Abstract—The importance of stochasticity within biological systems has been shown repeatedly during the last years and has raised the need for efficient stochastic tools. We present SABRE, a tool for stochastic analysis of biochemical reaction networks. SABRE implements fast adaptive uniformization (FAU), a direct numerical approximation algorithm for computing transient solutions of biochemical reaction networks. Biochemical reactions networks represent biological systems studied at a molecular level and these reactions can be modeled as transitions of a Markov chain. SABRE accepts as input the formalism of guarded commands, which it interprets either as continuous-time or as discrete-time Markov chains. Besides operating in a stochastic mode, SABRE may also perform a deterministic analysis by directly computing a mean-field approximation of the system under study. We illustrate the different functionalities of SABRE by means of biological case studies.

I. INTRODUCTION

Markov chains are an omnipresent modeling approach in the applied sciences. Often, they describe population processes, that is, they operate on a multidimensional discrete state space, where each dimension of a state represents the number of individuals of a certain type. Depending on the application area, “individuals” may be customers in a queuing network, molecules in a chemically reacting volume, servers in a computer network, actual individuals in a population, etc.

Here, we are particularly interested in dynamical models of biochemical reaction networks, such as signaling pathways, gene expression networks, and metabolic networks. Biochemical reaction networks operate on an abstraction level where a state of the system is given by an n-dimensional vector of chemical populations, that is, the system involves n different types of molecules and the i-th element of the state vector represents the number of molecules of type i. Molecules collide randomly and may undergo chemical reactions, which change the state of the system. Classical modeling approaches in biochemistry are based on a system of ordinary differential equations that assume a continuous deterministic change of chemical concentrations. During the last decade, stochastic analysis of biochemical reaction networks has seen growing interest because it captures molecular noise [13], which arises from the randomness of the discrete events in the cell. Molecular noise is of interest because it significantly influences fundamental biological processes such as gene expression [12, 35], decisions of the cell fate [2, 29], and circadian oscillations [3, 16].

Within the setting of stochastic analysis, biochemical reaction networks are modeled as discrete-state continuous-time Markov processes (CTMC) as suggested by Gillespie within the theory of stochastic chemical kinetics [14]. The evolution of a CTMC is given by a system of linear ordinary differential equations, known as the chemical master equation (CME). A single equation in the CME describes the time-derivative of the probability of a certain state at all times t ≥ 0. Thus, the solution of the CME is the probability distribution over all states of the CTMC at a particular time t, that is, the transient state probabilities at time t. The solution of the CME is then used to derive measures of interest such as the distribution of switching delays [30], the distribution of the time of DNA replication initiation at different origins [31], or the distribution of gene expression products [37]. Moreover, many parameter estimation methods require the computation of the posterior distribution because means and variances do not provide enough information to calibrate parameters [19].

Statistical estimation procedures such as Monte Carlo simulation are widely used to estimate the probability distribution of the underlying Markov process, because for realistic systems the size of the CME is very large or even infinite, and numerical methods become infeasible. Several tools for Monte Carlo simulation have been developed [18, 25, 32]. Recent work, however, indicates that numerical approximation methods for the CME can be used to compute the transient state probabilities more accurately and, depending on the measures of interest, with shorter running times [9]. Especially if the probabilities of interest are small, numerical approximations turn out to be superior to Monte Carlo simulation, because the later requires a large number of simulation runs in order to bound the statistical error appropriately. For estimating event probabilities, a higher precision level is necessary than for estimating cumulative measures such as expectations, and simulation based methods have a slow convergence because doubling the precision requires four times more simulation runs to be performed.

In the case of discrete-time Markov chains (DTMCs), the transient stochastic analysis gives the probability distribution over all states of the DTMC after k steps. For population
models, a step is interpreted as a triggered transition. The transient solution for DTMCs is the result of $k$ matrix-vector products, and can be used for the solution of CTMCs, as shown later in Section III-A.

