

Modeling Cellular Behavior with Hybrid Automata: Bisimulation and Collapsing

M. Antoniotti¹, B. Mishra^{1,2}, C. Piazza³, A. Policriti⁴, and M. Simeoni³

¹ Courant Institute of Mathematical Science, NYU, New York, U.S.A.

² Watson School of Biological Sciences, Cold Spring Harbor, New York, U.S.A.

³ Dept. of Computer Science, University Ca' Foscary of Venezia, Venezia, Italy

⁴ Dept. of Mathematics and Computer Science, University of Udine, Udine, Italy
marcoxa@cs.nyu.edu, mishra@cs.nyu.edu piazza@dsi.unive.it,
policrit@dimi.uniud.it simeoni@dsi.unive.it

Abstract. Many biological systems can be modeled using systems of ordinary differential algebraic equations (e.g., S-systems), thus allowing the study of their solutions and behavior automatically with suitable software tools (e.g., PLAS, Octave/Matlab[™]). Usually, numerical solutions (*traces* or *trajectories*) for appropriate initial conditions are analyzed in order to infer significant properties of the biological systems under study. When several variables are involved and the traces span over a long interval of time, the analysis phase necessitates automation in a scalable and efficient manner. Earlier, we have advocated and experimented with the use of automata and temporal logics for this purpose (XS-systems and Simpathica) and here we continue our investigation more deeply. We propose the use of *hybrid automata* and we discuss the use of the notions of *bisimulation* and *collapsing* for a “qualitative” analysis of the temporal evolution of biological systems. As compared with our previous proposal, hybrid automata allow maintenance of more information about the differential equations (S-system) than standard automata. The use of the notion of bisimulation in the definition of the *projection operation* (restrictions to a subset of “interesting” variables) makes possible to work with reduced automata satisfying the same formulae as the initial ones. Finally, the notion of collapsing is introduced to move toward still simpler and equivalent automata taming the complexity of the automata whose number of states depends on the level of approximation allowed.

1 Introduction

The emerging fields of *system biology* [21], and its sister field of bioinformatics, focuses on creating a finely detailed and “mechanistic” picture of biology at the cellular level by combining the part-lists (genes, regulatory sequences, other objects from an annotated genome, and known metabolic pathways), with observations of both transcriptional states of a cell (using micro-arrays) and translational states of the cell (using proteomics tools).

Recently, the need has arisen for more and more sophisticated and mathematically well founded computational tools capable to analyze the models that

are and will be at the core of *system biology*. Such computational models should be implemented in software packages faithfully while exploiting the potential trade-offs among usability, accuracy, and scalability dealing with large amounts of data. The work described in this paper is part of a much larger project still in progress, and thus only provides a partial and evolving picture of a new paradigm for computational biology.

Consider the following scenario. A biologist is trying to test a set of hypotheses against a corpus of data produced in very different ways by several *in vitro*, *in vivo*, and *in silico* experiments. The system the biologist is considering may be a piece of a *pathway* for a given organism. The biologist can access the following pieces of information:

- raw data stored somewhere about the temporal evolution of the biological system; this data may have been previously collected by *observing* an *in vivo* or an *in vitro* system, or by *simulating* the system *in silico*;
- some mathematical model of the biological system⁵.

The biologist will want to formulate *queries* about the evolution encoded in the data sets. For example, he/she may ask: *will the system reach a “steady state”?*, or *will an increase in the level of a certain protein activate the transcription of another?* Clearly the set of numerical *traces* of very complex systems rapidly becomes unwieldy to wade through for increasingly larger numbers of variables.

Eventually, many of these models will be available in large public databases (e.g. [6, 18–20, 28, 23]) and it is not inconceivable to foresee a biologist to test some hypotheses *in silico* before setting up expensive wet-lab experiments. The biologist will mix and match several models and raw data coming from the public databases and will produce large datasets to be analyzed.

To address this problem, we have proposed a set of theoretical and practical tools, XS-systems and Simpathica, that allow the biologist to formulate such queries in a simple way [3–5]. The computational tool Simpathica derives its expressiveness, flexibility, and power by integrating in a novel manner many commonly available tools from numerical analysis, symbolic computation, temporal logic, model-checking, and visualization. In particular, an *automaton-based* semantics of the temporal evolution of complex biochemical reactions starting from their representations as sets of differential equations is introduced. Then *propositional temporal logic* is used to qualitatively reason about the systems. Notice that here the adverb qualitatively does not mean that we completely abstract away from the quantities of the substances involved in the system, but that we concentrate on particular properties of the system, e.g. *will a certain protein reach level 0.5?*

In this paper we continue our research on the computational models at the core of our approach. We bring in several techniques from the fields of Verification, Logic and Control Theory, while maintaining a trade off between the need

⁵ We note that simulating a system *in silico* actually requires a mathematical model. However, we want to consider the case when such mathematical model is unavailable to both the biologist and the software system.

to manipulate large sets of incomplete data⁶ and the requirements arising from the needs to provide a mathematically well founded system. In particular, we propose the use of *hybrid automata* together with the notions of *bisimulation* and *collapsing*. Hybrid automata are equipped with states embodying time-flow, initial and final conditions, and therefore allow maintenance of more information about the differential equations (S-system). The use of the notion of bisimulation in the definition of the *projection operation* (restrictions to a subset of “interesting” variables) provides a way to introduce *reduced* automata satisfying the same formulae as the initial ones. Notice that the idea and potential behind this use of the notion of bisimulation can be exploited as fruitfully as here also in the context of standard automata. Finally, the notion of collapsing is introduced to improve along the direction of a qualitative study of the automata extracted from the analysis of traces as well as to tame its complexity of the automata (whose size depends on the level of approximation allowed).

The cellular and biochemical processes analyzed using XS-systems and Simpathica [4, 3] provide a large set of application examples for the framework we present here. In order to motivate the choices of our modeling framework, we give some details about one of such examples, the *repressilator system* described by Elowitz and Leibler in [13].

We conclude pointing out that the analysis presented in this paper is not limited to XS-Systems, but could be extended to more general hybrid system models.

