Trap spaces of Boolean networks are conflict-free siphons of their Petri net encoding

Van-Giang Trinh^a, Belaid Benhamou^a, Sylvain Soliman^{b,*}

^aLIS, Aix-Marseille University, Marseille, France ^bLifeware team, Inria Saclay center, Palaiseau, France

Abstract

Boolean network modeling of gene regulation but also of post-transcriptomic systems has proven over the years that it can bring powerful analyses and corresponding insight to the many cases where precise biological data is not sufficiently available to build a detailed quantitative model. Besides simulation, the analysis of such models is mostly based on attractor computation, since those correspond roughly to observable biological *phenotypes*. The recent use of trap spaces made a real breakthrough in that field allowing to consider medium-sized models that used to be out of reach. However, with the continuing increase in model size and complexity of Boolean update functions, the state-of-the-art computation of minimal trap spaces based on *prime-implicants* shows its limits due to the difficulty of the prime-implicant computation.

In this article we explore and prove for the first time a connection between trap spaces of a general Boolean network and siphons of its Petri net encoding. Besides important theoretical applications in studying properties of trap spaces, the connection enables us to propose an alternative approach to compute minimal trap spaces, and hence complex attractors, of a general Boolean network. It replaces the need for prime-implicants by a completely different technique, namely the enumeration of maximal siphons in the Petri net encoding of the original model. We then demonstrate its efficiency and compare it to the state-of-the-art methods on a large collection of real-world

^{*}Corresponding author.

Email addresses: trinh.van-giang@lis-lab.fr (Van-Giang Trinh), belaid.benhamou@lis-lab.fr (Belaid Benhamou), Sylvain.Soliman@inria.fr (Sylvain Soliman)

and randomly generated models.

Keywords:

22

23

Logical model, Boolean network, Trap space, Attractor computation, Petri net, Siphon, Systems biology

1. Introduction

From the observation that the transcriptional regulation behaved in a sigmoid step-like way, came the original idea to represent models of gene regulation as discrete event systems. Those Gene Regulation Networks (GRN) use thresholds or equivalently logical functions to represent the different regulations [1, 2, 3, 4].

Boolean net modeling has proven over the years that it can bring powerful analyses and corresponding insight to the many cases where precise biological data is not sufficiently available to build a detailed quantitative model [5], even for modelling post-transcriptional mechanisms. This is even more true for very large models where such data is frequently missing and led to a constant increase in size of logical models à la Thomas [6]. Besides simulation, the analysis of such models is mostly based on attractor computation, since those correspond roughly to observable biological phenotypes. The recent use of trap spaces [7] made a real breakthrough in that field allowing to consider medium-sized models that used to be out of reach. However, with the continuing increase in model size and complexity of Boolean update functions, the state-of-the-art computation of minimal trap spaces based on prime-implicants shows its limits. More specifically, the number of prime implicants of a Boolean function is in general exponential in the number of input nodes of this function [7]. Moreover, the computation of prime implicants is a demanding task, especially for complex Boolean functions.

It is worth noting that the recent method presented in [8] for computing minimal trap spaces avoids the prime-implicants computation by relying on the *most-permissive* semantics of Boolean networks. This method has been implemented in the tool mpbn¹ demonstrated in [9] for handling medium-sized models from the literature and very large synthetic models (up to 100,000 nodes). However, this method is only applicable for *locally-monotonic* Boolean networks, whereas the prime-implicants based method [7]

¹https://github.com/bnediction/mpbn

is applicable for *qeneral* Boolean networks (i.e., including both locally-monotonic and non-locally-monotonic ones). In addition, the bioLQM platform also provides another method using Binary Decision Diagrams (BDDs) in http:// colomoto.org/biolqm/doc/tools-trapspace.html. This method avoids the prime-implicants computation as it characterizes the set of generic trap spaces of a Boolean network by a BDD, then filters this set to get the set of all minimal trap spaces. By this approach, it requires the computation of all solutions, whereas the ASP-based methods [7, 9] can start enumerating them as they are found. Moreover, the main issue with the BDD-based method is that the number of generic trap spaces of a Boolean network may be extremely larger than the number of minimal trap spaces of this Boolean network. This issue limits the efficiency of the BDD-based method. The study [10] highlights the need for non-locally-monotonic Boolean networks in both biological and theoretical aspects. Hence, it is still necessary to develop efficient methods for computing minimal trap spaces of large-scale general Boolean networks.

Petri nets were introduced in the 60s as simple formalism for describing and analyzing information-processing systems that are characterized as being concurrent, asynchronous, non-deterministic and possibly distributed [11, 12]. The use of Petri nets for representing biochemical reaction systems, by mapping molecular species to places and reactions to transitions, hinted at already in [11, 12] was used more thoroughly quite late in [13], together with some Petri net concepts and tools for the analysis of metabolic networks. Siphons are such a concept, but they have not been used a lot for the study of biochemical systems [14, 15] even if the practical cost of computing their minimal/maximal elements appear much more manageable than the theoretical complexity would indicate [16, 17].

49

In this article we explore and prove for the first time a connection between trap spaces of a general Boolean network and siphons of its Petri net encoding. Not only having important theoretical applications in studying properties of trap spaces in Boolean networks, the connection has important practical applications in the trap space computation. Specifically, based on the connection, we propose an alternative approach to compute minimal trap spaces, and hence complex attractors, of a general Boolean network. It replaces the need for prime-implicants by a completely different technique, namely the enumeration of maximal siphons in the Petri net encoding of the original model. We then demonstrate its efficiency and compare it to the state-of-the-art methods for computing minimal trap spaces in Boolean net-

works on many real-world models from various sources in the literature and randomly generated models.