Numerical analysis tools for discrete-state Markov processes such as PRISM [28], INFAMY [17], ETMCC [22], MRMC [26], APNToolbox [5], SHARPE [23], SPNP [24], or Möbius [8] have been introduced (see Section VII). However, except for INFAMY, these tools do not accept models with possibly infinite state space. It is important to note that many population models have an infinite state space, that is, the number of reachable states is infinite. Even when in the real system the number of molecules, or more generally, individuals is finite, no a priori bound is known, and models do not include any constraints on the number of molecules, for example in production rules such as $\emptyset \rightarrow A$. Another issue is that existing tools usually implement algorithms that are not optimized specifically for population models, and do not scale well on such models.

SABRE is a tool for the transient analysis of Markov population models. In other words, SABRE analysis discrete-time, or continuous-time Markov processes that have a structured discrete state space and state-depended rate functions. In Section II we give more details on the space structure and the state dependency of rate functions that are present in Markov processes that represent population models.

SABRE offers both stochastic and deterministic analysis of population models. For stochastic analysis, SABRE implements three algorithms: standard uniformization, fast adaptive uniformization and Runge-Kutta fourth order method. The different configurations in which SABRE may operate are depicted in Figure 2. The focus of the tool is on the fast adaptive uniformization method, while the remaining methods are given for completeness and comparison.

Fast adaptive uniformization is a variant of the uniformization method [34, 36] which is, an efficient method to compute probability distributions if the number of states of the Markov process is manageable. However, the size of a Markov process that represents a biochemical reaction network is usually far beyond what is feasible.

Fast adaptive uniformization [10] improves the original uniformization method at the cost of a small approximation error. The main ideas for this improvement are the on-the-fly construction of the state space and the restriction imposed on the state space to contain only states with significant probabilities, e.g. states that have a probability larger than $10^{-15}$. Even though fast adaptive uniformization can treat larger models than the previous uniformization methods could, as expected, models with remarkably high expected populations remain unsolvable and should be studied using deterministic analysis of simulation tools. A second down side of fast adaptive uniformization is that, due to the approximation error, it can overlook rare events of the model, e.g. events that occur with a very small probability.

SABRE is available on-line at http://mtc.epfl.ch/~mateescu/sabre First, the user gives an input model (either in SBML format or in guarded commands format) and a time horizon and then the transient analysis of the system starts (see Figure 1). More details on the usage of the tool are given in Section IV.

II. Guarded Commands

Guarded-command models (GCM) is the input formalism of SABRE. GCMs are a textual description of processes and are given in the style of Dijkstra’s guarded-command language [11]. Their syntax has subsequently been used by languages such as Reactive Modules [1] and by the language for specifying PRISM models [33]. The basic unit within GCMs is a transition class, which is expressed as a guarded command that operates on the state variables of the system. A transition class encodes for a possibly infinite number of state transitions. Within population models, the state variables of the system are non-negative integers representing numbers of molecules for each species. A guarded command takes the form

$$\text{guard} \models \text{rate} \rightarrow \text{update}$$

where the guard is a Boolean predicate over the variables that determines in which states the corresponding transitions are enabled. The update is a rule that describes the change of the system variables if the transition is performed. Syntactically, update is a list of statements, each assigning to a variable an expression over variables. Assume that $x$ is a variable. If, for instance, the update rule is that $x$ is incremented by 1, we write $x := x + 1$. We assume that variables that are not listed in the update rule do not change if the transition is taken. Each guarded command also assigns a rate to the corresponding transitions, which is a function on the state variables. Within SABRE, rate is given in infix notation. In the case of population models, the update function is incrementing or decrementing each variable by a constant integer.

For a population model with $m$ reactions, the GCM description is a set of $m$ guarded commands, which we index as $\text{guard}_j \models \text{rate}_j \rightarrow \text{update}_j$, where each of the commands $j$, with $1 \leq j \leq m$, describe the $j$-th reaction of the model.

GCMs are used to express both CTMCs and DTMCs. The difference between the two interpretations comes from the semantic given to the rate function of each command. In the case of CTMCs, for a given reaction $j$, the rate function $\text{rate}_j$ assigns to each state $s$, a positive real value that represents the rate of the outgoing transition $j$.