2 Related Works

A survey on the different approaches for modeling and simulating genetic regulatory systems can be found in [11]: the author takes into consideration different mathematical methods (including ordinary and partial differential equations, qualitative differential equations and others) and evaluates their relative strengths and weaknesses.

The problem of constructing an automaton from a given mathematical model of a general dynamical system has been previously considered in the literature. In particular, it has been investigated by Brockett in [7]: our approach in [4] is certainly more focussed, since it deals with specific mathematical models (i.e. S-systems). Here we take the distance from pure discrete models, and propose the use of hybrid automata instead of standard automata, with the aim of taking advantage of their continuous component for allowing quantitative besides qualitative reasoning.

The use of hybrid automata for the modeling and simulation of biomolecular networks has been proposed also by Alur et. al. in [1]. In that paper the discrete component of an hybrid automaton is used to switch between two different behaviors (models) of the considered biological system, (for example) depending

⁶ The range of parameters tested in each *in vitro* and *in vivo* experiment is just too large, and coping with this combinatorial effects is a subject of research (c.f.r. [26]).

on the concentration of the involved molecules. In our case, the continuous component is used to regulate the permanence on a given state depending on the values of the involved variables (reactants), and the discrete component is used for enabling the transition to another state.

Moreover, in [1], as well as in other formalisms which model biochemical systems (e.g., [25, 10]), the notion of concurrency is explicitly used since the involved reactants are represented as processes running in parallel. In our case this kind of concurrency becomes implicit since in all the states of the automaton representing an S-system the values of all the reactants and their evolutions are represented.

3 Setting the Context

3.1 S-systems

We begin presenting the basic definitions and properties of S-systems. The definition of S-systems we use in this paper is basically the one presented in [27] augmented with a set of *algebraic constraints*. The constraints characterize the conditions under which a given set of equations is derived from a set of maps (see also [4]).

Definition 1 (S-system). *An S-system is a quadruple $S = (DV, IV, DE, C)$ where:*

- $DV = \{X_1, \dots, X_n\}$ is a finite non empty set of dependent variables ranging over the domains D_1, \dots, D_n , respectively;
- $IV = \{X_{n+1}, \dots, X_{n+m}\}$ is a finite set of independent variables ranging over the domains D_{n+1}, \dots, D_{n+m} , respectively;
- DE is a set of differential equations, one for each dependent variable, of the form

$$\dot{X}_i = \alpha_i \prod_{j=1}^{n+m} X_j^{g_{ij}} - \beta_i \prod_{j=1}^{n+m} X_j^{h_{ij}}$$

with $\alpha_i, \beta_i \geq 0$ called rate constants;

- C is a set of algebraic constraints of the form

$$C_j(X_1, \dots, X_{n+m}) = \sum (\gamma_j \prod_{k=1}^{n+m} X_k^{f_{jk}}) = 0$$

with γ_j called rate constraints.

In what follows we use \vec{X} to denote the vector $\langle X_1, \dots, X_n, X_{n+1}, \dots, X_{n+m} \rangle$ of variables and $\vec{d}(\vec{a}, \vec{b}, \dots)$ to denote the vector $\langle d_1, \dots, d_n, d_{n+1}, \dots, d_{n+m} \rangle \in D_1 \times \dots \times D_n \times D_{n+1} \times \dots \times D_{n+m}$ of values. Similarly given a set of variables $U = \{X_{U_1}, \dots, X_{U_u}\} \subseteq DV \cup IV$ we use $\vec{X} \upharpoonright U$ to denote the vector of variables of U , while $\vec{d} \upharpoonright U$ denotes the vector of values $\langle d_{U_1}, \dots, d_{U_u} \rangle \in D_{U_1} \times \dots \times D_{U_u}$.

The dynamic behavior of an S-system can be simulated by computing the approximate values of its variables at different time instants (*traces*). To determine a trace of an S-system it is necessary to fix an initial time (t_0), the values of the variables at the initial time ($\vec{X}(t_0)$), a final time (t_f), and a step (s).

Definition 2 (Trace). Let $S = (DV, IV, DE, C)$ be an S-system. Let $\vec{f}(t) = \langle f_1(t), \dots, f_{n+m}(t) \rangle$ be a (approximated) solution for the S-system S in the time interval $[t_0, t_f]$ starting with initial values $\vec{X}(t_0)$ in t_0 . Let $s > 0$ be a time step such that $t_f = t_0 + j * s$. The sequence of vectors of values

$$tr(S, t_0, \vec{X}(t_0), s, t_f) = \langle \vec{f}(t_0), \vec{f}(t_0 + s), \dots, \vec{f}(t_0 + (j - 1) * s), \vec{f}(t_0 + j * s) \rangle$$

is a trace of S . When we are not interested in the parameters defining the trace we use the notation tr .

Notice that $\vec{f}(t_0) = \vec{X}(t_0)$. A trace is nothing but a sequence of values of $D_1 \times \dots \times D_{n+m}$ representing a solution of the system in the time instants $t_0, t_0 + s, \dots, t_0 + j * s$. By varying the initial values of the variables, we obtain different system traces, for the same parameters t_0, s and t_f . Simulations of the behavior of an S-system can be automatically obtained by using the tool PLAS (see [27]). In fact, PLAS takes in input an S-system and approximates the values of the system variables, once the parameters in Definition 2 have been specified. The output is exactly a trace describing the behavior of the given system.

Example 1. The following feedback system is taken from [27], Chapter 6, and can be found in PLAS (see `\Book_Examples\Feedback.plc`). It represents a system in which the reactant X_1 is inhibited by X_2 , while X_3 is an independent input variable and X_4 an independent inhibitor for the degradation of X_2 . Hence, we have $DV = \{X_1, X_2\}$, $IV = \{X_3, X_4\}$, and

$$\dot{X}_1 = 0.5X_2^{-2}X_3^{0.5} - 2X_1 \quad \dot{X}_2 = 2X_1 - X_2^{0.5}X_4^{-1}$$

Let $t_0 = 0$ be the initial time, $\vec{X}(t_0) = \langle 1, 1, 4, 2 \rangle$ be the initial values of the reactants, $s = 1$ be the time step, and $t_f = 18$ be the final time. By simulating the system in PLAS with these values and setting the Taylor method with tolerance $1E - 16$ we obtain the following trace

$$\langle \langle 1, 1, 4, 2 \rangle, \langle 0.33, 1.59, 4, 2 \rangle, \langle 0.22, 1.48, 4, 2 \rangle, \dots \\ \dots, \langle 0.28, 1.31, 4, 2 \rangle, \langle 0.28, 1.31, 4, 2 \rangle, \langle 0.28, 1.31, 4, 2 \rangle \rangle$$

where we have not reported all the decimals and states for space reasons. In this trace, for instance, we can observe that the quantity of X_1 is 0.28 in the last three steps.

