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Attractor computation using interconnected Boolean networks: testing growth rate models in *E. Coli*

Madalena Chaves and Alfonso Carta*

Abstract

Boolean networks provide a useful tool to address questions on the structure of large biochemical interactions since they do not require kinetic details and, in addition, a wide range of computational tools and algorithms is available to exactly compute and study the dynamical properties of these models. A recently developed method has shown that the attractors, or asymptotic behavior, of an asynchronous Boolean network can be computed at a much lower cost if the network is written as an interconnection of two smaller modules. We have applied this methodology to study the interconnection of two Boolean models to explore bacterial growth and its interactions with the cellular gene expression machinery, with a focus on growth dynamics as a function of ribosomes, RNA polymerase and other “bulk” proteins inside the cell. The discrete framework permits easier testing of different combinations of biochemical interactions, leading to hypotheses elimination and model discrimination, and thus providing useful insights for the construction of a more detailed dynamical growth model.

1 Introduction

Large networks with complex interactions are hard to analyze in detail, but logical and discrete models can facilitate this task. Based essentially on the structure and topology of the network interactions, logical models provide qualitative information on the dynamical properties of system [1, 2], which can be used for model discrimination and guidance in model improvement. There are many recent examples of applications of discrete models including *Drosophila* embryo pattern formation [3, 4], yeast cell cycle [5], T-cell response [6], or an apoptosis network [7].

Boolean networks are a class of logical models whose variables are described in terms of only two levels (1 or 0; presence or absence; “on” or “off”), which have been useful for biochemical systems [8]. The dynamics of a Boolean model is determined by specifying an updating mode, most commonly synchronous (all nodes updated simultaneously) or asynchronous (only one node updated at any given instant). Since the state space is finite, the dynamics can be represented in terms of a transition graph, which can be studied using some classical algorithms from graph theory [9]. Other, more specific tools are available for an exact and rigorous analysis of the transition graph [10], computation of attractors (or asymptotic behavior) [11], and other properties [12]. In addition, a wide range of computer tools are available for simulation and analysis of discrete models [13], model reduction [14], or model checking [15].

It is clear that discrete models are not appropriate to finely describe the behavior of a system, since continuous effects are difficult to reproduce with such models (such as whether an oscillation is sustained or damped), but they are useful to verify whether a given network of interactions is feasible and compatible with known properties of the system. This is a first step towards the construction of a more detailed and informed model.

As an application, we will analyze a network of interactions involved in determining bacterial growth of *Escherichia Coli*, which varies nonlinearly with different factors, such as availability of nutrients or the concentration of the necessary enzymes and proteins needed for cell division [16, 17]. Mathematical models have been developed to describe and reproduce several regulatory modules and their response to nutrient availability [18, 19]. One of the least understood aspects in these studies remains the actual

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modeling of bacterial growth: while it is clear that growth depends on the general availability of “bulk” proteins, ribosomes, and RNA polymerase, it is difficult to find a reasonable mathematical model that reproduces all these effects [20]. In many cases, growth is considered to be a given constant and the model is designed to reproduce a single phase of bacterial growth.

Here, we propose to test and study a dynamical function for bacterial growth in terms of the major components involved in bacterial cell division, that is, gene transcription (RNA polymerase) and translation (ribosomes). To test the feasibility of mathematical growth functions, we will focus on a qualitative model of the network involved in the carbon starvation response [18] and its interconnection with a basic model describing the dynamics of ribosomes and RNA polymerase (see Section 3).

We will use two methods for analysis of qualitative systems (see Section 2): first, a method that transforms piecewise affine (PWA) systems into discrete and then Boolean models [21, 22]; and, second, a recently developed method to compute the attractors of an interconnection of two Boolean modules [11, 23]. Our analysis generates a general view of the dynamical properties of a model which is a first step towards verifying the feasibility of the model’s structure –by comparing to experimental observations– and facilitates hypotheses testing. The results indicate that at least two (positive) qualitative levels for growth rate (such as “high” and “intermediate” rates) are needed in order to reproduce both the stationary and exponential growth phases (see Section 4).

2 Methodology

In this section, we briefly recall two mathematical methods which are very useful for the analysis of qualitative systems and, in particular, interconnections of Boolean models.

2.1 From discrete to Boolean models

Although Boolean variables can only take the values 0 or 1, it is nevertheless possible to construct Boolean models that describe variables with a discrete number of values [24, 21]. Consider a discrete model $\Sigma_{disc} = (\Omega_d, F_d)$, with variables $V = (V_1, \dots, V_n)'$, state space $\Omega_d = \prod_{i=1}^n \{0, 1, 2, \dots, d_i\}$, where $d_i \in \mathbb{N}$ is the number of levels of variable V_i ($i = 1, \dots, n$), and a state transition table $F_d : \Omega_d \rightarrow \Omega_d$. The state of the system at the next instant $k + 1$ is given as a function of the state of the system at the current instant k , according to the rules F_d , using the notation:

$$V^+ = \tilde{F}_d(V).$$

Throughout this paper, the function $\tilde{F}_d$ is obtained from F_d by assuming an *asynchronous dynamic updating rule*, that is, exactly one variable is updated at any given time:

$$V^+ \in \{W \in \Omega_d : \exists k \text{ s.t. } W_k = (F_d)_k(V) \neq V_k \text{ and } W_j = V_j, \forall j \neq k\}. \quad (1)$$

Furthermore, for a more realistic model, we consider that each variable V_i can only switch from its current level to an immediately adjacent level [12], that is:

$$V_i^+ \in \{V_i - 1, V_i, V_i + 1\}, \quad \forall i. \quad (2)$$

The idea is to create an extended Boolean model $\Sigma_{bool} = (\Omega_b, F_b)$ where each discrete variable V_i is represented by d_i Boolean variables, for instance, $\{X_{i,1}, \dots, X_{i,d_i}\}$, so that the state space of the model is $\Omega_b = \{0, 1\}^{d_1 + \dots + d_n}$. There are several possible ways to convert the discrete into the Boolean variables, but here we chose to use the same criterion as in [21] which stipulates that

$$V_i = k \Leftrightarrow (X_{i,1} = \dots = X_{i,k} = 1, X_{i,k+1} = \dots = X_{i,d_i} = 0), \quad (3)$$

meaning that a variable i is at a state k if and only if all the first k Boolean variables are ON. In particular, note that this criterion implies the partition of the state space of the extended Boolean model into *permissible and forbidden regions*:

$$\Omega_p = \{X \in \Omega_b : k < l \Leftrightarrow X_{i,k} \geq X_{i,l}\}, \quad \Omega_f = \{0, 1\}^{d_1 + \dots + d_n} \setminus \Omega_p.$$

Thus, to generate the Boolean transition table F_b we need to guarantee that no transitions from a permissible to a forbidden state take place. The method described in [21] deals with this problem in a natural way, and guarantees that no transitions from permissible to forbidden states take place.

2.2 Dynamics of Boolean models

This section contains a brief summary of some useful objects that characterize the dynamics of a Boolean model. There are several possible ways of defining the dynamical updating rules [8] of a Boolean network $\Sigma = (\Omega, F_b)$, but here we will assume asynchronous updates, so the definitions and rules (1) stated for discrete systems also apply, with $d_i = 1$ for all i . Note that (2) is immediately satisfied for Boolean models.

The *asynchronous transition graph*, $G = (\Omega, E)$, of system Σ is a directed graph whose vertices (or nodes) are the elements of Ω , and the edges are given by E . There are thus 2^n nodes in G . Given any two elements $a, \tilde{a} \in \Omega$ the edge “ $a \rightarrow \tilde{a}$ ” is in E iff:

$$\tilde{a} \in \{w \in \Omega : \exists k \text{ s.t. } w_k = (F_b)_k(a) \neq a_k \text{ and } w_j = a_j, \forall j \neq k\}.$$

A *path* $a_1 \rightsquigarrow a_2$ in G is a sequence of edges linking a_1 to a_2 .

A *strongly connected component (SCC)* of G is a maximal subset $C \subset \Omega$, that contains a path joining any pair of its elements. In general, a SCC may have both incoming and outgoing edges. An SCC with no outgoing edges is called *terminal*.

An *attractor* $\mathcal{A}$ of G is a terminal strongly connected component, that is, once a trajectory enters $\mathcal{A}$ it cannot leave again. Therefore, the attractors can be said to characterize the asymptotic behavior of the network. An asynchronous transition graph always has at least one, but can have multiple, attractors. An attractor can be formed of a single state (we will call it a *singleton*) or of a subset of Ω .

2.3 Interconnection of Boolean models

To study the interconnection of the two systems, we will use a method based on control theory concepts recently developed by one of the authors [11, 23]. This method analyzes the asymptotic behavior of the interconnection of two systems directly from the behavior of the two subsystems, without having to construct or analyze the full interconnected system. The advantage is a much reduced computational cost, while still obtaining exact results: indeed, for large (e.g., $n \geq 15$) Boolean models, the computation of the asynchronous transition graph and its attractors is unfeasible, as it involves the analysis of a $2^n \times 2^n$ matrix. The idea is to first study each individual system for each set of inputs, obtain the corresponding attractors, and then construct a new object, the *asymptotic graph*. This new graph is much smaller than the state transition graph of the full model, but it contains all the information on its asymptotic dynamics, namely all the attractors of the full model correspond to attractors in the asymptotic graph. Some notation is next introduced.