Herein we revise and extend our previous work in [18] as follows. First, more formal definitions are given and the existing proofs are made more detailed. In particular, an updated proof provides another way to prove the independence of trap spaces of a Boolean network on its update scheme, which was originally proved in [7]. Second, we showcase a theoretical application of the connection between trap spaces in Boolean networks and conflict-free siphons in Petri nets. Third, beyond the proposed ASP method implementing the alternative approach [18], we propose several other possible methods for computing minimal trap spaces using Maximum Satisfiability (MaxSAT), Constraint Logic Programming (CLP), and Integer Linear Programming (ILP). Fourth, we discuss in detail how to compute several special types of trap spaces in a Boolean network. Fifth, we present the idea for using our Petri net approach to handle the problem of inconsistent and incomplete data in modeling biological systems. Sixth, regarding the implementation, we have developed a new converter that directly reads a .bnet file and builds the Petri net encoding, instead of using the PNML conversion of bioLQM [18]. Finally, we conduct a more comprehensive benchmark on more real-world models from various sources and randomly generated models to evaluate all the proposed methods (the benchmark conducted in [18] considers only real-world models).

The rest of this paper is organized as follows: Section 2 recalls the basic concepts including Boolean networks, attractors, trap spaces, Petri nets, and siphons. Section 3 presents the main finding, the connection between trap spaces in Boolean networks and siphons in Petri nets. Section 4 presents the alternative approach for computing minimal trap spaces and the four possible methods implementing it. Section 5 presents the idea to deal with problem of inconsistent and incomplete data. Section 6 shows an important biological case study showing the applicability of the new approach. Section 7 reports the experimental results for evaluating the efficiency of the proposed methods. Finally, Section 8 concludes the paper and draws future work.

2. Preliminaries

We shall briefly recall here some preliminaries on Boolean networks related to trap spaces and Petri nets. Remove this statement because there is not sure if the encoded Boolean network preserves the trap spaces of the original multi-level logical model.

2.1. Boolean networks

Definition 2.1. A Boolean Network (BN) is a pair $\mathcal{N} = (V, F)$ where:

- $V = \{v_1, \ldots, v_n\}$ is the set of nodes. We use v_i to denote both the node v_i and its associated Boolean variable.
- $F = \{f_1, \ldots, f_n\}$ is the set of update functions. Each function f_i is associated with node v_i and satisfies $f_i : \mathbb{B}^{|IN(v_i)|} \mapsto \mathbb{B}$ where $\mathbb{B} = \{0, 1\}$ and $IN(v_i)$ denotes the set of input nodes of v_i . Note that a node $v_i \in V$ is called a source node if and only if $f_i = v_i$.

A Boolean function is *locally-monotonic* if it can be represented by a formula in disjunctive normal form in which all occurrences of any given literal are either negated or non-negated [9]. A Boolean network is said to be locally-monotonic if all its Boolean functions are locally-monotonic. Otherwise, this model is said to be non-locally-monotonic.

A state $v \in \mathbb{B}^n$ is as a mapping $v: V \mapsto \mathbb{B}$ that assigns either 0 (inactive) or 1 (active) to each node. We denote the set of all possible states of a Boolean network \mathcal{N} by $\mathcal{S}_{\mathcal{N}} = \mathbb{B}^n$. At each time step t, node v_i can update its state by

$$v_i(t+1) = f_i(v(t))$$

where v(t) is the state of \mathcal{N} at time t and $v_i(t+1)$ is the state of node v_i at time t+1. Note that for simplicity, we write $f_i(v(t))$ even $IN(v_i) \subset V$ (i.e., $IN(v_i)$ does not contain some nodes of V). An update scheme of a Boolean network specifies the way that the nodes update their states through time evolution [4]. Following the update scheme, the Boolean network transits from a state to another state (possibly identical). This transition is called the state transition and denoted by $\to \subseteq \mathcal{S}_{\mathcal{N}} \times \mathcal{S}_{\mathcal{N}}$. Then the dynamics of \mathcal{N} is captured by the directed graph $(\mathcal{S}_{\mathcal{N}}, \to)$ called the State Transition Graph (STG). There are two main types of update schemes [4]: synchronous, where all the nodes are update simultaneously, and fully asynchronous, where only one node is nondeterministically selected to be updated.

2.2. Traps spaces

We recall here some definitions from [7] for the introduction of *trap spaces*. Minimal trap spaces prove to be a very good approximation of the attractors of a Boolean network under asynchronous update schemes and have become the *de facto* standard way to analyze models of a few tens of *genes* [19, 20].

An non-empty set $T \subseteq \mathcal{S}_{\mathcal{N}}$ is a trap set with respect to \to if for every $x \in T$ and $y \in S$ with $x \to y$ it holds that $y \in T$ [7]. An attractor of \mathcal{N} with respect to \to can be defined as an inclusion-wise minimal trap set of $(\mathcal{S}_{\mathcal{N}}, \to)$. An attractor can be also seen as a terminal strongly connected component of $(\mathcal{S}_{\mathcal{N}}, \to)$ [21]. An attractor of size 1 is called a fixed point, otherwise a cyclic attractor [7].

A subspace m of a Boolean network $\mathcal{N} = (V, F)$ is a mapping $m: V \mapsto \mathbb{B} \cup \{\star\}$. $m(v_i) \in \mathbb{B}$ means that the value of v_i is fixed in m and v_i is called a fixed variable. $m(v_i) \in \star$ means that the value of v_i is free in m and v_i is called a free variable. We denote D_m the set of all fixed variables of m. A subspace m is equivalent to a set of states:

$$\mathcal{S}_{\mathcal{N}}[m] := \{ s \in \mathcal{S}_{\mathcal{N}} \mid \forall v \in D_m : s(v) = m(v) \}.$$

For example, $m = \star \star 1$ (for simplicity, we write subspaces likes states) means that $D_m = \{v_3\}, m(v_3) = 1$, and it is equivalent to the set of states $\{001, 011, 101, 111\}$. We denote $\mathcal{S}_{\mathcal{N}}^{\star} = (\mathbb{B} \cup \{\star\})^n$ the set of all possible subspaces of \mathcal{N} . Note that $|\mathcal{S}_{\mathcal{N}}^{\star}| = 3^n$ and $S_{\mathcal{N}} \subset \mathcal{S}_{\mathcal{N}}^{\star}$ [7].