The solutions of an S-system have some nice properties. First of all they admit all the derivatives everywhere except when they intersect one of the hyperplane $X_i = 0$, for $i = 1, \dots, n + m$. There could be problems when $X_i = 0$ for $i \in \{1, \dots, m + n\}$ in the case one of the exponent is, for instance, of the form 0.5. As noticed in [1], this corresponds to the fact that at reasonably high molecular concentrations, one can adopt continuum models which lend themselves

to deterministic models, while at lower concentrations, the discrete molecular interactions become important and deterministic models are more difficult to obtain. However, the existence of all the derivatives implies that if at a given instant t_1 all the X_i , for $i = 1, \dots, n + m$, are different from 0, then there exists a unique solution in an interval $[t_1, t_1 + \epsilon]$ and this solution can be extended if it still holds that all the variables are different from 0. Moreover, if two solutions $\vec{f}(t)$ and $\vec{g}(t)$, obtained with different initial values, pass both in a point \vec{d} , possibly at different times, i.e., there exist two instants t_1 and t_2 such that $\vec{f}(t_1) = \vec{g}(t_2) = \vec{d}$, then from those instants on they always coincide, i.e., for all $p \geq 0$, $\vec{f}(t_1 + p) = \vec{g}(t_2 + p)$. This is a consequence of the fact that the *variable* time does not explicitly occurs in the differential equations. What we have just stated in mathematical terms can be restated from the biological point of view saying that if the biological system modeled by the S-system reaches a state \vec{d} , its evolution does not depend on the states in which the system was before reaching \vec{d} (i.e., the system is *without memory*). In particular, on a set of traces this last property has the following consequence.

Proposition 1. *Let $\langle \vec{a}_0, \dots, \vec{a}_j \rangle$ and $\langle \vec{b}_0, \dots, \vec{b}_i \rangle$ be two traces of an S-system S obtained by using the same time step s . If there exist h and k such that $\vec{a}_h = \vec{b}_k$, then for all $r \geq 0$ it holds $\vec{a}_{h+r} = \vec{b}_{k+r}$.*

Obviously in the above proposition we are assuming that we are using the same approximation method to obtain both traces. Moreover, it can be the case that the two traces are equal. This property of sets of traces of an S-system implies what is known in the area of Model Checking as *fusion closure* (see [14]). We anticipate here that all the results we present in the rest of this paper are consequences of Proposition 1, i.e., they hold every time we deal with a set of traces satisfying it. We formalize this as follows.

Definition 3 (Convergence). *A set of traces Tr is convergent if for all the traces $\langle \vec{a}_0, \dots, \vec{a}_j \rangle$ and $\langle \vec{b}_0, \dots, \vec{b}_i \rangle$ belonging to Tr , if there exist h and k such that $\vec{a}_h = \vec{b}_k$, then for all $r \geq 0$ it holds $\vec{a}_{h+r} = \vec{b}_{k+r}$.*

Corollary 1. *If Tr is a set of traces of an S-system S obtained by using the same time step s , then Tr is convergent.*

Example 2. Let us consider again the simple feedback system described in Example 1. If we simulate it using $\vec{X}(t_0) = \langle 0.33, 1.59, 4, 2 \rangle$, i.e., $\vec{X}(t_1)$ of the trace in Example 1, we obtain

$\langle \langle 0.33, 1.59, 4, 2 \rangle, \langle 0.22, 1.48, 4, 2 \rangle, \dots, \langle 0.28, 1.31, 4, 2 \rangle, \langle 0.28, 1.31, 4, 2 \rangle \rangle$
which is exactly the trace we had before without the first state.

3.2 XS-systems

The basic idea of XS-systems (introduced in [4]) is to associate an S-system S with a finite automaton, obtained by suitably encoding a set of traces on S .

Essentially, each trace on S can be encoded into a simple automaton, where states correspond to the trace elements (i.e., the values of the system variables observed at each step), and transitions reflect the sequence structure of the trace itself (i.e., there exists a transition from a state v_i to a state v_j if they are consecutive in the trace). When more than one trace is involved in the process, coinciding elements of different traces correspond to the same state in the automaton.

Consider an S-system and a set of traces on it: the automaton derived from the system traces is defined as follows.

Definition 4 (S-system Automaton). *Let S be an S-system and Tr be a set of traces on S . An S-system automaton is $\mathcal{A}(S, Tr) = (V, \Delta, I, F)$, where*

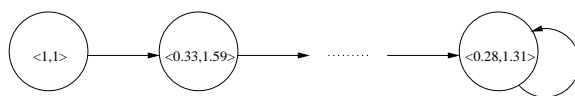
- $V = \{\vec{v} = \langle v_1, \dots, v_{n+m} \rangle \mid \exists tr \in Tr : \vec{v} \text{ is in } tr\} \subseteq D_1 \times \dots \times D_{n+m}$ is the set of states;
- $\Delta = \{(\vec{v}, \vec{w}) \mid \exists tr \in Tr : \vec{v}, \vec{w} \text{ are consecutive in } tr\}$ is the transition relation;
- $I = \{\vec{v} \mid \exists tr \in Tr : \vec{v} \text{ is initial in } tr\} \subseteq V$ is the set of initial states;
- $F = \{\vec{v} \mid \exists tr \in Tr : \vec{v} \text{ is final in } tr\} \subseteq V$ is the set of final states.

Automata can be equipped with labels on nodes and/or edges (see [17]). Labels on the nodes maintain information about the properties of the nodes, while labels on the edges are used to impose conditions on the action represented by the edge (see [8]). In the case of S-system automata edges are unlabelled, while the label we assign to each node is actually the name (identifier) of the node itself, i.e. the concentrations of the reactants for that state. In this way S-system automata maintain qualitative information about the system only in the instants corresponding to the steps.