Consider two asynchronous Boolean models, Σ_A and Σ_B , with a set of inputs (U_i) and a set of outputs (H_i):

$$\begin{aligned} \Sigma_A &= (\Omega_A, U_A, H_A, F_A) : \Omega_A = \{0, 1\}^{n_A}, \quad U_A = \{0, 1\}^{p_A}, \quad H_A = \{0, 1\}^{q_A}, \\ \Sigma_B &= (\Omega_B, U_B, H_B, F_B) : \Omega_B = \{0, 1\}^{n_B}, \quad U_B = \{0, 1\}^{p_B}, \quad H_B = \{0, 1\}^{q_B}. \end{aligned}$$

The following notation will be used: $a \in \Sigma_A$ and $b \in \Sigma_B$ denote the states of each system, $u \in U_A$ and $v \in U_B$ denote the inputs, and the output corresponding to state a will be represented by $h_A(a) \in H_A$ (resp., $h_B(b) \in H_B$ for state b). The synchronous rules are written:

$$a^+ = F_A(a; u), \quad \text{and} \quad b^+ = F_B(b; v).$$

For each fixed $u \in U_A$, there is a set of attractors of system Σ_A , and its elements will be represented by A_u^i , $i \in \mathbb{N}$. Similarly for system Σ_B , B_v^j , $j \in \mathbb{N}$.

The interconnection of these two systems is formed by letting the input of each system be the output of the other

$$v = h_A(a) \in U_B \quad u = h_B(b) \in U_A,$$

where it is assumed without loss of generality that $q_A = p_B$ and $q_B = p_A$. The new system will be represented by:

$$\Sigma = (\Omega, F_{bool}) : \Omega = \{0, 1\}^{n_A + n_B}, \quad F_{bool} : \Omega \rightarrow \Omega$$

with the Boolean rules F_{bool} given by the appropriate combination of F_A , F_B :

$$F_{bool}(a, b) = (F_A(a; h_B(b)), F_B(b; h_A(a))).$$

Note that F_A , F_B , and F_{bool} contain the synchronous table of state transitions. Here, we will consider that the dynamics is asynchronous, so that only one variable is updated at a given time. The asynchronous transition graphs of the two modules (one for each fixed input) and that of the full interconnected system will be called, respectively, $G^{A,u}$, $G^{B,v}$, and G .

Transition graphs and semi-attractors The first step of the method is to compute all the transition graphs $G^{A,u}$ and $G^{B,v}$, compute their attractors, and then divide each of these into subsets corresponding to a fixed output. These will be called *semi-attractors* of the individual system and are defined as follows:

$$\begin{aligned} A_{u\alpha}^i &= \text{the } i\text{-th semi-attractor of system } \Sigma_A, \text{ corresponding to input } u, \text{ with output } \alpha \\ B_{v\beta}^j &= \text{the } j\text{-th semi-attractor of system } \Sigma_B, \text{ corresponding to input } v, \text{ with output } \beta. \end{aligned}$$

Note that the standard attractor is the union of all corresponding “semi-attractors”:

$$A_u^i = \cup_{\alpha} A_{u\alpha}^i.$$

The asymptotic graph The second step of the method is to construct the asymptotic graph G^{as} whose nodes are the cross-products of semi-attractors:

$$A_{u\alpha}^i \times B_{v\beta}^j.$$

There is an edge between two of the nodes

$$A_{u\alpha}^i \times B_{v\beta}^j \rightarrow A_{u\alpha}^i \times B_{\alpha\beta}^j$$

if there is a path in the graph $G^{B,\alpha}$ that leads from some state in $B_{v\beta}^j$ to some state in $B_{\alpha\beta}^j$. Similarly for an edge $A_{u\alpha}^i \times B_{v\beta}^j \rightarrow A_{\beta\alpha}^i \times B_{v\beta}^j$. In order to satisfy an asynchronous updating scheme, only one set of variables is allowed to change for each edge. The computational cost can be further reduced by observing that all nodes with $u \neq \beta$ and $v \neq \alpha$ are transient (shown in [23]); hence, to compute the attractors of the asymptotic graph we only need to include the edges between nodes satisfying either $u = \beta$ or $v = \alpha$.

2.4 Attractors of an interconnection

The third step of the method is to compute all the attractors of G^{as} which contain, in fact, a representative of each of the attractors of G . This is theoretically proven in [11, 23]:

Theorem 1 [11] *If Q is an attractor of G , then there exists at least one corresponding attractor in G^{as} , $Q^{as} = Q^{as}(Q)$. Moreover, if $Q_1 \neq Q_2$ are two distinct attractors of G , then $Q^{as}(Q_1) \neq Q^{as}(Q_2)$.*

In other words, we recover *all the attractors of the interconnection, without explicitly constructing the interconnected system*. In broad terms, Theorem 1 says that any attractor of G generates an attractor in G^{as} , but the converse is not necessarily true and G^{as} may have more attractors than G .

To better illustrate Theorem 1, and show its advantages as well as limitations, a purely theoretical example is next given. For convenience, in the following examples, the attractors are labeled using the decimal representation for the Boolean inputs and outputs, that is:

$$000 \Leftrightarrow u = 1, \quad 001 \Leftrightarrow u = 2, \quad \dots, \quad 111 \Leftrightarrow u = 8, \quad \text{etc.}$$

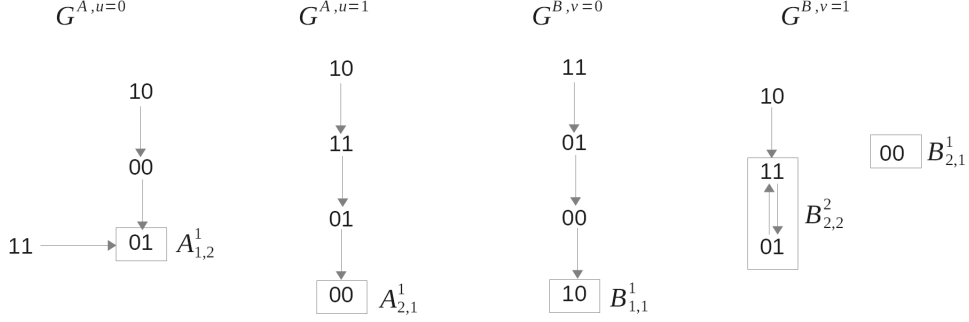


Figure 1: Example I: the asynchronous transition graphs that define the dynamics of the two systems (4).

Example I. Consider the following bi-dimensional systems A and B , with $n_A = n_B = 2$ and $p_A = p_B = 1$:

$$\begin{aligned}
 a_1^+ &= u \text{ and } (a_1 \text{ and not } a_2), \\
 a_2^+ &= [u \text{ and } (\text{not } a_1 \text{ or } a_2)] \text{ or } [\text{not } u \text{ and } a_1], \\
 h_A(a) &= a_2, \\
 b_1^+ &= [v \text{ and not } b_2] \text{ or } [\text{not } v \text{ and } (b_1 \text{ xor } b_2)], \\
 b_2^+ &= [v \text{ and } b_1 \text{ and } b_2] \text{ or } [\text{not } v \text{ and } (b_1 \text{ or } b_2)], \\
 h_B(b) &= b_2,
 \end{aligned} \tag{4}$$

whose asynchronous transition graphs $G^{A,u}$ and $G^{B,v}$ are shown in Fig. 1, for convenience. Note that the attractors in all graphs are singletons except for $B_{2,2}^2 = \{01, 11\}$. However, since the two states have the same output ($h_B(01) = h_B(11) = 1$), in this example the semi-attractors are in fact the actual attractors.

The corresponding asymptotic graph is shown in Fig. 2. To illustrate the computation of an edge, consider the product $A_{u\alpha}^i \times B_{v\beta}^j = A_{21}^1 \times B_{11}^1$: since $\alpha = 1 = v$, the system A does not induce any change in the variables b ; in contrast, the fact that $\beta = 1$ will induce a trajectory between a state in A_{21}^1 and an attractor in the graph $G^{A,2}$ (corresponds to Boolean input $u=1$). In the graph $G^{A,2}$, the state 00 is in the basin of attraction of $\{01\} = A_{12}^1$. Therefore, there is an edge $A_{21}^1 \times B_{11}^1 \rightarrow A_{12}^1 \times B_{11}^1$. All other edges are similarly computed.

Note that the full interconnected system has four variables and hence its dynamics is given by an asynchronous transition graph G with $2^4 = 16$ states. To compute the attractors of G we needed to compute a transition graph with only $2 \times 3 = 6$ states (2 attractors from system A and 3 from system B). Furthermore, as remarked above, the size of G^{as} can be further reduced by excluding the cross-product states known to be transient. In this example only $A_{21}^1 \times B_{21}^1$ satisfies the condition $u \neq \beta$ and $v \neq \alpha$, and can be excluded. For higher order systems, such a size reduction can represent very significant savings in computational cost.

The G^{as} of Example I has two attractors: $Q_1 = \{A_{12}^1 \times B_{21}^1\}$ and $Q_2 = \{A_{12}^1 \times B_{11}^1, A_{12}^1 \times B_{22}^2, A_{21}^1 \times B_{11}^1, A_{21}^1 \times B_{22}^2\}$. For this 4-dimensional example, it is easy to check that Q_1 is a true attractor of the full interconnected system (see also Prop. 1), while Q_2 is a “spurious” attractor, that is, not a real attractor of G . To see this, it suffices to note that there is a pathway that leads from a state within Q_2 to Q_1 , and which is not “covered” by G^{as} :

$$Q_2 \ni A_{21}^1 \times B_{22}^2 \ni (00, 01) \xrightarrow{G^{B,1}} (00, 00) \xrightarrow{G^{A,1}} (01, 00) \in Q_1$$

This Example shows that even very simple (and deterministic) individual asynchronous dynamics can lead to asymptotic graphs that exhibit spurious attractors. However, note that this example was specifically contrived to illustrate the generation of spurious attractors; its Boolean rules are not necessarily biologically plausible.