A trap space is defined as a subspace that is also a trap set. It is noted that trap spaces of a Boolean network are independent of the update scheme of this model [7]. Then, we define a partial order < on $\mathcal{S}_{\mathcal{N}}^{\star}$ as: m < m' if and only if $\mathcal{S}_{\mathcal{N}}[m] \subseteq \mathcal{S}_{\mathcal{N}}[m']$ and $\mathcal{S}_{\mathcal{N}}[m] \neq \mathcal{S}_{\mathcal{N}}[m']$. Consequently, a trap space m is minimal if and only if there is no trap space $m' \in \mathcal{S}_{\mathcal{N}}^{\star}$ such that m' < m.

For example, let us consider the Boolean network shown in Example 2.1. Figure 1(a) shows the dynamics of this model under the fully asynchronous update (i.e., only one node is nondeterministically selected in order to be updated at each time step). The model has all two trap spaces, $m_1 = 11$ and $m_2 = \star\star$. Since $m_1 < m_2$, m_1 is a minimal trap space of the Boolean network.

Example 2.1. We give a Boolean network $\mathcal{N} = (V, F)$, where $V = (x_1, x_2)$ and $F = (f_1, f_2)$ with $f_1 = (x_1 \land x_2) \lor (\neg x_1 \land \neg x_2)$, $f_2 = (x_1 \land x_2) \lor (\neg x_1 \land \neg x_2)$. Herein, \land , \lor , and \neg denote the conjunction, disjunction, and negation logical operators, respectively.



(a) State transition graph, under the fully asynchronous update.

171

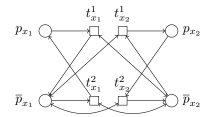
173

176

187

180

191



(b) Petri net encoding of the model. Circles denote places, whereas rectangles denote transitions.

Figure 1: Dynamics and encoding of the Boolean network of Example 2.1

2.3. Petri net encoding of Boolean networks

Definition 2.2. A Petri net is a weighted bipartite directed graph (P, T, W), where P is a non-empty finite set of vertices called places, T is a non-empty finite set of vertices called transitions, $P \cap T = \emptyset$, and $W : (P \times T) \cup (T \times P) \mapsto \mathbb{N}$ is a weight function attached to the arcs.

A marking for a Petri net is a mapping $m: P \mapsto \mathbb{N}$ that assigns a number of tokens to each place. A place p is marked by a marking m if and only if m(p) > 0. Marking m can be seen as a subset of P that contains all marked places by m. We shall write pred(x) (resp. succ(x)) to represent the set of vertices that have a (non-zero weighted) arc leading to (resp. coming from) x. In this work, we consider a class of Petri nets called 1-safe Petri nets where every place has at most 1 token and all arcs are of weight 1. In this case, weights are implicitly omitted in the arcs of a Petri net. Then, a transition $t \in T$ is enabled at a marking m if and only if $pred(t) \subseteq m$. A marking m is called a deadlock if there are no enabled transitions at m. The firing of t leads to a new marking m' specified by $m' = (m \backslash pred(t)) \cup succ(t)$. Note that when multiple transitions are enabled, we need to embed one firing scheme (similar to the update scheme of a Boolean network) to the Petri net. The classical firing scheme is that only one of the enabled transition is non-deterministically chosen to fire [12].

The link between Boolean networks \hat{a} la Thomas and Petri nets was originally established in [22] in order to make available formal methods like model-checking for the analysis of such systems. The basic encoding into 1-safe (i.e., never more than one token in each place) nets only holds for purely Boolean networks but was later extended to multivalued logical models in two ways, either in [23] with non 1-safe Petri nets or more recently in [21] with 1-safe nets but many more places.

Since our study is focused on Boolean networks, we briefly recall the original encoding here. Its basis is that every node (gene) v of the original model $\mathcal{N} = (V, F)$ is represented by two separate places $(p_v \text{ and } \overline{p}_v)$, corresponding to its two states, active, and inactive, respectively. Each conjunct of the logical function that activates the gene will lead to a transition t, consuming the inactive place (i.e., a directional arc from \overline{p}_v to t), producing the active place (i.e., a directional arc from t to p_v), and with all other literals both consumed and produced (i.e., a bidirectional arc). And conversely for the inactivation. Let s be a state of the Boolean network and m_s be its corresponding marking in the encoded Petri net. It holds that $\forall v \in V, s(v) = 0$ if and only if $m_s(\overline{p}_v) = 1$ and s(v) = 1 if and only if $m_s(p_v) = 1$. Note also that at any marking m of the Petri net encoding a Boolean network, it always holds that $m(p_v) + m(\overline{p}_v) = 1$.

The main property of this encoding is that it is completely faithful with respect to the update scheme of the original Boolean network. For each node v of \mathcal{N} , only transitions corresponding to v can change the current marking of p_v or \bar{p}_v . In addition, at any marking at most one of such transitions is enabled because $m(p_v) + m(\bar{p}_v) = 1$ holds. Hence, for any update scheme in \mathcal{N} , we have a corresponding firing scheme in \mathcal{P} , which preserves the equivalence between the dynamics of \mathcal{N} and \mathcal{P} [24].

For illustration, let us reconsider the Boolean network shown in Example 2.1. Figure 1(b) shows the Petri net encoding of this Boolean network. Place p_{x_1} (resp. \bar{p}_{x_1}) in \mathcal{P} represents the activation (resp. the inactivation) of node x_1 in \mathcal{N} . Marking $\{p_{x_1}, \bar{p}_{x_2}\}$ in \mathcal{P} represents state 10 in \mathcal{N} . Transitions $t_{x_1}^1$ and $t_{x_1}^2$ represent the update of node x_1 . Of course, in any marking $t_{x_1}^1$ and $t_{x_1}^2$ cannot be both enabled. Then, the fully asynchronous update scheme in \mathcal{N} corresponds to the classical firing scheme in \mathcal{P} where only one of the enabled transitions for a given marking will be fired [12].