In the case of DTMCs, the rate function $\text{rate}_j$ assigns to each state $s$, a positive real value that represents the transition probability from state $s$ to its successor on reaction $j$. The functions $\text{rate}_j$ must define probability distribution over the direct successor state, that is, for each state $s$ we impose that $\Sigma_j \text{rate}_j(s) = 1$. If the input is not given in this manner, SABRE will automatically normalize the rate functions such that the probability distribution condition to be fulfilled. Note that this is equivalent to interpreting the input as a CTMC and than considering its embedded DTMC.
GCMs are used to model systems that exhibit a finite number of transition types, but possibly an unbounded number of states. For example, in a computer network, the number of type of events is finite (send message, receive message, add node, etc.) but the number of states is countably infinite, because it depends on the number of nodes in the network and on the number of requests each of them has. The same holds for biochemical reaction networks, each reaction type generates a transition class, but the number of states is countably infinite, as we do not have any a-priori bound on the variables of the system, due to productions rules of the type \( \emptyset \to A \). We therefore conclude that GCMs are a natural formalism for describing population models.

Example 2.1: The bistable toggle switch is a prototype of a genetic switch with two competing repressor proteins and four reactions. We call the species \( A \) and \( B \) and we let \( x = (x_A, x_B) \in \mathbb{N}_0^2 \) be a vector describing a state of the system. The reactions are given in Table I.

| Reaction | Guarded command | Transition function |
|----------|-----------------|---------------------|
| \( \emptyset \to A \) | true | \( \gamma_1 \cdot (c_1 + x_A^2) \rightarrow x_A := x_A + 1 \) |
| \( A \to \emptyset \) | \( A > 0 \) | \( \gamma_2 \cdot x_1 \rightarrow x_A := x_A - 1 \) |
| \( \emptyset \to B \) | true | \( \gamma_3 \cdot (c_1 + x_A^2) \rightarrow x_B := x_B + 1 \) |
| \( B \to \emptyset \) | \( B > 0 \) | \( \gamma_4 \cdot x_2 \rightarrow x_B := x_B - 1 \) |

III. STOCHASTIC AND DETERMINISTIC ANALYSIS

SABRE performs a transient analysis of the input system, that is, SABRE computes the state of the system at time \( t \) given the state of the system at time 0. SABRE may execute either a stochastic analysis or a deterministic analysis of the input system; and in the first case the state of the system at time \( t \) is actually given as a probability distribution over the discrete states of the system. The second type of analysis—the deterministic analysis—is done over a continuous state space, and its result is a single state of this continuous space. The result of the deterministic analysis, also known as mean field analysis, is an approximation of the expectation of the stochastic analysis. Each of the two analysis (stochastic and deterministic) may be applied on each of the two semantics (CTMC and DTMC), and we will now give short interpretations for the results of the four possible combinations.

A. Stochastic Analysis

CTMC semantics: We note that the behavior of the CTMC is described as a differential equation (known in biochemistry as the chemical master equation) and that \( p(t) \) is the solution of that differential equation at time \( t \). The transient stochastic analysis at time \( t \), given the initial state \( y_0 \) with probability 1, computes the solution of the chemical master equation at time \( t \). Within SABRE, the solution \( p(t) \) may be computed either by uniformization or by Runge-Kutta explicit fourth order method.

We focus on two uniformization methods for CTMCs, standard uniformization and its generalization called adaptive uniformization. Standard uniformization splits the given CTMC into a discrete-time Markov chain (DTMC) and a Poisson process, whereas adaptive uniformization splits the CTMC into a DTMC and a birth process. SABRE implements the optimized algorithm called fast adaptive uniformization that has previously been proposed. One main strength of this algorithm is that it closely tracks the set of significant states of the state space, where by a significant state we mean a state with significant probability. Secondly, another strength of the fast adaptive uniformization lies in the on-the-fly construction of a non-explicit matrix used in the computation of the solution of the DTMC (remember that uniformization splits the given CTMC into a DTMC and a birth process).