We say that an automaton is *deterministic* if each node has at most one outgoing edge for each edge-label, i.e., in our case, at most one outgoing edge. From Proposition 1 we get the following result.

Proposition 2. *Let S be an S-system and Tr be a convergent set of traces on S . The automaton $\mathcal{A}(S, Tr)$ is deterministic.*

Example 3. The trace shown in Example 1 gives us the following automaton, where we omit the values of the independent variables.



The initial state is the one on the left, while final state is the one on the right. By using both the trace of Example 1 and the trace of Example 2 we obtain the same automaton, but with two initial states. The automaton represents the fact that all the simulations of this feedback system with initial values of the reactants equal to the values in one of the states of the automaton reach a steady state in which $X_1 = 0.28$ and $X_2 = 1.31$.

In [4], a language called ASySA (*Automata S-systems Simulation Analysis* language) has been presented to inspect and formulate queries on the simulation results of XS-systems. The aim of this language is to provide the biologists with a tool to formulate various queries against a repository of simulation traces. ASySA is essentially a *Temporal Logic* language (see [14]) (an English version of CTL) with a specialized set of predicate variables whose aim is to ease the formulation of queries on numerical quantities. The fusion closure of sets of traces (see Proposition 1 and Corollary 1) is necessary in order to reflect the behavior of the set of traces with temporal logic semantics (see [14]). This means that a formula is true on the S-system automaton if and only if it is true in the set of traces. Intuitively, the behavior of the traces is not approximated in the automaton because two traces which reach the same state always coincide in the future.

Example 4. The automaton in Example 3 satisfies the formula

$$\text{EVENTUALLY}(\text{ALWAYS}(X_2 > 1))$$

which means that the system admits a trace such that, from a certain point on, X_2 is always greater than 1. Similarly, it does not satisfies the formula

$$\text{ALWAYS}(\text{EVENTUALLY}(X_1 > X_2))$$

since it reaches a steady state in which X_1 is less than X_2 .

Unfortunately, in the practical cases the automata built from sets of traces have an enormous number of states. In [4] two techniques have been proposed to reduce the number of states of an S-system automaton, namely *projection* and *collapsing*.

Definition 5 (Projection). *Let S be an S-system and U be a subset of the set of variables of S . Given a trace $tr = \langle \vec{a}_0, \dots, \vec{a}_j \rangle$ of S the projection over U of tr is the sequence $tr \upharpoonright U = \langle \vec{a}_0 \upharpoonright U, \dots, \vec{a}_j \upharpoonright U \rangle$. Given a set of traces Tr the projection over U of Tr is the set of projected traces $Tr \upharpoonright U = \{tr \upharpoonright U \mid tr \in Tr\}$. The U -projected S-system automaton from Tr and S is $\mathcal{A}(S, Tr \upharpoonright U)$.*

The automaton $\mathcal{A}(S, Tr \upharpoonright U)$ has usually less states than $\mathcal{A}(S, Tr)$. However, the set of traces $Tr \upharpoonright U$ does not always satisfies neither convergence nor fusion closure and the automaton $\mathcal{A}(S, Tr \upharpoonright U)$ can be non-deterministic. This can introduce an approximation, i.e., the formulae satisfied by the automaton $\mathcal{A}(S, Tr \upharpoonright U)$ are not the same satisfied by the set of traces $Tr \upharpoonright U$.

Example 5. As a simple yet very interesting example, consider the repressilator system constructed by Elowitz and Leibler [13]. First the authors constructed a mathematical model of a network of three interacting transcriptional regulators and produced a trace of the interaction using a traditional mathematical package (Matlabtm). Subsequently, they constructed a plasmid with the three regulators and collected data from in vivo experiments in order to match them with the

predicted values. In particular, this contains three proteins, namely `lacI` (X_1), `tetR` (X_2), and `cl` (X_3). The protein `lacI` represses the protein `tetR`, `tetR` represses `cl`, whereas `cl` represses `lacI`, thus completing a feedback system. The dynamics of the network depend on the transcription rates, translation rates, and decay rates. Depending on the values of these rates the system might converge to a stable limit circle or become unstable. The following S-system represents⁷ the repressilator system: rate values have been set in such a way that the system converges to a stable limit circle.

$$\begin{aligned}\dot{X}_1 &= X_4 X_3^{-1} - X_1^{0.5} \\ \dot{X}_2 &= X_5 X_1^{-1} - X_2^{0.578151} \\ \dot{X}_3 &= X_6 X_2^{-1} - X_3^{0.5}\end{aligned}$$

If we simulate it in `PLAS`, with $t_0 = 0$, $\vec{X}(t_0) = \langle 0.01, 0.2, 0.01, 0.2, 0.2, 0.2 \rangle$, $s = 0.05$, and $t_f = 30$, we obtain a trace whose automaton reaches the loop shown on the left of Figure 1: we omit the independent variables and we use dotted lines to represent the fact that there are other intermediate states.

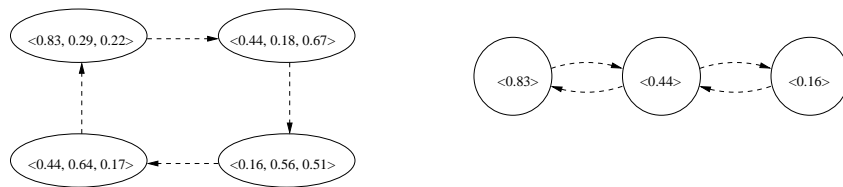


Fig. 1. Repressilator: automaton and projected automaton.

The automaton does not satisfy `EVENTUALLY(ALWAYS($X_1 \geq 0.3$))`. In fact in the limit circle reached by the repressilator, the values of X_1 range in the interval $[0.16, 0.83]$. Hence, the formula is false also in the projected trace. However, the formula is satisfied by the projected automaton, partially depicted on the right of Figure 1. In fact, the projected automaton represents a system in which it is possible that after a certain instant the variable X_1 assumes values in the interval $[0.44, 0.83]$.