Note that given a Boolean network in the standard SBML-Qual format [25], i.e., the package of SBML v3 [26] for such models, one can easily obtain its Petri net encoding in the Petri Net Markup Language (PNML)² standard using the bioLQM³ library. This piece of software extracted from GINsim [27] and part of the ColoMoTo⁴ [28] software suite allows for easy conversion

²https://www.pnml.org/

³http://www.colomoto.org/biolqm/

⁴http://colomoto.org/

between standard formats. It also accepts many other common formats for Boolean networks, notably the .bnet files of the BoolNet [29, 19] tools. The conversion is executed as follows:

java -jar GINsim.jar -lqm <input.{sbml,bnet,zginml,...}> <output.pnml> Note that transforming a Boolean network defined by its functions into its Petri net encoding roughly relies on obtaining conditions for the activation and inactivation of the states. In [22] this took the form of the whole truth table of the Boolean functions, but as shown in Appendix 1 of [21] computing Disjunctive Normal Forms (DNF) of each Boolean function is enough. Though this might appear quite computationally intensive it is important to remark first that contrary to the prime-implicants case, there is no need to find minimal DNFs. One way to look at this is to consider that this amounts to a similar approach as that used in [8] but with the encoding of both activation and inhibition functions as DNFs in order to take into account possible non-local-monotonicity. This does not change the worst-case-complexity (obtaining a single DNF being exponential) but might matter a lot in practice. As such, we will explore how this transformation, here using BDDs in bioLQM, and the one based on the most-permissive semantics compare in the Section 7 on evaluation.

2.4. Siphons

231

232

233

235

236

237

238

239

241

243

245

247

248

250

252

254

258

250

Siphons are a static and classical property of Petri nets [11]. Note however that the use of siphons for the analysis of biological models, though it is not new, has been mostly relevant to the ODE-based continuous semantics of Chemical Reaction Networks [30, 31, 32]. We recall here the basic definition establishing that to produce something in a siphon you must consume something from the siphon. This corresponds to the idea that a siphon is a set of places that once unmarked remains unmarked.

Definition 2.3. A siphon of a Petri net (P, T, W) is a set of places S such that:

$$\forall t \in T, S \cap succ(t) \neq \emptyset \Rightarrow S \cap pred(t) \neq \emptyset.$$

Note that \emptyset is trivially a siphon.

Let $pred(S) := \bigcup_{s \in S} pred(s)$ and $succ(S) := \bigcup_{s \in S} succ(s)$. If $S = \emptyset$, then conventionally $pred(S) = succ(S) = \emptyset$. We have an important property on siphons [33] as follows.

Proposition 2.1. Let S be a siphon of a Petri net (P, T, W). Then $pred(S) \subseteq succ(S)$.

3. Minimal trap spaces as maximal conflict-free siphons

First, we add a definition related to any set of places of a Petri net encoding a Boolean network, and notably a siphon of such a net.

Definition 3.1. A set of places of Petri net \mathcal{P} encoding Boolean network \mathcal{N} is conflict-free if it does not contain any two places corresponding to the active and inactive states of the same node of \mathcal{N} . Then, a conflict-free siphon S is said to be maximal if and only if there is no other conflict-free siphon S' such that $S \subset S'$.

Intuitively, a siphon is a set of places that once unmarked remains so. 271 If it is conflict-free then its dual corresponds to a partial-state of the model such that whatever update, the fixed values remain so (since the unmarked places remain unmarked). This is precisely the definition of a trap space and 274 maximality of the siphon is equivalent to as many fixed values as possible, hence minimality of the trap space. For example, the Boolean network given 276 in Example 2.1 has two trap spaces, $m_1 = 11$ and $m_2 = \star\star$. The Petri net encoding of this Boolean network has five generic siphons, $S_1 = \emptyset$, $S_2 = \emptyset$ 278 $\{p_{x_1}, \overline{p}_{x_1}\}, S_3 = \{p_{x_2}, \overline{p}_{x_2}\}, S_4 = \{\overline{p}_{x_1}, \overline{p}_{x_2}\}, \text{ and } S_5 = \{p_{x_1}, \overline{p}_{x_1}, p_{x_2}, \overline{p}_{x_2}\}.$ However, only S_1 and S_4 are conflict-free siphons and correspond to m_2 and m_1 , respectively. Since $S_1 \subset S_4$, S_4 is a maximal siphon corresponding to the minimal trap space m_1 . Hereafter, we formally prove that a (maximal) 282 conflict-free siphon is equivalent to a (minimal) trap space.

Definition 3.2. Let m be a subspace of Boolean network $\mathcal{N}=(V,F)$. A mirror of m is a set of places S in the Petri net encoding \mathcal{P} of \mathcal{N} such that:

$$\forall v \in D_m, m(v) = 0 \Leftrightarrow p_v \in S, m(v) = 1 \Leftrightarrow \overline{p}_v \in S$$

and

$$\forall v \in V \setminus D_m, p_v \notin S, \overline{p}_v \notin S.$$

Theorem 3.1. Let $\mathcal{N} = (V, F)$ be a Boolean network and \mathcal{P} be its Petri net encoding. A subspace m is a trap space of \mathcal{N} if and only if its mirror S is a conflict-free siphon of \mathcal{P} .

Proof. First, we show that if m is a trap space of \mathcal{N} , then S is a conflict-free siphon of \mathcal{P} (*). If $D_m = \emptyset$, then $S = \emptyset$ is trivially a conflict-free siphon of \mathcal{P} . Thus, we consider the case that $D_m \neq \emptyset$ (resp. $S \neq \emptyset$). Assume that S is