DTMC semantics: DTMC semantics are to be used when the number of triggered transitions, rather than the elapsed time, is of interest. Such situations may arise, for example, in population genetics models or as a part of the uniformization method. A transient analysis of a DTMC consists in a series of matrix-vector products:

\[
p(k + 1) = p(k) \cdot P,
\]

with \( P \) being the probability transition matrix of the DTMC and \( p(k) \) being a row vector representing the probability distribution after \( k \) steps. In our algorithm, the main phase of the DTMC transient analysis is called the propagation phase. The propagation phase completes the equivalent of a matrix-vector product by moving probability mass from on state \( x \) to all direct successors of \( x \) (including \( x \) itself if any self-loops are present). SABRE approximates the probability distribution over the states of the system after \( k \) reactions have happened, given the state \( y_0 \) of the system before any reaction happens. Formally, SABRE approximates the vector \( p(k) = \delta_{y_0} \cdot P^k \), where \( \delta_{y_0} \) is a dirac probability distribution in point \( y_0 \).

B. Deterministic Analysis

CTMC semantics: We can give an approximate solution of the mean field of the CTMC by using the fourth order Runge-Kutta method to solve a set of ordinary differential equations simpler than the CME. This set of equations are known as the reaction rate equations and express the change in the expectation of each variable over time. In the thermodynamic limit (that is, the number of molecules and the volume of the system approach infinity) the Markov model and the macroscopic ODE description are equal. Therefore, for large populations, the deterministic analysis can be used to approximate the mean field of the CTMC.

DTMC semantics: As in the case of CTMCs, for computing the first moment of the transient solution of a DTMC, we can directly solve a simpler set of equations that are written directly over variables that represent the expectancies of the stochastic solution of the input DTMC model.

The expected number of molecules changes deterministically over discrete time, as described by the following equation:

\[
x(k + 1) = x(k) \cdot A,
\]
where \( A \) is a probability matrix, and each of its entries \( a_{i,j} \) give the probability for a species \( i \) to modify into a species \( j \). Such analysis are useful for discrete-time models as those used to validate communication protocols.

IV. TOOL INTERFACE

From the tool’s interface, we have several ways of selecting a model for analysis. One can load an existing model, upload an SBML file or introduce a GCM text description of the system to analyze. SBML is a standardized format for representing models of biological processes, such as metabolism or cell signaling and is the input to SABRE’s core program. GCMs that have update functions with constant increment (or decrement) have a straight forward translation to SBML.

Example 4.1: We continue the toggle switch example with its SBML description. For brevity, we only give one reaction of the model. We observe that the rate function is not restricted to a particular template and is written following the mathML standard.

```xml
<sbml ...>
  <model>
    <listOfSpecies>
      <species id="A" initialAmount="133"/>
      <species id="B" initialAmount="133"/>
    </listOfSpecies>
    <listOfReactions>
      <reaction id="R1">
        <listOfProducts>
          <speciesReference species="A"/>
        </listOfProducts>
        <listOfModifiers>
          <speciesReference species="B"/>
        </listOfModifiers>
        <kineticLaw>
          <math ...
            <apply> <divide/>
              <ci> c1 </ci>
              <apply> <plus/>
                <ci> c2 </ci>
                <apply> <times/>
                  <ci> B </ci>
                  <ci> B </ci>
                </apply>
              </apply>
            </apply>
          <listOfParameters>
            <parameter id="c1" value="3000"/>
            <parameter id="c2" value="11000"/>
          </listOfParameters>
        </kineticLaw>
      </reaction>
    </listOfReactions>
  </model>
</sbml>
```

Once the model is chosen, we choose a configuration of the analysis by choosing the semantics, the mode and, if needed, the type of stochastic solution. Finally, we choose a time horizon, or the number of steps for which we want the system to run. We also give as an input a dump time \( t_d \), which corresponds to the intermediate results, that is, the system will compute the distributions for \( t_d, 2 \cdot t_d, \cdots \cdot t \). The program computes the intermediates and the final results which are then dynamically plotted for each species, as the computation runs (see Figure [1]). If the uniformization method is selected, the user also needs to provide an estimate of the maximal exit rate over all reachable states. If the estimate is too small, the computation needs to be restarted, and if the estimate is too large, the computation is likely to take longer. It is standard uniformization which is especially touched by choosing a too large upper bound on the maximal exit rate. Estimating this upper bound by heuristics such as those used for the sliding window algorithm[21] is an ongoing work.