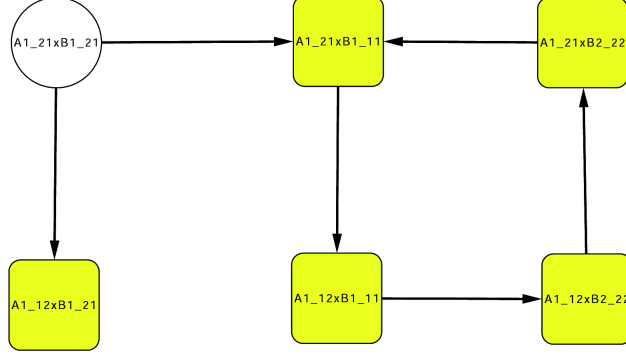


Figure 2: Example I: the asymptotic graph of the interconnection of the two systems (4). The cross-products inside shaded squares belong to an attractor. The cross-product inside a white circle represents a transient state that can be excluded from the computation.

In view of Example I, it would be useful to complement Theorem 1 by conditions permitting to decide whether an attractor of G^{as} is also an attractor of G . An exact result was also proved in [23] –i.e., recovering exactly all the attractors in G from the cross-products of semi-attractors, with no spurious generation,– by computing the *cross graph* which is similar to G^{as} but involves *cross-products of all (semi-)SCCs* (as opposed to considering only semi-attractors). However, depending on the number of SCCs, the cross graph can often be more costly to compute than the full graph G , hence the usefulness of establishing sufficient conditions for deciding whether an attractor of G^{as} is a “true” attractor.

Some preliminary results were presented in Proposition 1 of [23], which are improved below in Prop. 1. To state this, we need to introduce projection functions, for $V = A_{u\alpha}^i \times B_{v\beta}^j$, and $R = \{V_1, \dots, V_r\}$:

$$\begin{aligned}\pi(V) &= \{(a, b) \in \Omega : a \in A_{u\alpha}^i, b \in B_{v\beta}^j\}, \\ \pi(R) &= \cup_{V \in R} \pi(V), \\ \pi^A(R) &= \{a \in \Omega_A : \exists b \text{ such that } (a, b) \in \pi(R)\}.\end{aligned}$$

The *A-output* of R is the set:

$$\text{A-output} = \{h_A(a) : a \in \pi^A(R)\} \subset H_A$$

Similar definitions apply for the projection $\pi^B(R)$ and the *B-output* of R .

Recall that we are assuming $q_A = p_B$ and $q_B = p_A$, hence $H_A \equiv U_B$ and $H_B \equiv U_A$ and the *A-output* (resp., *B-output*) of R is also contained in U_B (resp., U_A). The new result of Prop. 1 is in parts (ii), (iii), which previously stated “for all $u \in U_A$ ” or “for all $v \in U_B$ ”. The new conditions are much less restrictive, although the proof is similar. If Proposition 1 is not applicable, then one may still verify *a posteriori* whether R represents an attractor of G by simulating all trajectories starting from all states in $\pi(R)$ and checking whether any of them leaves R (however, this “direct force” procedure may also involve some computational costs).

Proposition 1 *Let R be a terminal SCC of G^{as} . If either one of the following conditions is satisfied:*

- i) R is a singleton (i.e., contains a single product V);*
- ii) the A-output of R is a singleton and the set $\pi^A(R)$ is an attractor of $G^{A,u}$ for all u in the B-output of R ;*
- iii) the B-output of R is a singleton and the set $\pi^B(R)$ is an attractor of $G^{B,v}$ for all v in the A-output of R ;*

then $R^{as} = \pi(R)$ is an attractor of G .

Proof: We will use the notation $(a, b) \rightsquigarrow_G (a', b')$ to denote a path connecting the two elements in the transition graph G and $(a, b) \rightarrow_G (a', b')$ to denote a one-step transition.

Part (i) is unchanged from [23]. Parts (ii) and (iii) are very similar, so we will only prove part (iii). If the B -output of R is a singleton, say $\{\alpha\}$, then any $V \in R$ must be of the form

$$A_{\alpha(\cdot)}^j \times B_{v\alpha}^{(\cdot)}, \quad \text{for some } v \text{ in the } A\text{-output of } R.$$

In particular, (see definition of semi-SCC) all $A_{\alpha(\cdot)}^j$ belong to the same attractor A_α^j of $G^{A,\alpha}$.

Suppose now that the set $\pi^B(R)$ is an attractor for all v in the A -output of R . Then, to show that $\pi(R)$ is an attractor of G^{as} , it suffices to show that: (1) $\pi(R)$ is a strongly connected set, and (2) $\pi(R)$ contains all its successors. If (1) and (2) hold, then $\pi(R)$ is indeed a terminal SCC.

To show (1), let (a, b) and (a', b') be any two elements of $\pi(R)$. Then

$$\begin{aligned} (a, b) &\rightsquigarrow_G (a, b'), \quad \text{since } \pi^B(R) \text{ is an attractor of } G^{B, h_A(a)} (v = h_A(a) \in A\text{-output}) \\ (a, b') &\rightsquigarrow_G (a', b'), \quad \text{since } a, a' \text{ belong to the same attractor } A_\alpha^j \text{ of } G^{A,\alpha}. \end{aligned}$$

To show (2), observe that there are two forms of successors: either $(a, b) \rightarrow_G (a', b)$ or $(a, b) \rightarrow_G (a, b')$. We want to prove that both (a', b) and (a, b') are in $\pi(R)$. In the first case, since a, a' belong to the same attractor A_α^j , it is immediate to see that $(a', b) \in \pi(R)$. In the second case, since $b' \in \pi^B(R)$ and $\pi^B(R)$ is an attractor of $G^{B, h_A(a)}$, by definition of $\pi^B(R)$ there some exists a' such that $(a', b') \in \pi(R)$. Recall that the B -output is a singleton so $h_B(b') = \alpha$. This implies

$$(a, b') \rightsquigarrow_G (a', b') \rightsquigarrow_G (a, b'), \quad \text{since } a, a' \text{ belong to the same attractor } A_\alpha^j \text{ of } G^{A,\alpha}.$$

Therefore, $(a, b') \in \pi(R)$ as wanted. ■

Remark. The generalization of points (ii) and (iii) of Proposition 1 to multiple A -outputs and B -outputs is not clear, due to Example I where the spurious attractor Q_2 satisfies $A\text{-output}=B\text{-output}=\{1,2\}$. Other examples exist where an attractor of G^{as} of the same form as Q_2 is indeed and attractor of G (see Example 2 in [11]).

If Proposition 1 cannot be applied, there may be other methods to decide whether an attractor of G^{as} is a true attractor, such as identifying invariant sets of the system that contain the given attractor: examples of this are given below in Propositions 3 and 4.

Example II. To illustrate the relevance of Prop. 1, another theoretical example is now given. The two systems A and B are more conveniently represented by their asynchronous transitions graphs, one for each fixed input (Fig. 3). The dimensions are $n_a = 2$, $n_B = 3$, $p_A = 1$, $p_B = 2$ and their outputs are as follows:

$$h_A(a) = (a_1, a_2)', \quad h_B(b) = b_1.$$

Note that attractor A_2^1 splits into two semi-attractors, A_{21}^1 and A_{23}^2 , and the attractor B_2^1 splits into B_{21}^1 and B_{22}^2 . The full interconnected system has five variables and hence its dynamics is given by an asynchronous transition graph G with $2^5 = 32$ states. To compute the attractors of G we needed to compute a transition graph with $4 \times 7 - 8 = 20$ states: 4 semi-attractors from system A and 7 from system B , and 8 transient cross-products (see also Fig. 4).

The G^{as} of Example II (Fig. 4) has two attractors: $Q_1 = \{A_{11}^1 \times B_{11}^1\}$ and $Q_2 = \{A_{21}^1 \times B_{32}^1, A_{21}^1 \times B_{12}^2, A_{23}^2 \times B_{12}^1, A_{23}^2 \times B_{32}^1\}$. It is easy to check that $Q_1 = \{00000\}$ is an attractor of G , by Prop. 1(i). Likewise

$$Q_2 = \{10111, 10101, 10100, 00111, 00101, 00100\}$$

is also an attractor of G , by Prop. 1(iii): the B -output is a singleton since $\{h_B(b) : b \in \pi^B(Q_2)\} = \{1\}$; the A -output of Q_2 is $\{h_A(a) : a \in \pi^A(Q_2)\} = \{10, 00\}$; and, finally, the set $\pi^B(Q_2) = \{111, 101, 100\}$ is indeed an attractor of both $G^{B, v=10}$ and $G^{B, v=00}$.

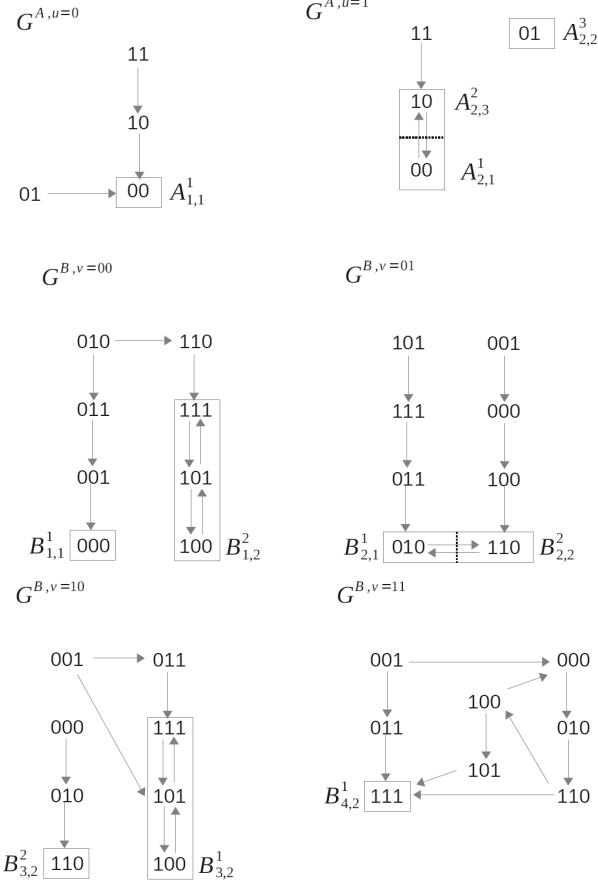


Figure 3: Example II: the asynchronous transition graphs of systems A and B , for each fixed input.