not a siphon of \mathcal{P} . Then, there is a transition $t \in T$ such that $S \cap succ(t) \neq \emptyset$ but $S \cap pred(t) = \emptyset$. This implies that there is a place $p \in S$ such that $p \in succ(t)$ but $p \notin pred(t)$. Let v be the corresponding node in \mathcal{N} of p. By 295 the characteristics of the encoding [22], there is a directional arc from t to pand a directional arc from the complementary place of p to t. Without loss 297 of generality, we assume that $p = p_v$, then there is a directional arc from t 298 to p_v and a directional arc from \overline{p}_v to t. We follow the following procedure to find a state $s \in \mathcal{S}_{\mathcal{N}}[m]$ such that $m_s(p') = 1, \forall p' \in pred(t)$ where m_s is 300 the corresponding marking in \mathcal{P} of s. For every place $p' \in pred(t)$, let p'' be 301 the complementary place of p' and v' be the corresponding node in \mathcal{N} of p'and p''. If $p'' \notin S$, then $v' \notin D_m$ and we can always set a Boolean value to 303 s(v') such that $s \in \mathcal{S}_{\mathcal{N}}[m]$ and $m_s(p') = 1$. If $p'' \in S$, then $v' \in D_m$ and we set s(v') = m(v'). In this case, if $p' = p_{v'}$ then s(v') = m(v') = 1 leading to $m_s(p')=1$, if $p'=\overline{p}_{v'}$ then s(v')=m(v')=0 leading to $m_s(p')=1$. For the remaining nodes of \mathcal{N} , we can always set Boolean values to these nodes to preserve that $s \in \mathcal{S}_{\mathcal{N}}[m]$. We also have $m_s(p_v) = 0$ by the characteristics of the encoding [22]. Now, t is enabled at marking m_s . Its firing leads to a new marking m'_s such that $m'_s(p_v) = 1$ and $m'_s(\overline{p}_v) = 0$. Let s' be the 310 corresponding state in \mathcal{N} of m_s' . We have s'(v)=1 because $m_s'(p_v)=1$ and m(v) = 0 because $p_v \in S$. This implies that $s' \notin \mathcal{S}_{\mathcal{N}}[m]$. For any firing 312 scheme of \mathcal{P} , the firing of t always happens. Since a firing scheme of \mathcal{P} is equivalent to an update scheme of \mathcal{N} , s can escape from the trap space m for any update scheme of \mathcal{N} , which contradicts to the property of a trap space. Hence, S is a siphon of \mathcal{P} . By the definition of a mirror, S is also a 316 conflict-free one. 317

Second, we show that if S is a conflict-free siphon of \mathcal{P} , then m is a trap space of \mathcal{N} (**). By the definition of a mirror, m is a subspace of \mathcal{N} . Let s be an arbitrary state in $\mathcal{S}_{\mathcal{N}}[m]$ and m_s be its corresponding marking in \mathcal{P} . Assume that there is a place $p \in S$ such that $m_s(p) = 1$. Let v be the corresponding node in \mathcal{N} of p. Since $p \in S$, $v \in D_m$ and m(v) = s(v). If $p = p_v$, then $m_s(p_v) = 1$ leading to m(v) = s(v) = 1 by the characteristics of the encoding [22]. By the definition of a mirror, m(v) = 0 because $p_v \in S$, which is a contradiction. It is symmetric for the case that $p = \overline{p}_v$. Hence, $m_s(p) = 0, \forall p \in S$. In any marking m'_s reachable from m_s regardless of the firing scheme of \mathcal{P} , we have $m'_s(p) = 0, \forall p \in S$ by the dynamical property on markings of a siphon [33]. Let s' be the corresponding state in \mathcal{N} of m'_s . For every node $v \in D_m$, we have all two cases as follows. Case 1: $p_v \in S$, then $m'_s(p_v) = 0$, thus s'(v) = 0 = m(v). Case 2: $\overline{p}_v \in S$, then $m'_s(\overline{p}_v) = 0$, thus

s'(v) = 1 = m(v). Hence, s'(v) = m(v) for every $v \in D_m$. Then, $s' \in \mathcal{S}_{\mathcal{N}}[m]$.

By the definition of a trap space and the arbitrariness of s, m is a trap space of \mathcal{N} .

334

335

336

337

339

341

352

355

361

363

364

From (*) and (**), we can conclude the proof.

From the proof of Theorem 3.1, we can see that this theorem still holds for any update scheme of the Boolean network. Since the Petri net encoding of a Boolean network is independent of its update scheme and siphons are a static property of a Petri net, we can imply that trap spaces of a Boolean network are independent of its update scheme. Note that the original proof for this property of trap spaces (see Theorem 1 of [7]) only considers the two popular update schemes (i.e., synchronous and fully asynchronous). This exhibits the very first theoretical application of the connection between trap spaces of Boolean networks and siphons of Petri nets.

Theorem 3.2. Let \mathcal{N} be a Boolean network and \mathcal{P} be its Petri net encoding. A subspace m is a minimal trap space of \mathcal{N} if and only if its mirror S is a maximal conflict-free siphon of \mathcal{P} .

Proof. First, we show that if m is a minimal trap space of \mathcal{N} , then S is a maximal conflict-free siphon of \mathcal{P} (*). Since m is a trap space of \mathcal{N} , S is a conflict-free siphon of \mathcal{P} by Theorem 3.1. Assume that S is not maximal. Then, there is another conflict-free siphon S' such that $S \subset S'$. By Theorem 3.1, there is a trap space m' corresponding to S'. Following the definition of a mirror, $D_m \subset D_{m'}$ and $m(v) = m'(v), \forall v \in D_m$. It follows that $S_{\mathcal{N}}[m'] \subset S_{\mathcal{N}}[m]$, thus m' < m. This contradicts to the minimality of m. Hence, S is a maximal conflict-free siphon of \mathcal{P} .

Second, we show that if S is a maximal conflict-free siphon of \mathcal{P} , then m is a minimal trap space of \mathcal{N} (**). Since S is a conflict-free siphon of \mathcal{P} , m is a trap space of \mathcal{N} by Theorem 3.1. Assume that m is not minimal. Then, there is another trap space m' such that m' < m. By the definition of the partial order < on subspaces, $\mathcal{S}_{\mathcal{N}}[m'] \subset \mathcal{S}_{\mathcal{N}}[m]$. Let S' be the mirror of m'. S' is a conflict-free siphon by Theorem 3.1. Following the definition of a mirror, $S \subset S'$, which contradicts to the maximality of S. Hence, m is a minimal trap space of \mathcal{N} .

From (*) and (**), we can conclude the proof.