V. SOFTWARE ARCHITECTURE

SABRE is available on line, assuring this was a fast and portable release of our implementation. The core of our tool is implemented in C++, while the website that hosts it is implemented using PHP and Javascript. The user provides the desired input through the web interface, than a query is generated to the 3GHz Linux machine on which SABRE is installed. The server sends back to the user intermediate results which are then plotted as we show in Section IV.

A. Components

SABRE’s different components are activated as shown in Figure [2]. Depending on the chosen semantics, analysis mode and, if necessary, stochastic solution type, SABRE calls the corresponding method. Some of the functionalities are shared among different methods, for example the DTMC solution is accessed either directly from choosing the DTMC semantics, either indirectly, by the uniformization algorithm. As well, Runge-Kutta method, is used both as a solver of the CME or as the solver of the reaction rate equations.

B. Data Structure

We present an efficient data structure used by SABRE when used in stochastic analysis mode. SABRE’s main focus is on a fast implementation of the fast adaptive algorithm, so we will use this algorithm to motivate the choice of our data structure. However, the same kind of reasoning works if one wants to optimize the Runge-Kutta implementation. The most computationally demanding part of fast adaptive uniformization is the probability propagation phase, which performs the equivalent of one matrix-vector product in a DTMC. We therefore need a data structure that is efficient during this step.

First, we mention that, for each state, along with the state description, we need to record additional information about
the state space would have the following characteristics. Ideally, the data structure used for storing information about a case they are added to the state space data structure. Therefore, $n$ direct successors of $n$ to the state space as they are discovered. That is, some of the space has a single state, and that state is dynamically added along all of its outgoing transitions. Note that, initially the state is in the state space, and for each node we move probability mass to the probability of the state, about its successors, and about the rates/probabilities of the reactions that lead to those respective states.

We summarize the comparison between arrays and hashes in Table II. Arrays allow fast sequential access, fast add but slow search and delete operations. Hashes allow fast add, delete, search, but slow sequential access. We propose a hybrid solution that has the advantages of each data structure at the expense of extra memory usage.

Our hybrid data structure is composed of:

- **array** nodes that acts as a function from index $\rightarrow$ node
- **hash** index that acts as a function from state $\rightarrow$ index
- **vector** inactive_nodes of indices of nodes that have become inactive as a result of a delete.

This mixture of structures lets us give fast implementations for each of the required operations:

- **Sequential access** Simple iteration over the elements of nodes.
- **Search** Search within index followed by an access in nodes.
- **Delete state** The nodes array is allocated statically, so physically erasing a node would be expensive. The alternative is to mark the node for deletion by inactivating it – setting its probability to zero – and adding it to the inactive_nodes vector. Because of their zero probability, inactive nodes are ignored when iterating over all states. An inactive node has two possible futures: either it will be reoccupied by a newly added state, either it will be deleted during a compress phase. The compress phase is initiated when the number of inactive nodes covers more than 20% of the number of both active nodes and inactive nodes and it consists of eliminating all inactive nodes and rearranging the active nodes in a contiguous region.
- **Add state** When we add a state to the state space, we need to assign it to a node within the nodes array. The nodes array is allocated statically and during the program’s initialization phase, it is initialized to $2^{20}$ free nodes. When we add a new state, if inactive_nodes is non-empty, that is, if an inactive node exist, assign the state to this node, which now becomes active. If inactive_nodes is empty, we check whether we still have free allocated nodes, that is, we check whether the

- The probability of the state, about its successors, and about the rates/probabilities of the reactions that lead to those respective successors. We gather all this information in a structure called $\lambda$. During the propagate phase we iterate over all nodes of the state space, and for each node we move probability mass along all of its outgoing transitions. Note that, initially the state space has a single state, and that states are dynamically added to the state space as they are discovered. That is, some of the direct successors of $n$ may be newly discovered, and in this case they are added to the state space data structure. Therefore, ideally, the data structure used for storing information about the state space would have the following characteristics.