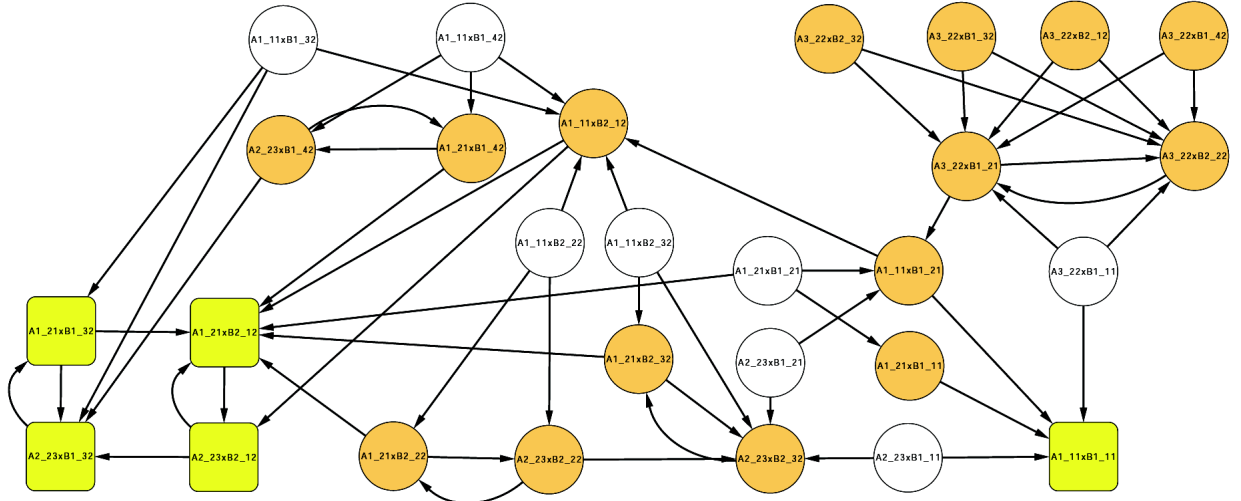


Figure 4: Example II: the asymptotic graph of the interconnection of systems A and B defined in Fig. 3. States inside light shaded squares belong to some attractor; there are two attractors in this graph. States inside white circles represent known transient state, which can be discarded from the computation. States inside light shaded circles represent all other states.

3 Application: a model for *E. Coli* growth mechanism

The bacteria *Escherichia Coli* are unicellular micro-organisms (present in the human gut, for instance) which grow and divide in the presence of a carbon source, such as glucose or other sugars. In typical experiments, in a carbon rich medium, the bacteria are observed to grow at a constant growth rate, which is referred to as the *exponential phase* [25]. In the absence of carbon, the bacteria enter a *stationary phase*, with no cellular growth or division. *E. Coli* use a network of genes and proteins to detect the presence or absence of carbon sources and respond accordingly, by adjusting their gene expression levels.

The major players in this nutritional response network are well characterized (see, for instance, [16, 19, 18] and references therein) but, in contrast, it has been difficult to find an appropriate dynamical expression for modeling the growth rate of *E. Coli* [25]. In other words, if one wishes to add a model variable to describe growth rate, what should its mathematical rule be? To overcome this problem, models often focus on either the exponential or the stationary phases, thereby considering growth rate to be either constant or zero, respectively [20]. However, such models are not able to describe the transition from one phase to the other, thus failing to provide intuition on a crucial cellular mechanism.

Growth should depend on the capacity of the bacteria to produce all the different proteins necessary to its development and cellular division. In its turn, the synthesis of any protein depends on the transcription and translation steps, which are limited, respectively, by the concentrations of RNA polymerase and ribosomes. To model the many proteins involved in bacterial growth, we will therefore distinguish between three “classes”: RNA polymerase, ribosomal proteins, and all others will be collectively denoted as “bulk” proteins (as a reference see also [17], where a distinction is made between ribosomal and nonribosomal proteins). Some models have thus tried to include these effects to obtain a more accurate expression for growth rate. For instance, one may have a dependence on one step:

$$\text{Growth rate} \sim \text{RNA polymerase} \quad (5)$$

as tested previously in [26], or in two (or more) steps, each of them separately limiting growth rate, hence the use of the *minimum* function:

$$\text{Growth rate} \sim \min\{ \text{ribosomal proteins, bulk proteins} \} \quad (6)$$

as considered in [17], or

$$\text{Growth rate} \sim \min\{ \text{ribosomal proteins, RNA polymerase} \} \quad (7)$$

as we considered in [27]. In this Section, our goal is to test these expressions, by interconnecting a well known nutritional response module with a basic transcription/translation model, using the Boolean interconnection method described in Section 2.3.

3.1 *E. Coli* nutritional stress response module

The nutritional stress response network developed in [18] involves three groups of variables, each representing a different regulatory effect: DNA supercoiling (determined by the enzymes GyrAB and TopA), carbon response (involving the proteins Crp, Cya), and a global regulator (protein Fis) that sends the carbon availability signal down to the stable RNAs (*rrn*). The latter are limiting factors in ribosome production, and are thus a measure of the growth of the bacteria.

A very brief description of the main biological steps in response to nutritional stress is as follows (see [18] and references therein): in answer to carbon depletion, the bacteria increase their cyclic AMP concentration (cAMP); this small molecule will bind to Crp (cAMP receptor protein) to form a complex that controls the expression of different genes, some involved in the synthesis of enzymes that allow the bacteria to make use of other carbon sources, others involved in morphological changes and motility. The complex cAMP-Crp also activates the enzyme Cya (adenylate cyclase), which contributes to produce cAMP from ATP, and represses the global regulator Fis, a protein which is available at high concentration during the exponential phase, and is responsible for the control of many other genes. The protein Fis also represses the complex cAMP-Crp and, among others, it controls two enzymes

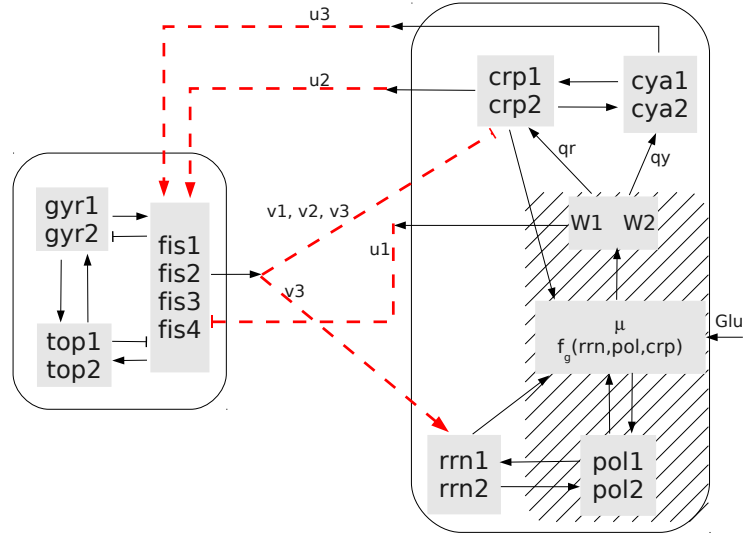


Figure 5: The interconnection of the *fis* global regulatory module (left rectangle) and a basic cellular growth model (right rectangle). Each module has three inputs and outputs: $u = (u_1, u_2, u_3)$, $v = (v_1, v_2, v_3)$. The dashed lines represent the interconnection: i.e., the output of one system becomes the input of the other. Bacterial growth rate is internally computed as a function of the external nutrient sources (Glu), ribosomes (here represented by rrn_i), RNA polymerase (pol_i) or “bulk” proteins (which will be basically represented by *crp*). Growth rate is first translated into two qualitative levels, W_1 and W_2 , which signal downstream. The region under hatching represents the new variables and interactions added to the original model in [18]. Several different forms for f_g will be tested (see text).

involved in DNA topology regulation: Gyrase AB (GyrAB) which induces negative supercoils in the DNA and Topoisomerase A which restores supercoiling to “normal” levels. Finally, Fis also stimulates the transcription of stable RNAs, a necessary condition for the production of ribosomes and hence necessary for bacterial growth.

The model developed in [18] includes a constant external input named “Signal” that represents nutritional stress, that is, the presence (“Signal” = 0) or absence (“Signal” = 1) of carbon sources, while the variable *rrn* is simply an output, as it does not influence the other variables. Growth rate was summarized into the effect of the complex cAMP-Crp on the other variables, namely Fis, Crp, and Cya. Depending on the value of “Signal”, the network reproduced two steady states corresponding to the stationary or exponential phases of *E. coli*, characterized in Table 1. The two states predicted by this model are consistent with experimental observations: in the exponential phase, Fis is present at high levels, as well as stable RNAs, and the cAMP receptor protein is not strongly present. The opposite happens in the stationary phase.

In our model, the interactions are reorganized in order to include the explicit effect of growth rate. It is known that the complex cAMP-Crp is growth dependent [28], so we replaced this complex by an equivalent expression that depends on Crp, Cya, and growth, now represented by the arrows u_1 , q_r and q_y in Fig. 5. The components inside the hatched region in Fig. 5 were not present in model [18], and the *rrn* variable did not influence the system. The objective in this paper is thus to refine the effect of growth in the system, as described below in Section 3.2.