The collapsing operation is defined in such a way that a state is removed from a trace when it behaves similarly to the previous one, i.e., when the derivatives computed in it can be approximated by the derivatives computed in the previous state (see [4] for the formal definition). Also this operation can introduce approximation as shown in the following example.

⁷ To be precise the system described in [13] is not an S-system. However, it can be reasonably approximated through an S-system, as proved by the general theory presented in [27]. Notice that our automaton-model can be built using directly traces of the system in [13].

Example 6. Let S be an S-system with dependent variables X_1 and X_2 . Let us assume that S admits a trace of the form $\langle \langle 1, 5 \rangle, \langle 2, 4 \rangle, \langle 3, 3 \rangle, \langle 4, 2 \rangle, \langle 5, 1 \rangle \rangle$.

We also assume that the derivative \dot{X}_1 is 1 in all the states except the last one, and, similarly, \dot{X}_2 is -1 in all the states except the last one. By applying the definitions presented in [4] we can collapse some of the states obtaining the reduced trace $\langle \langle 1, 5 \rangle, \langle 5, 1 \rangle \rangle$. The formula $\text{EVENTUALLY}(|X_1 - X_2| \leq 3)$ is true in the trace of S , but is false in the collapsed one.

Consider again the repressilator system of Example 5, whose automaton is partially represented on the left of Figure 1. If all the intermediate states on the dotted lines are collapsed, then we obtain an automaton with 4 states which does not satisfy the formula $\text{EVENTUALLY}(|X_1 - X_2| \leq 0.1)$, while it is easy to check that the same formula is satisfied by the repressilator system.

In order to avoid these approximations and to obtain a more powerful and flexible framework in the next sections we propose the use of hybrid automata together with a reformulation of projection and collapsing.

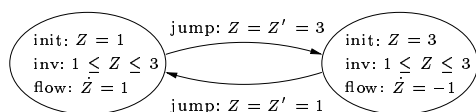
4 Hybrid Automata to model S-systems

The notion of hybrid automata has been first introduced in [2] as a model and specification language for hybrid systems, i.e., systems consisting of a discrete program within a continuously changing environment.

Definition 6 (Hybrid automata). An hybrid automaton $H = (Z, V, \Delta, I, F, \text{init}, \text{inv}, \text{flow}, \text{jump})$ consists of the following components:

- $Z = \{Z_1, \dots, Z_k\}$ a finite set of variables; $\dot{Z} = \{\dot{Z}_1, \dots, \dot{Z}_k\}$ denotes the first derivatives during continuous change; $Z' = \{Z'_1, \dots, Z'_k\}$ denotes the values at the end of discrete change;
- (V, Δ, I, F) is an automaton; the nodes of V are called control modes, the edges of Δ are called control switches;
- each $v \in V$ is labeled by $\text{init}(v)$, $\text{inv}(v)$, and $\text{flow}(v)$; the labels $\text{init}(v)$ and $\text{inv}(v)$ are constraints whose free variables are in Z ; the label $\text{flow}(v)$ is a constraints whose free variables are in $Z \cup \dot{Z}$;
- each $e \in E$ is labeled by $\text{jump}(e)$, which is a constraint whose free variable are in $Z \cup Z'$.

Example 7. Consider the following simple hybrid automaton.



The initial state is the left one, with $Z = 1$. In this state Z grows with constant rate 1. After 3 instants we have $Z = 3$ and we jump on the right state. In this second state Z decreases and when Z becomes 1 we jump again in the state on the left.

The usefulness of hybrid automata has been widely proved in the area of verification (see, e.g., [22]). In order to exploit the expressive power of hybrid automata their properties have been deeply studied (see [15]) and model checkers have been developed to automatic test temporal logic properties on them. Among the model checkers which deal with hybrid automata we mention HyTech (see [16]) developed at Berkeley University, Charon (see [1]) developed at the University of Pennsylvania.

In the S-system automata introduced in the previous section the only quantitative information maintained is the values of the variables in the instants corresponding to the steps. The values at instants between two steps were lost. This becomes particularly dangerous when we intend to apply a reduction operation like the collapsing one. Our idea here is to use the continuous component of hybrid automata to maintain also some approximate information about the values of the variables between two steps.

Let us introduce some notations which simplify the definition of an hybrid automaton modeling a convergent set Tr of traces of an S-system. Given the vectors $\vec{X} = \langle X_1, \dots, X_{n+m} \rangle$ and $\vec{v} = \langle v_1, \dots, v_{n+m} \rangle$ we use the notation $\vec{X} = \vec{v}$ to denote the conjunction $X_1 = v_1 \wedge \dots \wedge X_{n+m} = v_{n+m}$. The notation $\vec{v} \leq \vec{X} \leq \vec{w}$ has a similar meaning, while $\vec{X} = (\vec{w} - \vec{v})/s$ stands for $X_1 = (w_1 - v_1)/s \wedge \dots \wedge X_{n+m} = (w_{n+m} - v_{n+m})/s$.

Definition 7 (S-system Hybrid Automaton). *Let S be an S-system and Tr be a convergent set of traces on S . Consider the S-system automaton $\mathcal{A}(S, Tr)$. The S-system hybrid automaton built on $\mathcal{A}(S, Tr)$ is $\mathcal{H}(S, Tr) = (X, V, \Delta, I, F, init, inv, flow, jump)$, where:*

- $X = \{X_1, \dots, X_{n+m}\} = DV \cup IV$;
- (V, Δ, I, F) is the automaton $\mathcal{A}(S, Tr)$;
- for each $\vec{v} \in V$ let $init(\vec{v}) = \vec{X} = \vec{v}$;
- for each $\vec{v} \in V$ such that $(\vec{v}, \vec{w}) \in \Delta$ let⁸ $inv(\vec{v}) = \vec{v} \leq \vec{X} \leq \vec{w}$;
- for each $\vec{v} \in V$ such that $(\vec{v}, \vec{w}) \in \Delta$ let $flow(\vec{v}) = \vec{X} = (\vec{w} - \vec{v})/s$;
- for each $(\vec{v}, \vec{w}) \in \Delta$ let $jump((\vec{v}, \vec{w})) = \vec{X} = \vec{X}' = \vec{w}$.