- **Fast sequential access.** For enumerating all nodes. We note that this is a property of the array primitive type of most programming languages.
- **Fast search.** For quickly finding the successors of a node. We note that this is a property of map or hash type of many programming languages.
- **Fast add.** For dynamically adding newly discovered states to the current state space.

- **[Fast delete.]** For dynamically removing states that have close to zero probability.

- We summarize the comparison between arrays and hashes in Table II. Arrays allow fast sequential access, fast add but slow search and delete operations. Hashes allow fast add, delete, search, but slow sequential access. We propose a hybrid solution that has the advantages of each data structure at the expense of extra memory usage.

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  - **Add state** When we add a state to the state space, we need to assign it to a node within the nodes array. The nodes array is allocated statically and during the program’s initialization phase, it is initialized to $2^{20}$ free nodes. When we add a new state, if inactive_nodes is non-empty, that is, if an inactive node exist, assign the state to this node, which now becomes active. If inactive_nodes is empty, we check whether we still have free allocated nodes, that is, we check whether the
number of active nodes has reached the size allocated to nodes. If free nodes exist, we assign the new state to a free node, if free nodes do not exist we need to allocate extra $2^{20}$ nodes to nodes and then pick a newly created free node. We note that the reallocation operation is expensive but happens only rarely, e.g. when the state space first reaches one million, two millions, three millions states and so on.

VI. CASE STUDIES

We present case studies for stochastic and deterministic analysis of CTMCs and for the stochastic analysis of DTMCs. For more and larger experiments on stochastic analysis of CTMCs we refer the reader to the paper giving the fast adaptive uniformization algorithm\cite{10}. All our experiments are performed on a 3GHz Intel Linux PC, with 6 GB of RAM. We give the results of our experiments in Table III.

| Analysis          | Model                  | Time | Error | States |
|-------------------|------------------------|------|-------|--------|
| Stochastic        | Exclusive switch       | 94s  | 9e-8  | 3047   |
| Deterministic     | Enzymatic reaction     | <1s  | -     | 1      |
| Stochastic        | Moran’s model          | 49s  | 0     | 1001   |

A. Genetic exclusive switch

The exclusive genetic switch we analyze involves two species of proteins that may bond to the same promoter site. We denote the unbonded proteins by $N_1$ and $N_2$ and the bonded ones by $r_1$ and $r_2$.\cite{4} The guarded commands for this model are given in Table IV. The rate functions are evaluated for the state $(x_{N_1}, x_{r_1}, x_{N_2}, x_{r_2})$, where $x_{N_1}$ is the number of molecules of type $N_1$ and so on.

When it is bonded to the promoter site, a protein represses the production of the other protein. And so, for example, production of $N_1$ only happens if no $N_2$ molecule is bounded to the promoter site (see rate function of first reaction). $N_1$ or $N_2$ may bond only to a free promoter site (see rate functions of the third and seventh reaction). Note that it always holds that $x_{r_1} + x_{r_2} \leq 1$.

We run the system from initial state $(25, 0, 0, 0)$ for a period of time of 10000 units with constants: $g_1 = g_2 = 0.05, d_1 = d_2 = 0.005, b_1 = b_2 = 0.1, u_1 = u_2 = 0.005$, and present the solution in Figure 3.

B. Enzymatic reaction

We use enzyme-catalyzed substrate conversion to exemplify how to perform a deterministic analysis under continuous-time semantics. The enzymatic reaction is described by three reactions (see Table V), that involve four chemical species, namely, enzyme ($E$), substrate ($S$), complex ($C$), and product ($P$) molecules. The state of the system is described by the vector $(x_E, x_S, x_C, x_P)$, which gives the existing number of molecules of each type.