The model [18] consists of a piecewise affine system on six variables, it was further studied in [29, 30, 31] and has been written as an extended Boolean model in [21], using the procedure briefly described in Section 2. The first Boolean module is formed by the 8 variables corresponding to genes *fis*, *gyr*, and *top*, since *fis* is described by 4 Boolean variables and *gyr*, *top* by 2 each (see Fig. 5). The rules for this module are given in the Appendix.

Since each variable may have several discrete values, the Boolean models will use var_i , $i \in \{1, 2, \dots, d\}$ to denote the corresponding d Boolean variables (see Section 2.1) (similarly for the other variables). The discrete variable can be recovered simply by adding the Boolean variables:

$$var = \sum_{i=1}^d var_i. \quad (8)$$

Table 1: The two *E. coli* modes reproduced by the model [18]. If a variable has more than one value, this means that the asymptotic solution is oscillatory among those values.

“Signal”	<i>fis</i>	<i>gyr</i>	<i>top</i>	<i>crp</i>	<i>cya</i>	<i>rrn</i>	Phase
0	1,2,3,4	1,2	0	1	2	0,1	Exponential
1	0	2	0	2	2	0	Stationary

3.2 The cellular growth module

To test the dependence of growth rate on some of the major model components, we will study a “closed-loop system”: that is, use the state of the system to construct a mathematical expression for bacterial growth rate and then feed it back to the system, by letting proteins Cya and Crp depend on it. Thus, cellular growth rate (represented by μ) now appears explicitly in the model, as an internal variable that depends dynamically on the state of the system at each instant (see Fig. 5). In agreement with the variables of the system, growth rate will have two positive discrete levels (translated to W_1 and W_2 , see equation (9) below). This also implies that the effect of growth on *fis*, *crp* and *cya* has to be updated relative to the original model [18]. In Fig. 5, there are thus three links (respectively, u_1 , q_r and q_y) which are not fixed for now, but for which several possible combinations will be tested, with a view to better understand growth signaling (see Section 4). The motivation for building this closed-loop system is to test the dynamical dependence of bacterial growth rate on the system’s variables, a question which is

still not well understood. Thus, for this example, *an expression for growth rate will be considered valid if the refined system in Fig. 5 is able to reproduce the same results as the (more schematic) model [18].*

As indicated in Fig. 5, the second Boolean module will describe the expression of the genes encoding for *crp*, *cya*, *rrn*, and will further include *pol*, to represent the expression of RNA polymerase, the enzyme responsible for gene transcription (2 Boolean variables each). The presence of carbon sources will be represented by the external input Glu. Previously [26], we have studied a mathematical expression for bacterial growth rate that is dependent only on RNA polymerase, for a simple 2-dimensional model. However, experimental data [17] suggests that ribosomes play a major role, hence we wish to improve our results by analyzing models that consider different combinations of limiting factors, and checking their compatibility with known results.

The growth variable, μ , and its downstream signals will be given by:

$$\begin{aligned}\mu &= \text{Glu} \text{ and } f_g(rrn_1, rrn_2, pol_1, pol_2, crp_1, crp_2); \\ W &= 2 - \mu; \\ W_1 &= \text{sign}(W); \\ W_2 &= \max(0, W - 1);\end{aligned}\tag{9}$$

where $\text{sign}(W) = 1$ if $W > 0$ and $\text{sign}(W) = 0$ if $W = 0$ (by construction, $\text{sign}(W)$ is never negative). The variables W_1 and W_2 correspond, respectively, to:

$$W_1 = 1 \Leftrightarrow \mu \leq 1, \quad W_2 = 1 \Leftrightarrow \mu = 0.$$

and satisfy $W_1 \geq W_2$. Following (5)-(7) and (8), different expressions for the function f_g will be tested, namely:

$$\begin{aligned}f_g^r &= rrn_1 + rrn_2, \\ f_g^p &= pol_1 + pol_2, \\ f_g^b &= crp_1 + crp_2, \\ f_g^{rp} &= \min(rrn_1 + rrn_2, pol_1 + pol_2) \\ f_g^{rb} &= \min(rrn_1 + rrn_2, crp_1 + crp_2),\end{aligned}\tag{10}$$

where the protein Crp is used as a surrogate for the level of expression of “bulk” proteins. In addition, to describe how the growth rate affects the genetic machinery, two functions need to be chosen: these correspond to the arrows labeled q_y and q_r (see below), which will also be a function of W_1 and W_2 . Several possible combinations will be tested and the final results compared to the original model.

3.3 System interconnection

The full discrete system will thus have 7 variables,

$$V = (fis, gyr, top, crp, cya, rrn, pol)',$$

with discrete levels $d_1 = 4$, $d_j = 2$ for $j = 2, \dots, 7$ and state space:

$$\Omega_d = \{0, 1, \dots, 4\} \times \{0, 1, 2\} \times \dots \times \{0, 1, 2\}.$$

The extended Boolean model will have 16 variables. As described in Section 2.3, the interconnection of two input/output asynchronous Boolean networks such as systems (13) and (14), is obtained by setting $u = h_B(b)$ and $v = h_A(a)$. Most of the input/output functions are already fixed by model [18]. There is a new interaction between the two modules, due to the effect of the growth rate in *fis*, which is represented by u_1 in Fig. 5:

$$\begin{aligned}u_1 &\in \{W_1, W_2\}, & v_1 &= fis_1, \\ u_2 &= crp_1 \text{ or } crp_2, & v_2 &= fis_2 \text{ or } fis_4, \\ u_3 &= cya_1 \text{ or } cya_2, & v_3 &= fis_3.\end{aligned}$$

The goal in this paper is the discrimination between different variants of the model in Fig. 5, in order to choose the mechanism that better represents bacterial response. The variants cover:

- models for growth rate: f_g^r , f_g^b , f_g^{rp} , and f_g^{rb} ;
- interactions between growth signals and the genetic machinery response: q_r , q_y , and u_1 .

As remarked above (Section 3.1), the interactions q_r , q_y , and u_1 in some sense replace the effect of the complex cAMP-Crp on the system, by including an explicit dependence on growth rate. To evaluate the new rules we will consider that there are two signaling stages, corresponding to the response of Cya/cAMP (the initial steps in the case of nutritional stress) and of Fis (global regulator). The response of *crp* will be timed with one or the other:

$$q_r = u_1 \quad \text{or} \quad q_r = q_y.$$

The following distinct combinations for q_y , q_r and u_1 will be tested:

$$\begin{aligned}
(I) \quad & q_y = W_1, \quad q_r = W_1, \quad u_1 = W_1 \\
(II) \quad & q_y = W_1, \quad q_r = W_1, \quad u_1 = W_2 \\
(III) \quad & q_y = W_2, \quad q_r = W_2, \quad u_1 = W_1 \\
(IV) \quad & q_y = W_1, \quad q_r = W_2, \quad u_1 = W_2 \\
(V) \quad & q_y = W_2, \quad q_r = W_2, \quad u_1 = W_2 \\
(VI) \quad & q_y = W_2, \quad q_r = W_1, \quad u_1 = W_1
\end{aligned} \tag{11}$$

4 Results

As discussed above (cf. Section 3), the goal is to recover the behavior of the system as described in Ropers et al [18] (Table 1) but now with growth rate “actually computed” by the bacteria, for the system in closed loop form which uses the state of the system. Various combinations of interactions and growth rate functions were tested, with the results summarized in Table 2 and discussed below.

Table 2: The attractors for each combination of growth rate function and interactions u_1 , q_r , q_y . Attractors σ_i satisfy $rrn = pol = 0$, while attractors α_j , $j \in \{2, 4, 24, 48, 52, 72\}$, satisfy $rrn \geq 1$ and $pol \geq 1$ (full characterizations are given in Sections 4.2 and 4.3). The indexes i, j denote the number of distinct states contained in the attractor. All the attractors of G^{as} are also attractors of G : either they satisfy Prop. 1 and/or other methods, as indicated. The highlighted row (***) represents the model variants which better reproduce Table 1 results (see Section 4.4).

Growth rate function	Interactions (11)	Attractors, G^{as} (Glu=1)		Attractors, G^{as} (Glu=0)
		Stationary	Exponential	Stationary
$f_g^r, f_g^p, f_g^{rp}, f_g^{rb}$	I	σ_4 [Prop. 1(i)]	α_4 [Prop. 1(i)]	σ_4 [Prop. 2]
	II	σ_4 [Prop. 1(i)]	α_{72} [Prop. 3]	σ_4 [Prop. 2]
	III,VI	σ_4 [Prop. 1(i)]	α_2 [Prop. 1(i)]	σ_4 [Prop. 2]
	IV,V	σ_4 [Prop. 1(i)]	α_{24} [Prop. 1(iii)]	σ_4 [Prop. 2]
***	I,III,VI	σ_{52} [Prop. 4(i)]	α_{52} [Prop. 4(ii)]	σ_4 [Prop. 2]
	II	σ_{48} [Prop. 4(i)]	α_{48} [Prop. 4(ii)]	σ_4 [Prop. 2]
	IV,V	σ_{24} [Prop. 1(iii)]	α_{24} [Prop. 1(iii)]	σ_4 [Prop. 2]

As an indication of the computational costs, application of the method presented in Section 2.3 to compute the attractors for model f_g^{rp} , case IV, gave the following results:

- there are eight constant-input asynchronous transition graphs for each system ($G^{A,u}$, or $G^{B,v}$);
- on these graphs there are a total of 22 semi-attractors for system Σ_A and 20 for Σ_B ;
- the total number of vertices in the asymptotic graph is thus $20 \times 22 = 440$;

- as remarked in Section 2.3 (and [23]), the number of vertices in G^{as} can be further reduced by eliminating those which are known to have no incoming arrow. This leads to only 90 vertices;
- the computational cost of finding the attractors of the interconnected system Σ has therefore been reduced from analysis of a size $2^{16} = 65536$ to a size 90 matrix;
- one should nevertheless consider the cost of computing this size 90 matrix, which involves reachability calculations in the 2×8 individual asynchronous transition graphs (the full process was very fast here, taking between 30-60 seconds for each model variant).