Notice from the above definition that being in a state \vec{v} does not necessarily mean that the values of the variables are exactly \vec{v} : they can in fact assume values between \vec{v} and \vec{w} . In particular, they grow linearly in this interval and when they reach \vec{w} the system jump on a new state.

The automaton $\mathcal{H}(S, Tr)$ is a rectangular singular automaton and the temporal logic CTL is decidable for this class of automata (see [15]). The model checker HyTech can be used to check whether a temporal formula is satisfied by $\mathcal{H}(S, Tr)$. Moreover, $\mathcal{H}(S, Tr)$ is deterministic, since we require Tr to be convergent and hence $\mathcal{A}(S, Tr)$ is deterministic. Notice also that all the information needed to build $\mathcal{H}(S, Tr)$ is already encoded in $\mathcal{A}(S, Tr)$, i.e., it is possible to work on $\mathcal{H}(S, Tr)$ by only maintaining in memory $\mathcal{A}(S, Tr)$.

⁸ We invert the interval when $w_i < v_i$.

Example 8. From the traces of the feedback system of Examples 1 and 2 we obtain the hybrid automaton shown in Figure 8. In the first state (the one on

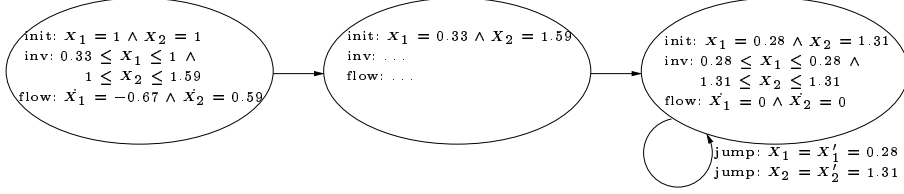


Fig. 2. Feedback hybrid automaton.

the left) variable X_1 starts with value 1 and decreases until it reaches value 0.33, while variable X_2 starts with value 1 and grows until it reaches value 0.59. Then, we jump on the second state. When we reach the last state the values of the variables become stable and the system loops forever.

The additional quantitative information stored in each state of an S-system hybrid automaton allow to deeply investigate the behavior of the system in the instants within a step. This becomes really relevant when we apply a collapsing technique to reduce the number of states as we will see in Section 6.

5 Bisimulation and Projection

As pointed out in Example 5 the projection operation can cause incorrect prediction on the reduced automaton. In order to avoid this problem, we define in this section a projection operator based on the notion of bisimulation. Since bisimulation is an equivalence relation preserving temporal logic formulae (see, e.g., [9]), the obtained projected automata will satisfy the same formulae of the original one.

Let us fix some notations. Given a condition $init(\vec{v})$ ($inv(\vec{v})$, $flow(\vec{v})$, resp.) and $U \subseteq DV \cup IV$ we use $init(\vec{v}) \upharpoonright U$ ($inv(\vec{v}) \upharpoonright U$, $flow(\vec{v}) \upharpoonright U$, resp.) to denote that we consider only the conditions relative to the variables in U .

Definition 8 (U -bisimulation). Let $\mathcal{H}(S, Tr)$ be an S-system hybrid automaton. Let $U \subseteq \{X_1, \dots, X_{n+m}\}$ be a subset of variables. A relation $R \subseteq V \times V$ is a U -bisimulation if

- if $\vec{v}R\vec{w}$, then $init(\vec{v}) \upharpoonright U = init(\vec{w}) \upharpoonright U \wedge inv(\vec{v}) \upharpoonright U = inv(\vec{w}) \upharpoonright U \wedge flow(\vec{v}) \upharpoonright U = flow(\vec{w}) \upharpoonright U$;
- if $\vec{v}R\vec{w}$ and $(\vec{v}, \vec{v}') \in \Delta$, then $(\vec{w}, \vec{w}') \in \Delta$ and $\vec{v}'R\vec{w}'$;
- if $\vec{v}R\vec{w}$ and $(\vec{w}, \vec{w}') \in \Delta$, then $(\vec{v}, \vec{v}') \in \Delta$ and $\vec{v}'R\vec{w}'$.

The idea is that two states \vec{v} and \vec{w} are U -bisimilar if not only the variables in U have the same values in \vec{v} and \vec{w} , but from \vec{v} and \vec{w} the system evolves in the

same way with respect to the variables in U . In fact, for instance, it is possible that there are two states in which the variables in U have the same values, but the first state evolves into a state in which the variables are incremented while the second one evolves into a state in which the variable are decremented. Intuitively we do not want to consider equivalent these two states.

Lemma 1. *There always exists a unique maximum U -bisimulation \approx_U which is an equivalence relation. Moreover, if $\vec{v} \approx_U \vec{w}$ and $(\vec{v}, \vec{v}') \in \Delta$, then $(\vec{w}, \vec{w}') \in \Delta$ and $\text{jump}((\vec{v}, \vec{v}')) \upharpoonright U = \text{jump}((\vec{w}, \vec{w}')) \upharpoonright U$.*

Proof. The first part follows immediately from the fact that a U -bisimulation on $\mathcal{H}(S, Tr)$ is nothing but a strong bisimulation on $\mathcal{A}(S, Tr)$ whose nodes have been labeled using part of the conditions defining the hybrid automaton $\mathcal{H}(S, Tr)$.

The second fact is a consequence of the fact that *jump* is uniquely defined once we know *init* and *inv*. \square

Definition 9 (Projected Hybrid automaton $\mathcal{H}(S, Tr, U)$). *Let $\mathcal{H}(S, Tr) = (X, V, \Delta, I, F, \text{init}, \text{inv}, \text{flow}, \text{jump})$, be an S -system hybrid automaton and U be a subset of variables. The projected hybrid automaton $\mathcal{H}(S, Tr, U) = (U, V_U, \Delta_U, I_U, F_U, \text{init}_U, \text{inv}_U, \text{flow}_U, \text{jump}_U)$ is defined as follows:*

- $V_U = V / \approx_U$;
- $\Delta_U = \{([\vec{v}], [\vec{w}]) \mid \exists \vec{v}' \in [\vec{v}], \vec{w}' \in [\vec{w}] : (v, w) \in \Delta\}$;
- for each $[\vec{v}] \in V_U$ let $\text{init}_U([\vec{v}]) = \text{init}(\vec{v}) \upharpoonright U$;
- for each $[\vec{v}] \in V_U$ let $\text{inv}_U([\vec{v}]) = \text{inv}(\vec{v}) \upharpoonright U$;
- for each $[\vec{v}] \in V_U$ let $\text{flow}_U([\vec{v}]) = \text{flow}(\vec{v}) \upharpoonright U$;
- for each $([\vec{v}], [\vec{w}]) \in \Delta_U$ such that $(\vec{v}', \vec{w}') \in \Delta$ let $\text{jump}_U(([\vec{v}], [\vec{w}])) = \text{jump}((\vec{v}', \vec{w}')) \upharpoonright U$.