For our experimental results, we chose the same parameters as in [6], that is, initial state $y = (1000, 100, 0, 0)$, time horizon $t = 70$, and rate constants $c_1 = c_2 = 1$ and $c_3 = 0.1$. For the case deterministic analysis we can not give any error bounds, as shown in Table III.

C. Moran’s population model

As a simple example of how SABRE operates on DTMC models we choose Moran’s genetic population model, which can be seen as a set of biochemical reactions, more specifically as one reversible reaction.

For a population of $N$ individuals, with two alleles, $A_1$ and $A_2$, we are interested to find the probability of fixation of $A_1$, that is, the probability for $A_1$ individuals to be equal to $N$ after a certain time. We have two reactions: $A_2 \rightarrow A_1$ and $A_1 \rightarrow A_2$. For $x_{A_1}$ individuals with $A_1$ allele and $x_{A_2}$ individuals with $A_2$ allele, the probability of the first reaction is $\frac{1}{2} + s \cdot \frac{x_{A_1}}{N}$, where $s$ is a small constant. As for the second reaction, its probability is $\frac{1}{2} + s \cdot \frac{x_{A_2}}{N}$.

We choose $N = 1000$ and $s = 2e - 3$, the initial state of $x_{A_1} = 1$ and we perform a transient analysis until time $k = 10^6$, at this time, the probability of fixation is 0.00049. In this case the error we obtain is 0 because no cutting is performed, the state space is kept at its complete size of 1001.
VII. COMPARISON WITH OTHER TOOLS

Several tools for stochastic analysis of Markov chains have been developed by communities such as probabilistic verification, computational biology and performance evaluation among others. Here, we provide a comparison with the tools that are the closest to SABRE. The PRISM tool [28], which is widely used in probabilistic verification, considers a more general class of Markov processes than population models. For instance, it does not restrict the update function such that it allows only a constant change of the state variables. The models addressed by PRISM are less structured and typically they do not have state dependent rate functions. PRISM uses powerful minimization techniques such as bisimulation that do not result in significant reductions in the case of population model. PRISM requires that upper bounds on the state variables are given as an input by the user. As opposed to that the SABRE tool finds appropriate bounds automatically and avoids an exhaustive state space exploration. The drawback is that the SABRE tool cannot guarantee the validity of properties such as “Is the probability to reach state $x$ within $t$ time greater than $p$?” but gives an approximate solution. As opposed to that PRISM can guarantee such properties. On the other hand, since SABRE avoids an exhaustive state space exploration it is able to handle much larger models with state-dependent rates. Infamy is a model-checking tool for infinite-state CTMCs by Zhang et al. [17]. Depending on the desired precision, their algorithm simply explores the reachable states up to a finite path depth. In contrast, our approach takes into account the direction into which the probability mass moves, and constructs a sequence of abstract models “on-the-fly” during the verification process. Similar approaches have also been used in the context of biochemical reaction networks [6].

Other tools for stochastic analysis of Markov chains, such as ETMCC [22], MRMC [26], APNtoolbox [5], SHARPE [23], SPNP [24], and Möbius [8], are conceived for answering performance analysis questions and as PRISM, due to their exhaustive state space exploration can not be applied to infinite models.

Dizzy [32], Snoopy [18] and Copasi [25] are tools for stochastic simulation alone and not do not compute probability distributions over states. Bio-PEPA [7] is a language for modeling and analysis of biochemical networks. For numerical analysis and verification problems Bio-PEPA uses PRISM’s engine.

VIII. CONCLUSION

We have introduced SABRE, a tool for stochastic analysis of biochemical reaction networks and of population models in general. We have motivated the choice of guarded commands as input formalism for our tool and the need for a stochastic analyzer specialized on biological systems. SABRE currently has the form of an accessible web tool, which was chosen out of the need to deliver our algorithms and optimizations in a fast and portable way. However, an offline version release is planned for the future. For completeness and comparison, SABRE also performs deterministic analysis of the input system.

ACKNOWLEDGMENT

We thank Marius Mateescu for valuable advices on the web interface and Nick Barton for an introduction to population genetics.

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