4.1 General properties

Some immediate observations from the results are:

- a common point to all model variants is that, in the presence of nutrient (Glu=1), G^{as} always has two attractors which are both attractors of G , by application of Prop. 1(i) or (iii), or other methods (see Prop. 3, 4).
- for all model variants, the first attractor ($\sigma_i, i \in \{4, 24, 48, 52\}$) has $rrn = pol = 0$ and the second attractor ($\alpha_j, j \in \{2, 4, 24, 48, 52, 72\}$) $rrn = pol = 1$. The first may be said to represent stationary phase, while the second stands for exponential phase.
- also for all model variants, in the absence of nutrient (Glu=0), there is only one attractor, σ_4 ; this can be verified directly (see Prop. 2 below). The *stationary phase* attractor σ_4 has four states and is characterized by:

$$\sigma_4 : \quad fis = 0, \quad gyr \in \{1, 2\}, \quad top = 0, \quad crp = 2, \quad cya \in \{1, 2\}, \quad rrn = 0, \quad pol = 0. \quad (12)$$

coinciding with the stationary attractor of model [18] (see Table 1) with the exception of gyr and cya which oscillate between 1 and 2 (instead of being fixed at 2).

- all models involving the ribosomes or RNA polymerase as a growth rate limiting factor exhibit the same stationary phase and a similar exponential phase attractors, depending only on the choice of feedback interactions.

As an example, for model variant f_g^{rp} , case IV, the basins of attraction for σ_4 and α_{24} are disconnected. The stationary phase attractor is formed of a single vertex, while the exponential phase attractor is composed of 9 vertices, as shown in Fig. 6. However, all system B semi-attractors coincide (Prop. 1(iii) is satisfied):

$$B_{54}^{12} = B_{74}^{17} = B_{84}^{19} = \{10111010\},$$

while the A system semi-attractors are characterized by the levels of fis , with either $fis = 1, fis = 2$ or $fis \geq 3$:

$$\begin{aligned} A_{45}^{10} &= \{10000000, 10000010, 10001000, 10001010, 10001100, 10001110\}, \\ A_{47}^{11} &= \{11000000, 11000010, 11001100, 11001010, 10001100, 11001110\}, \\ A_{48}^{12} &= \{11100000, 11100010, 11101100, 11101010, 11101100, 11101110, \\ &\quad 11110000, 11110010, 11111100, 11111010, 11111100, 11111110\}. \end{aligned}$$

In practice, the attractor in Fig. 6 can be reduced to (either) one of the horizontal rows, with three components only. All concentrations are fixed, except for fis , gyr , and top which are allowed to oscillate in any given increasing or decreasing order, provided that $fis \geq 1$ and $top \leq 1$.

In the case where no carbon sources are present, it can be shown that all model variants become the same, and hence exhibit the same stationary phase attractor. This is essentially due to the direct effect of growth rate on the synthesis of RNA polymerase.

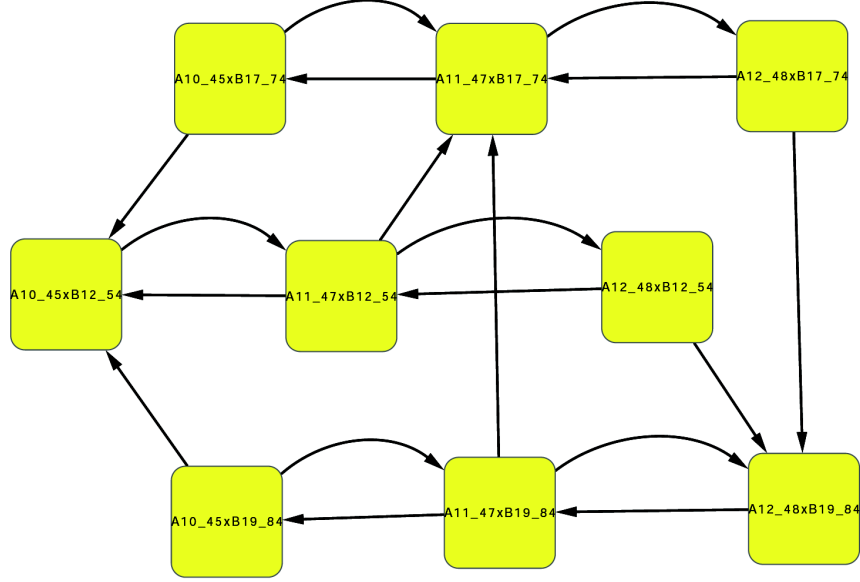


Figure 6: The exponential growth phase attractor (model f_g^{rp} , case IV). Since the $B_{\alpha 4}^j$ components are all equal, this attractor can be reduced to (either) one of the horizontal rows, with three components only (see text for more details).

Proposition 2 *Assume that $Glu=0$. Then, the asymptotic graph for all model variants exhibits only one attractor, σ_4 .*

Proof: In the case $Glu=0$, we immediately have the steady state values for rrn and pol :

$$\mu = 0 \Rightarrow pol_1 = pol_2 = 0 \Rightarrow rrn_1 = rrn_2 = 0$$

For the interactions W , q_y , q_r , and u_1 it also follows that:

$$W_1 = W_2 = 1 \Rightarrow q_y = q_r = u_1 = 1.$$

Together with the rules in Appendix, this leads to:

$$cya_1 = crp_1 = 1 \Rightarrow u_2 = u_3 = 1 \Rightarrow fis_i = hf_i$$

which simplifies to

$$fis_1^+ = fis_2, \quad fis_2^+ = fis_3, \quad fis_3^+ = fis_4, \quad fis_4^+ = fis_3 \text{ and not } fis_4.$$

Thus, at steady state, the values for fis satisfy $fis_i = 0$, for all i , which in turn imply that all the outputs of system A are zero: $v_i = 0$ for all i . The remaining concentrations can now be easily established from the Boolean rules, so it follows that there is only one attractor and that it is σ_4 (12). ■

4.2 Growth Rate limited by ribosomes or RNA polymerase

For the model variants using f_g^r , f_g^p , f_g^{rp} , or f_g^{rb} , the *stationary phase* attractor σ_4 is always the same (as described in Section 4.1). The *exponential phase* attractor, α_j , $j \in \{2, 4, 24, 72\}$, depends on the

wiring and has j states characterized by :

$$\alpha_2 : fis = 0, gyr \in \{1, 2\}, top = 0, crp = 2, cya = 2, rrn = pol = 1,$$

$$\alpha_4 : fis = 0, gyr \in \{1, 2\}, top = 0, crp = 2, cya \in \{1, 2\}, rrn = pol = 1,$$

$$\alpha_{24} : fis \in \{1, 2, 3, 4\} \text{ } gyr \in \{1, 2\}, top = 0, crp = 1, cya = 2, rrn = pol = 1,$$

$$\alpha_{72} : fis \in \{1, 2, 3, 4\}, gyr \in \{0, 1, 2\}, top = 0, crp \in \{1, 2\}, cya \in \{1, 2\}, rrn = pol = 1.$$

Note that cases IV and V (α_{24}) are similar to the exponential phase attractor of [18] (see Table 1) (the only difference is rrn now fixed at 1, which seems reasonable for the exponential phase). Cases I, III, VI (α_2, α_4) fail to reproduce the levels of fis during exponential phase (here they are fixed at zero). Case II (α_{72}) also exhibits oscillations in crp and cya , which are not observed in Table 1. This attractor does not fit into Proposition 1, but an alternative way to show that it is not a spurious attractor, is to note that the set of states with ribosomes, RNA polymerase and Fis all at discrete level 1 is invariant, so trajectories cannot leave this set; therefore, an attractor with such properties must exist, with the only possible candidate being α_{72} .

Proposition 3 Assume that $f_g \in \{f_g^r, f_g^p, f_g^{rp}, f_g^{rb}\}$. The set $Q = \{x \in \Omega_d : rrn = 1, pol = 1, fis \geq 1\}$ is invariant.

Proof: From the Boolean rules (see Appendix), it suffices to note that:

$$rrn = pol = 1 \Rightarrow \mu = 1 \Rightarrow W_1 = 1, W_2 = 0 \Rightarrow q_y = q_r = 1, u_1 = 0$$

and also

$$rrn = pol = 1, \mu \geq 1 \Rightarrow rrn = pol = 1.$$

And then:

$$u_1 = 0 \Rightarrow fis_1 = h01 \equiv 1 \Rightarrow fis \geq 1.$$

Therefore, the set Q is invariant. ■

4.3 Growth Rate limited by bulk proteins

For the model variants using f_g^b , the exponential phase attractors are characterized as follows:

$$\alpha_{24} : fis \in \{1, 2, 3, 4\} \text{ } gyr \in \{0, 1, 2\}, top = \{0, 1\}, crp = 1, cya = 2, rrn = pol = 1,$$

$$\alpha_{48} : fis \in \{1, 2, 3, 4\} \text{ } gyr \in \{0, 1, 2\}, top = \{0, 1\}, crp \in \{1, 2\}, cya = 2, rrn = pol = 1,$$

$$\alpha_{52} : fis \in \{0, 1, 2, 3, 4\}, gyr \in \{0, 1, 2\}, top = \{0, 1\}, crp \in \{1, 2\}, cya = 2, rrn = pol = 1.$$

The stationary phase attractors are similar in all variables except that $rrn = pol = 0$. Comparison with Table 1 shows many differences with respect to model [18]. These attractors are also true attractors of G , as shown by application of the following result.