We here showcase a theoretical application of the connection between trap spaces in Boolean networks and conflict-free siphons in Petri nets. We use it to prove a property of minimal trap spaces, which has surprisingly not been formally proved. Specifically, all minimal trap spaces of a Boolean network are mutually disjoint. This property is important because we can use it to approximate the set of attractors of the Boolean network [7].

Theorem 3.3. Let $\mathcal{N} = (V, F)$ be a Boolean network. For any two distinct minimal trap spaces m_1 and m_2 of \mathcal{N} , we have that $\mathcal{S}_{\mathcal{N}}[m_1] \cap \mathcal{S}_{\mathcal{N}}[m_2] = \emptyset$.

Proof. Let \mathcal{P} be the Petri net encoding of \mathcal{N} . If \mathcal{N} has only one minimal trap space, then the theorem trivially holds. Note that by Theorem 3.2, \mathcal{N} always has at least one minimal trap space because \mathcal{P} has at least one maximal conflict-free siphon. Hence, we consider the case that \mathcal{N} has at least two minimal trap spaces.

375

376

382

384

388

389

390

394

396

Consider two any distinct minimal trap spaces m_1 and m_2 . Assume that $\mathcal{S}_{\mathcal{N}}[m_1] \cap \mathcal{S}_{\mathcal{N}}[m_2] \neq \emptyset$. Let S_1 and S_2 be the mirrors of m_1 and m_2 , respectively. By Theorem 3.2, S_1 and S_2 are maximal conflict-free siphons of \mathcal{P} . We have that $S = S_1 \cup S_2$ is also a siphon because of Proposition 2.1. For every node $v \in V$, assume that $p_v \in S$ and $\overline{p}_v \in S$ hold. Since S_1 and S_2 are conflict-free, there are all two cases. Case 1: $p_v \in S_1$ and $\overline{p}_v \in S_2$. Case 2: $p_v \in S_2$ and $\overline{p}_v \in S_1$. These two cases lead to $m_1(v) \neq m_2(v), m_1(v) \neq \star, m_2(v) \neq \star$, then $\mathcal{S}_{\mathcal{N}}[m_1] \cap \mathcal{S}_{\mathcal{N}}[m_2] = \emptyset$. This is a contradiction. Hence, for every node $v \in V$, $p_v \in S$ and $\overline{p}_v \in S$ cannot hold together. Therefore, S is conflict-free. Now, we have that S is a conflict-free siphon but $S_1 \subset S$ or $S_2 \subset S$ holds because $S_1 \neq S_2$. This contradicts to the maximality of S_1 and S_2 . Hence, $\mathcal{S}_{\mathcal{N}}[m_1] \cap \mathcal{S}_{\mathcal{N}}[m_2] = \emptyset$ holds.

One naturally computational application of Theorem 3.1 is that we can efficiently decide whether a subspace m is a trap space. In PyBoolNet [19], this is checked by using the percolation on the prime-implicants of the Boolean functions. As we have mentioned at the beginning of this article, the computation of prime-implicants is a demanding task for complex Boolean networks, even is sometimes intractable. Hence, the checking method in [19] shows its limitations. Instead, we can first compute the mirror S_m of m in the Petri net encoding. Then, by Proposition 2.1 and Theorem 3.1, we can check if $pred(S_m) \subseteq succ(S_m)$. Note that the Petri net construction is less computationally demanding than the prime-implicant computation because it only requires computing generic (not prime) implicants of the Boolean func-

tions [21]. In addition, the time complexity of the above checking method is quadratic in the number of transitions of the Petri net in worst cases.

Furthermore, by Theorem 3.2, we can reduce the problem of computing all minimal trap spaces of a Boolean network to the problem of computing all maximal conflict-free siphons of its Petri net encoding. Note that in the case of special types of trap spaces (e.g., fixed points), this can be put in regard to special types of siphons in Petri nets. See Subsection 4.5 for more discussions about many special types of trap spaces. It might actually be possible to generalize our result to any 1-safe place-complementary Petri net to define a notion of trap spaces that might be useful for the analysis of Petri nets, but this is out of the scope of this present article.

It is noted that there are no existing methods specifically designed for computing maximal conflict-free siphons (even maximal siphons) of a Petri net. The reason might be that researchers mainly focus on minimal generic siphons [33] in the field of Petri nets. Hence, we here propose several methods for computing maximal conflict-free siphons of a Petri net. The details of the proposed methods shall be given in the next section.

4. Computation methods

4.1. Characterization

First, we show the characterization of all conflict-free siphons of the encoded Petri net $\mathcal{P}=(P,T,W)$. Suppose that S is a generic siphon of \mathcal{P} . If a place p should belong to S, then by Proposition 2.1 all the transitions in pred(p) must belong to succ(S). A transition t belongs to succ(S) if and only if there is at least one place p' in S such that $p' \in pred(t)$. Hence, for each transition $t \in pred(p)$, we can state that

$$p \in S \Rightarrow \bigvee_{p' \in pred(t)} p' \in S.$$
 (1)

The system of all the rules of the above form with respect to all pairs (p,t) where $p \in P, t \in T, t \in pred(p)$ fully characterizes all generic siphons of a Petri net and has been used with SAT solvers in [16, 17]. To make S to be a conflict-free siphon, we need to add to the system the rule

$$p_v \in S \Rightarrow \overline{p}_v \notin S \land \overline{p}_v \in S \Rightarrow p_v \notin S$$
 (2)

for each node $v \in V$. By definition, the final system fully characterizes all conflict-free siphons of the encoded Petri net.

32 4.2. Constraint satisfaction problem

The following Boolean Constraint Satisfaction Problem (CSP) directly derives from the above characterization:

Definition 4.1. Given a Petri net $\mathcal{P} = (P, T, W)$ encoding a Boolean network $\mathcal{N} = (V, F)$. The CSP $\mathcal{C}(\mathcal{P})$ is the triple (R, D, C) where

- R = P, i.e., a variable is introduced for each place of \mathcal{P} ,
- $D(p) = \mathbb{B}$ for all $p \in R$, i.e., the variables are Boolean,
- $C = \{ \neg p_v \lor \neg \overline{p}_v = 1 \mid \forall v \in V \} \bigwedge \{ (p = 1 \to \bigvee_{p' \in pred(t)} p' = 1) \mid p \in P, t \in pred(p) \}.$

Proposition 4.1. $\mathcal{C}(\mathcal{P})$ is satisfied by a valuation r if and only if

$$\{p \in P \mid r(p) = 1\}$$

is a conflict-free siphon of \mathcal{P} .