The above definition does not depend on the representative element of each class. This is a consequence of the definition of \approx_U as far as the *init*, *inv*, and *flow* conditions are concerned, and of Lemma 1 as far as the *jump* conditions are concerned. For those who are familiar with automata and bisimulation reductions we can say that the hybrid automaton $\mathcal{H}(S, Tr, U)$ is nothing but the hybrid automaton built on the bisimulation reduced automaton $\mathcal{A}(S, Tr) / \approx_U$ with conditions defined only on the variables of U .

The automaton $\mathcal{H}(S, Tr, U)$ is still a rectangular singular automaton, hence CTL is still decidable on it. Moreover, $\mathcal{H}(S, Tr, U)$ is deterministic, since bisimulation preserves determinism. The fact that we are working on deterministic automata implies that the bisimulation relation \approx_U can be computed in linear (see [12]) time using the procedure defined in [24].

As far as the correctness of the reduction is involved, we have the following result.

Proposition 3. *Let TL be a temporal logic which is a fragment of the μ -calculus. Let φ a formula of TL involving only the variables in U . $\mathcal{A}(S, Tr)$ satisfies φ if and only if $\mathcal{A}(S, Tr) / \approx_U$ satisfies φ . $\mathcal{H}(S, Tr)$ satisfies φ if and only if $\mathcal{H}(S, Tr, U)$ satisfies φ .*

Proof. The first part is a consequence of the fact that \approx_U is a strong bisimulation and strong bisimulations preserve all the formulae of the μ -calculus.

The second part is a consequence of the first part and of the fact that $\mathcal{H}(S, Tr, U)$ is basically the hybrid automaton built on $\mathcal{A}(S, Tr)/\approx_U$. \square

In the following example we show the difference between $\mathcal{A}(S, Tr \upharpoonright U)$ and $\mathcal{A}(S, Tr)/\approx_U$. This difference is at the basis of the correctness of $\mathcal{H}(S, Tr, U)$.

Example 9. Consider again the repressilator system of Example 5. Part of the projected automaton we obtain by applying bisimulation is shown on the left of Figure 3. The two states in which $X_1 = 0.44$ do not coincide when we use bisimulation. In fact, the first state in which X_1 is 0.44 evolves to a state in which X_1 is 0.43 (the protein is decreasing), while the second state in which X_1 is 0.44 evolves to a state in which X_1 is 0.47 (the protein is growing). Hence, the projected automaton does not satisfy the formula $\text{EVENTUALLY}(\text{ALWAYS}(X_1 \geq 0.3))$. This is correct, since the repressilator system we are simulating reaches a steady loop in which X_1 oscillates between 0.16 and 0.83. Part of the projected

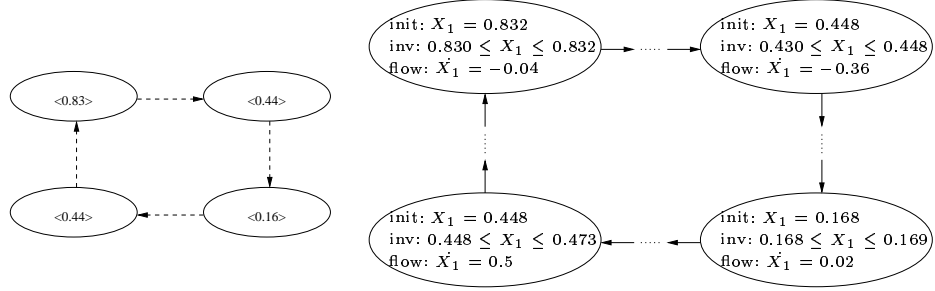


Fig. 3. Repressilator: bisimulation quotiented automata.

hybrid automaton is shown on the right of Figure 3.

6 Collapsing States

In this section we introduce the definition of collapsing of a trace. The definition we present is similar but not equivalent to the one given in [4]. In fact, we do not consider the difference between the derivatives calculated in the states, but the degree of growth within a step. This is inspired by hybrid automaton in which in the *flow* condition of a state we do not use the derivatives calculated at the beginning of a time step, but the coefficients of the lines connecting the values at the beginning to the ones at the end of a time step. In the following collapsing definition we use a compact notation similar to the one already introduced in Section 4.

Definition 10 (Collapsing). Let $\vec{\delta} = \langle \delta_1, \dots, \delta_{n+m} \rangle$ be a $n + m$ -vector of values. Let $tr = \langle \vec{a}_0, \dots, \vec{a}_j \rangle$ be a trace obtained by simulating the S -system S with time step s . A $\vec{\delta}$ -collapsing of tr is a partition of the states of tr such that:

- the blocks are sub-traces of tr ;
- if a block is formed by the states from \vec{a}_i to \vec{a}_{i+h} , and \vec{a}_j, \vec{a}_{j+1} belong to the block, then $|(\vec{a}_{j+1} - \vec{a}_j)/s - (\vec{a}_{i+1} - \vec{a}_i)/s| \leq \vec{\delta}$.

The collapsing operation in [4] is based on the difference between the first derivatives computed in the elements of the trace. Here, instead, we consider as collapsing criterion the degree of growth within a step. In practice the definition requires that the lines connecting \vec{a}_i to \vec{a}_{i+1} are good approximations of the lines connecting \vec{a}_j to \vec{a}_{j+1} . As a consequence we obtain that the lines connecting \vec{a}_i to \vec{a}_{i+h} are good approximations of all the small lines. In particular, the following result holds.