Proposition 4 Define the sets P_0 and P_1 :

$$P_0 = \{x \in \Omega_d : rrn = 0, pol = 0\}, \quad P_1 = \{x : rrn = 1, pol = 1\}.$$

Then:

(i) The set P_0 is invariant independently of the function f_g ;

(ii) The set P_1 is invariant if $f_g = f_g^b$.

Proof: Invariance of P_0 follows directly from the Boolean rules for rrn_i and pol_i . For P_1 , it suffices to note that the Boolean rules imply (see Appendix): $cya \geq 1$ and $crp \geq 1$ which imply $\mu \geq 1$. Then, $rrn = pol = 1$ and $\mu \geq 1$ imply $rrn = pol = 1$. ■

4.4 Model discrimination

Based on the observations above and comparison of Tables 1 and 2, it seems clear that the growth rate function should depend on the ribosomes and/or RNA polymerase. With this model, at steady state there is no difference between a dependence on ribosomes or RNA polymerase, although the transient dynamics do depend differently on these two species (simulations in Section 4.5). (This may be due to a very simplified model for the transcription/translation steps which is, however, not our aim to study here.) The model variants corresponding to I,III,VI do not satisfy the properties of the exponential phase attractor and can thus be eliminated. The interconnection of type II has most of the correct properties, but it allows the concentration of *crp* and *cya* to oscillate, in contrast to the original model. The cases that better fit the original model are IV and V, whose asymptotic behavior is indistinguishable. This is consistent with the observations (summarized in Ropers et al. [18]) that: immediately upon carbon starvation, or absence of carbon source, transcription of the gene *cya* is activated, which leads to production of the protein Cya. Phosphorylation of Cya leads to synthesis of cyclic AMP, which in turn will bind to Crp and produce a complex [cAMP-Crp]. This complex will then control a variety of genes which are directly involved in the adaptive response of *E. coli* to a deprivation of carbon. Among others, it activates *crp*, and inactivates *cya* and the global regulator *fis*.

Furthermore, to establish the transition to exponential phase, and guarantee the presence of global regulator *fis*, the wiring interactions should be as in cases II, IV, and V, which all satisfy $u_1 = W_2$: in other words, since $W_2 = 1$ corresponds to $\mu = 0$, *fis* is inhibited only at low growth rate, as also observed in [18]. The levels of *crp* and *cya* are in agreement with those of Table 1 if *crp* is activated at high growth rate level and the inhibition effect on *cya* is not later than the activation of *crp*, i.e., $q_r = W_2$, $q_y \geq q_r$ (recall that $W_1 \geq W_2$).

In conclusion, to develop a more detailed continuous model, bacterial growth rate should depend on the ribosomes. For simplicity, one may even consider the ribosomes to be the only variable influencing growth rate (besides the external input), because no differences were observed between models with f_g^r , f_g^{rp} , or f_g^{rb} .

4.5 Dynamical behavior

By virtue of Theorem 1 and Propositions 1 to 4, we know that any trajectory of the interconnected model will eventually reach either the exponential or stationary phase attractors, depending on the initial condition and path in the graph G . To illustrate possible dynamical behaviors, we can generate trajectories in G , by randomly choosing the variable to be updated at the next instant, according to the rules. Note that this simulation does not need the graph G to be constructed; but, on the other hand, such simulations cannot characterize the full behavior of the network. Hence the usefulness of the asymptotic graph G^{as} , which can now be completed with some statistical results on initial conditions and attractors reached.

For the statistical analysis we choose interconnection model IV, and growth rate models f_g^{rp} , f_g^r , and f_g^p . It must be noted that the asymptotic graph “looses” some trajectories of the full interconnected system as, to construct G^{as} , the system is assumed to evolve in one of the constant-input/constant-output graphs $\{a\} \times G^{B,\alpha}$ or $G^{A,\beta} \times \{b\}$ until reaching an attractor. In simulations, however, the system is allowed to switch before reaching an attractor, meaning that the basins of attraction are not really disconnected as might be suggested by the asymptotic graph.

Monte Carlo simulations of the full model (10^4 randomly generated trajectories) assume that all transitions in G are equally probable and show that the two attractors are reached with similar frequencies: for f_g^{rp} , a fraction of 0.59 (0.57 for f_g^r , or 0.58 for f_g^p) trajectories converge to the exponential phase attractor.

Note that the invariance results in Propositions 3 and 4 already provide an idea of the basins of attractions, since they imply notably that initial conditions of the form $rrn = pol = 0$ (resp., $rrn = pol = 1$) lead immediately to the stationary (resp., exponential) phase attractor. To obtain more information on the distribution of the basins of attraction, we have further analyzed the probability that the system converges to either attractor given an initial condition with variable $var_i = \ell$ (where var_i runs over the sixteen Boolean variables of the system and $\ell \in \{0, 1\}$). We found that the convergence to either attractor depends essentially on the initial concentrations of RNA polymerase and ribosomes,

while all other concentrations play minor roles (in agreement with Propositions 3 and 4). An interesting observation is that, for all variables except the polymerase, and for any initial condition, the probability of converging to exponential phase is higher than to stationary phase. It is also evident that the absence of polymerase immediately prevents convergence to exponential phase. In addition, we observe that all trajectories converging to exponential phase need to start with an intermediate (or higher) level of RNA polymerase ($pol \geq 1$). Table 3 summarizes the statistics obtained from the Monte Carlo simulations. Our studies lead to the conclusion that RNA polymerase and ribosomes are both crucial for bacterial

Table 3: Initial conditions and attractor reached, for some model variants, with interconnection of type IV.

Initial conditions	Attractor reached f_g^{rp}, f_g^r, f_g^p
$pol = 0$	stationary
$pol \geq 1$ and $rrn = 0$	either
$pol \geq 1$ and $rrn \geq 1$	exponential

growth, but exert their roles at different times: initially, the presence of RNA polymerase is necessary to grow and reach the exponential phase (otherwise, if RNA polymerase is absent at time zero, the bacteria enter the stationary phase even in the presence of carbon sources), while ribosomes can be absent; at later times, the presence of ribosomes is essential to guarantee the entry into exponential phase.

5 Conclusions

Several dynamic model variants for bacterial growth rate that consider limitation by availability of the proteins needed for cell division (RNA polymerase for transcription, ribosomes for translation, or other “bulk” proteins) were tested and compared to a well established model. The main goal was to analyze (qualitative) feasibility of the wiring network, as well as the logical coherence of each model variant. This was accomplished by using a Boolean version of the model for nutritional stress response in [18], coupled with a basic cellular growth module.

We can conclude that Boolean models provide a useful framework for analysis of a system’s dynamical behavior, convenient for hypotheses testing and model discrimination. This framework presents several advantages from a computational point of view, as many tools and algorithms are available for the study and rigorous analysis of the networks. In particular, using the interconnection of two Boolean modules, it is possible to compute the attractors of a large network at a much lower cost than with classical graph theoretical tools. However, the drawbacks of this methodology include problems related to identifying the two (or more) Boolean modules as well as the corresponding inputs and outputs, which are not always obvious (see also [23]). As the number of modules and inputs increases, also the computational cost will increase and a balance must be found. This is a topic that should be further developed in future work.

A number of interesting points arise from our qualitative analysis. First, it was clear that limitation of growth rate by the ribosomes is needed in order to correctly reproduce the asymptotic modes, as well as transient dynamics, of the original model [18]. Second, in the presence of nutrient, our closed-loop model—where bacteria internally compute their growth rate, rather than responding to an already fixed signal—has the capacity for bistability (i.e., two asymptotic modes, representing exponential and stationary phases). Thus the closed-loop model also recovers the correct response to initial conditions: if both ribosomes and RNA polymerase concentration is very low, then the bacteria cannot grow even in the presence of nutrient. In the absence of nutrient, only the stationary phase attractor remains, as should be expected. Finally, by comparison to [18], we were able to discard most of the model variants and retain several properties necessary to reproduce the original model’s attractors.

Since our main goal was essentially theoretical, we have not fully explored the directions for model improvement suggested by our analysis. For instance, a more detailed module for transcription/translation

including other components besides ribosomes and RNA polymerase, or the modeling of the “bulk” proteins in a more precise way. To conclude, although discrete models are, of course, not appropriate for a detailed description of a system or to answer more specific questions, this analysis constitutes a very useful preliminary study of growth rate models. It provides many indications and clues for future work on constructing a more detailed, continuous model of the system.

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A Boolean rules of the two *E. coli* modules

The Boolean model for the Fis module is defined by a set of rules which use some auxiliary expressions of the form $h-$ given below:

$$\begin{aligned}
fis_1^+ &= (\text{not } u_1 \text{ and } h01) \text{ or } (u_1 \text{ and } h11); \\
fis_2^+ &= (\text{not } u_1 \text{ and } h02) \text{ or } (u_1 \text{ and } h12); \\
fis_3^+ &= (\text{not } u_1 \text{ and } h03) \text{ or } (u_1 \text{ and } h13); \\
fis_4^+ &= (\text{not } u_1 \text{ and } h04) \text{ or } (u_1 \text{ and } h14); \\
gyr_1^+ &= (\text{not } fis_3 \text{ and not } fis_4) \text{ or } (gyr_2 \text{ and } hf_3); \\
gyr_2^+ &= \text{not } fis_3 \text{ and not } fis_4 \text{ and } gyr_1 \text{ and } (\text{not } gyr_2 \text{ or } top_1 \text{ or } top_2); \\
top_1^+ &= (\text{not } fis_3 \text{ and not } fis_4 \text{ and } top_2) \text{ or} \\
&\quad (hf_3 \text{ and } ((\text{not } gyr_2 \text{ and } top_2) \text{ or } (gyr_2 \text{ and } (\text{not } top_1 \text{ or } top_2)))); \\
top_2^+ &= 0.
\end{aligned} \tag{13}$$

with the auxiliary expressions:

$$\begin{aligned}
hf_2 &= fis_1 \text{ and } fis_2; \\
hf_3 &= fis_1 \text{ and } fis_2 \text{ and } fis_3; \\
hf_4 &= fis_1 \text{ and } fis_2 \text{ and } fis_3 \text{ and } fis_4; \\
hf_{4n} &= fis_1 \text{ and } fis_2 \text{ and } fis_3 \text{ and not } fis_4; \\
h01 &= 1; \\
h02 &= (fis_1 \text{ and } gyr_1 \text{ and not } top_2) \text{ or } hf_3; \\
h03 &= (hf_2 \text{ and } gyr_1 \text{ and not } top_2) \text{ or } hf_4; \\
h04 &= hf_{4n} \text{ and } gyr_1 \text{ and not } top_2; \\
h11 &= ((u_2 \text{ or } u_3) \text{ and } hf_2) \text{ or } ((\text{not } u_2 \text{ or not } u_3) \text{ and } h01); \\
h12 &= ((u_2 \text{ or } u_3) \text{ and } hf_3) \text{ or } ((\text{not } u_2 \text{ or not } u_3) \text{ and } h02); \\
h13 &= ((u_2 \text{ or } u_3) \text{ and } hf_4) \text{ or } ((\text{not } u_2 \text{ or not } u_3) \text{ and } h03); \\
h14 &= (\text{not } u_2 \text{ or not } u_3) \text{ and } h04;
\end{aligned}$$