437

438

439

440

447

449

451

453

455

456

457

459

Proof. By the former part $\neg p_v \lor \neg \overline{p}_v = 1$ of C, the conflict-freeness is imposed because for any satisfable valuation r, $r(p_v) = r(\overline{p}_v) = 1$ is impossible for all $v \in V$. As shown in [17], the latter part of C can characterize the set of all generic siphons of \mathcal{P} . Hence, we can conclude the proof.

In [17], the set of all siphons of a given Petri net is characterized by a similar Boolean CSP except the conflict-freeness constraint. From the encoded CSP, the set of all *minimal* siphons of the Petri net can be enumerated in the set inclusion order. For enumerating siphons in the set inclusion order, the proposed method by [17] uses the technique that labels directly the Boolean variables with increasing value selection (i.e., to test first the absence, then the presence of a place in the candidate solution). The method has two implementations, one uses an iterated SAT procedure and the other uses Constraint Logic Programming (CLP) with backtracking.

One natural question is that how to use the CSP-based method for enumerating all the maximal conflict-free siphons of a Petri net encoding a Boolean network? Of course, the set of all conflict-free siphons of the Petri net can easily characterized by the CSP model presented in [17] along with the additional constraint $\neg p_v \vee \neg \overline{p}_v = 1$, for each $v \in V$, which represents

the conflict-freeness. However, the main concern is to enumerate all the maximal ones, which is not trivial to adapt from the CSP-based method. By Proposition 4.1, the set of all maximal conflict-free siphons of \mathcal{P} can be 464 enumerated in the (maximality) set inclusion order, by restarting the search each time a conflict-free siphon S is found, with the following additional con-466 straint for disallowing any subset of that conflict-free siphon: $\bigvee_{p \notin S} p = 1$. 467 For enumerating conflict-free siphons in the set inclusion order, we can use the same technique as used in [17] but with the opposite setting, i.e., labeling 469 directly the Boolean variables with decreasing value selection. The correct-470 ness of this technique comes from the fact that once S is found, it is the 471 conflict-free siphon of maximum cardinality among all the remaining feasible 472 conflict-free siphons. Similar to [17], the newly CSP-based method can also 473 be implemented with SAT and CLP solvers.

For the SAT-based method, however a more direct method is to use a MaxSAT solver. We construct a MaxSAT problem with the following hard clauses:

$$(\neg p_v \lor \neg \overline{p}_v), \forall v \in V$$

478 and

475

483

485

$$(\neg p \lor \bigvee_{p' \in pred(t)} p'), \forall p \in P, \forall t \in pred(p).$$

We set a soft clause for each variable of the CSP and then use a "minimal correction subset" blocking strategy, which will ensure set-inclusion maximality of the solutions. This is what is implemented in Trappist using the RC2 MaxSAT solver [34] available through the python-sat package⁵.

For the CLP-based method, we use TODO: ...

4.3. Answer set programming-based method

Another possible method is to translate the characterization shown in Subsection 4.1 into the ASP \mathcal{L} as follows. We introduce atom p-v (resp. n-v) to denote place p_v (resp. \overline{p}_v), $\forall v \in V$. The set of all atoms in \mathcal{L} is given as $\mathcal{A} = \bigcup_{v \in V} \{p-v, n-v\}$. For each pair (p,t) where $p \in P, t \in T, t \in pred(p)$, we translate the rule (1) into the ASP rule

⁵https://pysathq.github.io/docs/html/api/examples/rc2.html

where $a \in \mathcal{A}$ is the atom representing place p and $\{a_1, \ldots, a_k\} \subseteq \mathcal{A}$ is the set of atoms representing places in pred(t). The rule (2) is translated into the ASP rule

for each $v \in V$. This ASP rule guarantees that two places representing the same node in \mathcal{N} never belong to the same siphon of \mathcal{P} , representing the conflict-freeness. Naturally, a Herbrand model (see, e.g., [35]) of \mathcal{L} is equivalent to a conflict-free siphon of \mathcal{P} . To guarantee that a Herbrand model is also a stable model (an answer set), we need to add to \mathcal{L} the two choice rules

$$\{p-v\}. \{n-v\}.$$

500

501

503

507

508

510

511

512

514

516

for each $v \in V$. Note that the number of atoms of \mathcal{L} is only 2n, whereas the ASP encoding shown in [7] has as many atoms as the number of prime-implicants of the Boolean network and that number might be exponential in n. In [8], there is an ASP characterization of trap spaces that does not rely on minimal DNFs either and thus seems very similar to our ASP encoding. Remarkably it only requires the DNF for the activation part, using the information that it will only be used for locally-monotonic Boolean networks. We would therefore expect that, when available, it will have comparable performance on the ASP part (the ASP program would be approximately twice smaller, though redundancy is not always bad in that field), but can also avoid combinatorial explosion of the Petri net encoding for some formula where the activation DNF is simple but the inhibition is not. Since mpbn is included in our benchmark this will be evaluated in our experiments.

Now, a solution (simply an answer set) $A \subseteq \mathcal{A}$ of \mathcal{L} is equivalent to a conflict-free siphon S of \mathcal{P} , thus a trap space m of \mathcal{N} . The conversion from A to m is straightforward. If $\mathbf{p}-\mathbf{v} \in A$ then $v \in D_m$ and m(v) = 0. Conversely, if $\mathbf{n}-\mathbf{v} \in A$ then $v \in D_m$ and m(v) = 1. Otherwise, $v \notin D_m$. Computing multiple answer sets is built into ASP solvers and the solving collection POTASSCO [35] also features the option to find set-inclusion maximal answer sets with respect to the set of atoms. Naturally, a set-inclusion maximal answer set of \mathcal{L} is equivalent to a maximal conflict-free siphon of \mathcal{P} , thus a minimal trap space of \mathcal{N} . By using this built-in option, we can compute all the set-inclusion maximal answer sets of \mathcal{L} (resp. all the minimal trap spaces of \mathcal{N}) in one execution.

