through 10 elements before accessing the correct element.

Using a std::list has a huge impact on performance. Therefore, in BoundChemical (and a lot of other places in the program), we replace std::list by a VectorList, where elements are placed in a std::vector. There is one trick to use: every time an element is removed, it is replaced by the last element in the vector, so that elements remain contiguous in memory. This means that order of elements is lost, but in the example above and every time we VectorList, order is not important. The size of std::vector is automatically adjusted by C++. Most of the time it will be larger than the number of elements it contains, but we prefer using a little more memory than contantly reallocating nodes. What is more, accessing the nth element is instantaneous.

The same general idea applies for VectorQueue except we need to know in advance how large the queue will be. It is used to update nodes in RateTree because we know how many nodes there are in the tree and they are updated at most once.

# A.5. Handler classes for memory handling

C++ has no garbage collector and this program was designed according to old standards (that is without smart pointers). Therefore, we need to be extremely careful to delete elements at the right time. This is achieved by using as few storage places as possible. As described earlier, CellState is used to store all reactions and reactants read from files. We designed a Handler class that effectively stores objects to their definitive location and distributes references or pointers to this constant location. Similarly EventHandler and BoundUnitFactory are used to store Events and BoundUnits which are the other dynamical elements stored in the program. At the end of the simulation, all these handlers have to be carefully deleted. Observer patterns are particularly dangerous, as we must be sure that at destruction, there is no message sent to a non-existent observer. Every time an observer pattern is used, we made sure that observers are correctly unsuscribed when they are destructed or the object they observe is destructed.

# **B.** Tests

# **B.1.** Testing philosophy

We use three layers of tests: programming by contract, unit tests, integration tests (Tab. 1). They are integrated in an automated framework to detect bugs rapidly and at the lowest possible level.

**Programming by contract** These tests typically apply to attributes of classes and arguments of methods. They are usually divided into three subcategories: *preconditions*, *postconditions* and *invariants*. They check whether the class interacts correctly with the outside world, generally other classes. We left invariants out because they are hard to check in a language that does not support them natively. In the simulator, we defined two macros REQUIRE and ENSURE to test pre- and postconditions. Each time a precondition or a postcondition is broken, the program is interrupted and the condition that was

Test type	Preconditions Postconditions Invariants	Unit Tests	Integration Tests
Test level	Implementation details	Class interface	Systemic
Time per test	a few instructions (ns)	ms to a few seconds	seconds to several minutes
Use frequency	Permanent	Very frequent	Less frequent

Table 1: Comparisons of tests used to develop the simulator

broken is displayed (using assert()). Preconditions and postconditions are extremely useful for detecting simple typing mistakes, numerical issues and so on (Fig. 33).

```
int RateTree::find (double value) const
{
    /** @pre value must be smaller than total tree rate. */
    REQUIRE (value <= total_rate());
    /** @pre value must be strictly positive. */
    REQUIRE (value > 0);
    int index = _root->find (value);
    // rarely, the algorithm will fail because of rounding problems and
    // return a leaf with zero rate. We just take the next nonzero rate.
    while (_leaves [index]->rate() == 0)
        { ++index; if (index == _leaves.size()) { index = 0; } }
    /** @post Rate of returned leaf must be strictly positive. */
    ENSURE (_leaves [index]->rate() > 0);
    return index;
}
```

Figure 33: Example of the use of preconditions with REQUIRE and postconditions with ENSURE. Here the postcondition helped discover a rare bug due to rounding problems that is now adressed in the code.

**Unit tests** Unit tests test the behavior at the class level. We used some simple guidelines to try and write useful and maintainable tests. We only wrote tests for the most important classes of the simulator.

**Integration tests** This is the last layer of test. The whole simulator or large pieces of it are used. The idea is to test more systemic behaviors, in which the interaction of classes is crucial. Typical examples are:

- If we provide known DNA and define RNAs and proteins correctly according to simulator input, the sequence of proteins as processed by the simulator should match known proteins.
- If we provide DNA, define RNAs, provide a transcription pathway but activate only one promoter, only the RNA associated to that promoter should be transcribed.

# B.2. Organizing and running tests

Tests are associated to the source code of the simulator in order to be run as frequently and as simply as possible. Tests are driven by Unix Autotools and use the BOOST Test Framework. Preconditions and postconditions are written inside the code, unit tests are regrouped in a tests/unit\_tests directory and integration tests are stored in tests/integration.

Options of ./configure are used to activate every layer of test individually. By default, all tests are turned off. Several layers can be activated simultaneously.

- --enable-pre-check enables preconditions.
- --enable-post-check enables postconditions.
- --enable-unit-tests enables unit tests and the possibility to create mock objects.
- --enable-integration-tests enables integration tests and the possibility to create mock objects.

Code needs to be recompiled after it has been configured.

- Preconditions and postconditions are automatically checked every time the program is run (for unit tests, integration tests or any kind of other run). Remember to turn them off for real simulations as they are very time consuming.
- Unit tests and/or integration tests are run by running make check. BOOST automatically generates useful and human readable messages about tests that failed or clearly indicates that all tests have passed.

# C. Formats and Conventions

# C.1. Input format description

- A plain word indicates a tag, that needs to be written.
- <...> indicates a variable that has to be completed with an existent element of the specified type.
- [...] indicates an optional part.

- [...]^{0..n} indicates an optional part that can be repeated an arbitrary number of times.
- [...]^{1..n} indicates an part that can be repeated an arbitrary number of times, at least once.
- [...,]^{0/1..n} indicates a part that can be repeated an arbitrary number of times, each repetition being separated by a , (but there is actually no , after the last repetition).

# C.2. UML

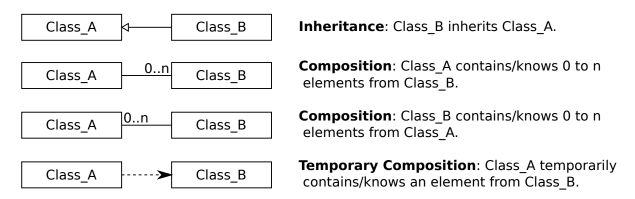


Figure 34: UML format used.

# References

Marc Dinh, Stephan Fischer, and Anne Goelzer. CATI MIAGO: comparing algorithms for gillespie based simulations. 2016.