The rules for the cellular growth module can be written as follows:

$$\begin{aligned}
crp_1^+ &= 1; \\
crp_2^+ &= (\text{not } q_r \text{ and } crp_1 \text{ and not } v_1) \text{ or } (q_r \text{ and } crp_1 \text{ and not } (v_2 \text{ or } v_3)); \\
cya_1^+ &= 1; \\
cya_2^+ &= (\text{not } q_y \text{ and } cya_1) \text{ or } (q_y \text{ and } (hy1 \text{ or } hy2)); \\
rrn_1^+ &= pol_1 \text{ or } rrn_2; \\
rrn_2^+ &= pol_2 \text{ and } rrn_1 \text{ and } v_3; \\
pol_1^+ &= (\text{sign}(\mu) \text{ and } rrn_1 \text{ and } pol_1) \text{ or } pol_2; \\
pol_2^+ &= \text{sign}(\mu) \text{ and } rrn_2 \text{ and } pol_2;
\end{aligned} \tag{14}$$

where the auxiliary expressions are

$$\begin{aligned}
hy1 &= cya_1 \text{ and } (\text{not } crp_1 \text{ or not } crp_2); \\
hy2 &= cya_1 \text{ and not } cya_2 \text{ and } crp_1 \text{ and } crp_2;
\end{aligned}$$

References

- [1] L. Glass, S. Kauffman, The logical analysis of continuous, nonlinear biochemical control networks, J. Theor. Biol. 39 (1973) 103–129.
- [2] R. Thomas, Boolean formalization of genetic control circuits, J. Theor. Biol. 42 (1973) 563–585.
- [3] L. Sánchez, D. Thieffry, A logical analysis of the *drosophila* gap-gene system, J. Theor. Biol. 211 (2001) 115–141.
- [4] R. Albert, H. G. Othmer, The topology of the regulatory interactions predicts the expression pattern of the *Drosophila* segment polarity genes, J. Theor. Biol. 223 (2003) 1–18.
- [5] V. Sevim, X. Gong, J. Socolar, Reliability of transcriptional cycles and the yeast cell-cycle oscillator, PLoS Comput. Biol. 6 (2010) e1000842.
- [6] J. Saez-Rodriguez, L. Simeoni, J. A. Lindquist, R. Hemenway, U. Bommhardt, B. Arndt, U.-U. Haus, R. Weismantel, E. D. Gilles, S. Klamt, B. Schraven, A logical model provides insights into T cell receptor signaling, PLoS Comput. Biol. 3 (8) (2007) e163.
- [7] L. Calzone, L. Tournier, S. Fourquet, D. Thieffry, B. Zhivotovsky, E. Barillot, A. Zinovyev, Mathematical modelling of cell-fate decision in response to death receptor engagement, PLoS Comput. Biol. 6 (3) (2010) e1000702.
- [8] R. S. Wang, A. Saadatpour, R. Albert, Boolean modeling in systems biology: an overview of methodology and applications, Physical Biology 9 (2012) 055001.
- [9] T. Cormen, C. Leiserson, R. Rivest, C. Stein, Introduction to algorithms, MIT Press and McGraw-Hill, 2001.
- [10] T. Lorenz, H. Siebert, A. Bockmayr, Analysis and characterization of asynchronous state transition graphs using extremal states, Bull. Mathematical Biology 75(6) (2013) 920–938.
- [11] M. Chaves, L. Tournier, Predicting the asymptotic dynamics of large biological networks by interconnections of Boolean modules, in: Proc. 50th Conf. Decision and Control and European Control Conf., Orlando, Florida, USA, 2011.
- [12] R. Thomas, R. D’Ari, Biological feedback, CRC Press, 1990.

- [13] A. Gonzalez, A. Naldi, L. Sánchez, D. Thieffry, C. Chaouiya, GINsim: a software suite for the qualitative modelling, simulation and analysis of regulatory networks, *BioSystems* 84 (2) (2006) 91–100.
- [14] A. Naldi, E. Rémy, D. Thieffry, C. Chaouiya, Dynamically consistent reduction of logical regulatory graphs, *Theor. Comput. Sci.* 412 (21) (2011) 2207–18.
- [15] F. Fages, S. Soliman, N. Chabrier-Rivier, Modelling and querying interaction networks in the biochemical abstract machine BIOCHAM, *J. Biological Physics and Chemistry* 4(2) (2004) 64–73.
- [16] K. Bettenbrock, T. Sauter, K. Jahreis, J. Lengeler, E.-D. Gilles, Analysis of the correlation between growth rates, EIACrr phosphorylation, and intracellular camp levels in *Escherichia coli* K-12, *J. Bacteriol.* 189 (19) (2007) 6891–6900.
- [17] I. Shachrai, A. Zaslaver, U. Alon, E. Dekel, Cost of unneeded proteins in *E. coli* is reduced after several generations in exponential growth, *Molecular Cell* 38 (5) (2010) 758–767.
- [18] D. Ropers, H. de Jong, M. Page, D. Schneider, J. Geiselmann, Qualitative simulation of the carbon starvation response in *Escherichia coli*, *Biosystems* 84 (2) (2006) 124–152.
- [19] T. Hardiman, K. Lemuth, M. Keller, M. Reuss, M. Siemann-Herzberg, Topology of the global regulatory network of carbon limitation in *Escherichia coli*, *J. Biotechnology* 132 (2007) 359–374.
- [20] A. Goelzer, V. Fromion, Bacterial growth rate reflects a bottleneck in resource allocation, *Biochim. Biophys. Acta* 1810 (10) (2011) 978–988.
- [21] M. Chaves, L. Tournier, J. L. Gouzé, Comparing Boolean and piecewise affine differential models for genetic networks, *Acta Biotheoretica* 58(2) (2010) 217–232.
- [22] S. Jamshidi, H. Siebert, A. Bockmayr, Comparing discrete and piecewise affine differential equation models of gene regulatory networks, in: M. Lones, S. Smith, S. Teichmann, F. Naef, J. Walker, M. Trefzer (Eds.), *Information Processing in Cells and Tissues*, Vol. 7223 of LNCS, Springer, 2012, pp. 17–24.
- [23] L. Tournier, M. Chaves, Interconnection of asynchronous Boolean networks, asymptotic and transient dynamics, *Automatica* 49(4) (2013) 884–893.
- [24] P. van Ham, How to deal with variables with more than two levels, in: R. Thomas (Ed.), *Kinetic Logic: A Boolean Approach to the Analysis of Complex Regulatory Systems*, Vol. 29 of Lecture Notes in Biomathematics, Springer-Verlag, 1979, pp. 326–343.
- [25] A. Marr, Growth rate of *Escherichia coli*, *Microbiol. Rev.* 55 (1991) 316–333.
- [26] A. Carta, M. Chaves, J.-L. Gouzé, A simple model to control growth rate of synthetic *E. coli* during the exponential phase: model analysis and parameter estimation, in: D. Gilbert, M. Heiner (Eds.), *CMBS 2012, Lecture Notes in Computer Science* 7605, Springer, 2012, pp. 107–12.
- [27] A. Carta, M. Chaves, J.-L. Gouzé, A class of switched piecewise quadratic systems for coupling gene expression with growth in bacteria, in: *Proc. 9th IFAC Symp. on Nonlinear Control Systems (NOLCOS’13)*, Toulouse, France, 2013.
- [28] S. Berthoumieux, H. de Jong, G. Baptist, C. Pinel, C. Ranquet, D. Ropers, J. Geiselmann, Shared control of gene expression in bacteria by transcription factors and global physiology of the cell, *Molecular Systems Biology* 9 (2013) 634.
- [29] F. Grogard, J.-L. Gouzé, H. de Jong, Piecewise-linear models of genetic regulatory networks: theory and example, in: I. Queinnec, S. Tarbouriech, G. Garcia, S. Niculescu (Eds.), *Biology and control theory: current challenges*, *Lecture Notes in Control and Information Sciences (LNCIS)* 357, Springer-Verlag, 2007, pp. 137–159.

- [30] L. Tournier, J.-L. Gouzé, Hierarchical analysis of piecewise affine models of gene regulatory networks, *Theory Biosci.* 127 (2008) 125–134.
- [31] F. Corblin, S. Tripodi, E. Fanchon, D. Ropers, L. Trilling, A declarative constraint-based method for analyzing discrete genetic regulatory networks, *BioSystems* 98 (2) (2009) 91–104.