Lemma 2. *If a block of a $\vec{\delta}$ -collapsing is formed by the sequence of states from \vec{a}_i to \vec{a}_{i+h} and \vec{a}_j, \vec{a}_{j+1} belong to the block, then $|(\vec{a}_{j+1} - \vec{a}_{j+r})/s - (\vec{a}_{i+h} - \vec{a}_i)/(h*s)| \leq 2 * \vec{\delta}$.*

Proof. It is not restrictive to prove that the result holds on the first component. Let $(a_{i+r+1,1} - a_{i+r,1})/s = \text{coef}_{r+1}$ for $r = 0, \dots, h-1$. It is easy to prove that $(a_{i+h,1} - a_{i,1})/(h*s) = (1/h)*((a_{i+h,1} - a_{i+h-1,1})/s + (a_{i+h-1,1} - a_{i+h-2,1})/s + \dots + (a_{i+1,1} - a_{i,1})/s) = (1/h)*\sum_{r=0}^{h-1} \text{coef}_{r+1}$. By hypothesis we have $\text{coef}_1 - \delta_1 \leq \text{coef}_{r+1} \leq \text{coef}_1 + \delta_1$, hence we obtain $(1/h)*h*(\text{coef}_1 - \delta_1) \leq (a_{i+h,1} - a_{i,1})/(h*s) \leq (1/h)*h*(\text{coef}_1 + \delta_1)$, i.e. $\text{coef}_1 - \delta_1 \leq (a_{i+h,1} - a_{i,1})/(h*s) \leq \text{coef}_1 + \delta_1$. From this last we get $(\text{coef}_1 - \delta_1) - (\text{coef}_1 + \delta_1) \leq \text{coef}_{r+1} - (a_{i+h,1} - a_{i,1})/(h*s) \leq (\text{coef}_1 + \delta_1) - (\text{coef}_1 - \delta_1)$, i.e. $-2*\delta_1 \leq \text{coef}_{r+1} - (a_{i+h,1} - a_{i,1})/(h*s) \leq 2*\delta_1$, which is equivalent to our thesis. \square

Given the trace tr and the vector $\vec{\delta}$ the partition in which each state constitutes a singleton class is a $\vec{\delta}$ -collapsing.

Definition 11 (Maximal Collapsing). *Let C_1 and C_2 be two $\vec{\delta}$ -collapsing of tr . We say that C_1 is coarser than C_2 if each block of C_2 is included in a block of C_1 . We say that the $\vec{\delta}$ -collapsing C_1 is maximal if there does not exist another $\vec{\delta}$ -collapsing coarser than C_1 .*

The uniqueness of a coarsest $\vec{\delta}$ -collapsing is not guaranteed. However, we can give an algorithm to find a maximal $\vec{\delta}$ -collapsing. The algorithm performs the following steps: it starts from \vec{a}_0 , it check if \vec{a}_1 can be collapsed with \vec{a}_0 , if this is the case it goes on with \vec{a}_2 , and so on. Assume that \vec{a}_i is the first state which does not collapse with \vec{a}_0 , then the algorithm starts another block from \vec{a}_i and it goes on in the same way.

The following proposition states that if we use maximal δ -collapsing, then two traces which match in one state always match in the future.

Proposition 4. *Let Tr be a convergent set of traces of an S -system S . Let $Tr/\vec{\delta}$ be the set of collapsed traces obtained by applying to each trace of Tr a maximal $\vec{\delta}$ -collapsing. The set $Tr/\vec{\delta}$ is convergent.*

This property is sufficient to guarantee that taking a set of traces and collapsing them using maximal δ -collapsing, the set of collapsed traces can be used to build automata and hybrid automata as defined in the previous sections. In fact, as pointed out along the paper, the correctness of our framework holds whenever we use convergent sets of traces. This does not mean that the automaton we build from a set of collapsed traces satisfies the same formulae of the original set of traces, but that it satisfies the same formulae of the set of collapsed of traces.

Example 10. Consider again the S-system and the trace of Example 6. The collapsed trace we obtain is again $\langle \langle 1, 5 \rangle, \langle 5, 1 \rangle \rangle$. The hybrid automaton we build from this trace has two states. In the first state, let us call it \vec{v} , we have the conditions

$$inv(\vec{v}) = 1 \leq X_1 \leq 5 \wedge 1 \leq X_2 \leq 5 \quad flow(\vec{v}) = \dot{X}_1 = 1 \wedge \dot{X}_2 = -1$$

which make $EVENTUALLY(|X_1 - X_2| \leq 3)$ true in the automaton, as it was in the original trace.

Similarly, we can safely collapse the states of the repressilator system and obtain an hybrid automaton with four states which correctly satisfy the formula $EVENTUALLY(|X_1 - X_2| \leq 0.1)$.

This last example shows that the additional information maintained in the hybrid automaton is particularly useful when we use techniques as collapsing to reduce the number of states.

7 Conclusions

In this paper we have described how hybrid automata can be used to model and analyze set of traces representing the behavior of a biological system. Automata give a qualitative view of a set of traces by abstracting from the time instants, thus allowing a compact representation in which the properties of the system can be easily investigated. The use of hybrid automata, instead of standard ones, improves along the direction of a qualitative, but *complete*, modeling of the biological system. In fact, powerful techniques such as (bisimulation-)projections and collapsing can be “safely” applied to hybrid automata in order to reduce the number of states. In particular, while the bisimulation based projection we present could be applied also to standard automata, the “good” behavior of the collapsing operation w.r.t. the verification of temporal formulae strongly depends on the information which is stored in each state of hybrid automata. Notice that, although we have presented a construction of hybrid automata from standard S-systems, it is not difficult to modify our framework in order to deal with more complicated systems, e.g., systems whose differential equations change during the evolution of the system itself.

As future work, we intend to extend our tool set in order to (1) integrate some Time-Frequency analysis tools to identify the XS-system states by analyzing the

traces, and (2) incorporate a *learning* scheme in our approach that keeps track of what automata (read: model) is being manipulated at a given stage of the analysis carried out by a biologist, and that amends it semi-automatically before stepping in the next stage. This could be particularly useful in the evaluation phase for the constant rates involved in the system.

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