4.4. Integer linear programming-based method

524

538

530

541

We first show how an Integer Linear Programming (ILP) \mathcal{I} can define a set of all conflict-free siphons of the encoded Petri net \mathcal{P} . We introduce binary variable p-v (resp. n-v) to denote place p_v (resp. \overline{p}_v), $\forall v \in V$. The set of all binary variables in \mathcal{I} is $\bigcup_{v \in V} \{p-v, n-v\}$. For each pair (p,t) where $p \in P, t \in T, t \in pred(p)$, we translate the rule (1) into the ILP inequality

$$a \le a_1 + ... + a_k$$

where a is the binary variable representing place p and $\{a_1, ..., a_k\}$ is the set of binary variable representing places in pred(t). The rule (2) is translated into the ILP inequality

$$p-v + n-v \le 1$$

for each $v \in V$. This inequality forbids both p-v and n-p receive the value 1, thus representing the conflict-freeness. Since we only consider feasible solutions, the objective function is set to max p-v for some $v \in V$. Naturally, a solution I of \mathcal{I} is equivalent to a conflict-free siphon S of \mathcal{P} . The conversion is that

$$S = \{ p \in P \mid I(\mathtt{a-p}) = 1 \}$$

where a-p is the binary variable presenting place p.

We can see the similarity between \mathcal{I} and the encoded ASP shown in the previous subsection. However, due to the nature of solutions of an ILP, it is hard to compute all the set-inclusion maximal solutions of \mathcal{I} in one execution of an ILP solver. Hence, we propose an iterative approach as follows.

The conflict-free siphon of maximum cardinality is of course maximal. Therefore, we impose the following objective function:

$$\max \sum_{v \in V} (\mathtt{p} \mathtt{-v} + \mathtt{n} \mathtt{-v}).$$

Now, \mathcal{I} can be solved using a general purpose ILP solver. If it admits any solution I^* , the corresponding conflict-free siphon (say S^*) is maximal. Hence, it makes sense that it does not need to find any other conflict-free siphon of the net that is strictly contained in S^* . To do this, we add to \mathcal{I} a new inequality

$$1 <= \sum_{p \in P \setminus S^*} a - p$$

where a-p is the binary variable presenting place p. Now, we solve \mathcal{I} again to find a new solution. If a new solution I' exists, then let S' be its corresponding conflict-free siphon. Indeed, abide by the newly added inequality, we have $S' \cap (P \setminus S^*) \neq \emptyset$ because there is some a-p with $p \in P \setminus S^*$ such that I'(a-p) = 1. This implies that it is impossible that $S' = S^*$ or $S' \subset S^*$. By the objective function, it means that S' is the conflict-free siphon of maximum cardinality among the conflict-free siphons that are not contained in S^* . Hence, S' is also a maximal conflict-free siphon. Again, we add to \mathcal{I} a new inequality with respect to the newly found siphon. The above process is iterated until \mathcal{I} becomes unfeasible, this means that there is no further maximal conflict-free siphon. Thus, all the maximal conflict-free siphons of the Petri net have been found. We use ... as the ILP solver. TODO: ...

4.5. Computation of special types of trap spaces

In the field of systems biology, biologists may want to compute more special types of trap spaces beyond minimal trap spaces [19]. We shall show that our proposed methods can be easily adjusted to compute popular types of trap spaces. We illustrate the adjustments via the ASP-based method (see Subsection 4.3), but these adjustments are completely applicable for other approaches such as MaxSAT, CLP, and ILP.

First, the work by [36] uses the concept of stable motifs to build the succession diagram of a Boolean network, a summary of the decisions in the network dynamics that lead to successively more restrictive nested stable motifs. The succession diagram is useful for control and decision making on this Boolean network. In particular, the proposed control methods are independent to the update scheme. It has been shown that a stable motif of a Boolean network is equivalent to a maximal trap space of this Boolean network [36]. Hence, it is necessary to develop an efficient method for computing maximal trap spaces of a Boolean network. We shall show how to adjust the ASP-method presented in Subsection 4.3 to compute maximal trap spaces.

We first provide the definition of maximal trap spaces. Let ε be the special trap space of $\mathcal N$ where all the nodes are free. Of course, ε corresponds to the special conflict-free siphon \emptyset . A trap space m is called maximal if $m \neq \varepsilon$ and there is no other trap space m' such that $m' \neq \varepsilon$ and m < m'. Analogously, a conflict-free siphon S is called minimal if $S \neq \emptyset$ and there is no other trap space S' such that $S' \neq \emptyset$ and $S' \subset S$. By using the reasoning similar to the proof of Theorem 3.2, we can easily conclude that a maximal trap space of $\mathcal N$ is equivalent to a minimal conflict-free siphon of its encoded

Petri net \mathcal{P} . Let \mathcal{L} be the ASP characterizing all conflict-free siphons of \mathcal{P} (see Subsection 4.3). Naturally, we need to exclude \emptyset from the solution space of \mathcal{L} (equivalently exclude ε from the set of trap spaces). To do this, we add to \mathcal{L} the ASP rule

$$p-v_1; n-v_1; ...; p-v_n; n-v_n.$$

that ensures that every answer set of \mathcal{L} cannot be empty. Then a set-inclusion minimal answer set of \mathcal{L} is equivalent to a minimal conflict-free siphon of \mathcal{P} , thus a maximal trap space of \mathcal{N} .

591

593

595

602

603

605

606

609

Second, we consider fixed points in Boolean networks. Let s be a fixed point of a Boolean network \mathcal{N} . We have a subspace m corresponding to s as follows: $\forall v \in V, m(v) = s(v)$, i.e., all nodes are fixed in m. Clearly, s is a trap set of \mathcal{N} regardless of the update scheme. Hence, m is a trap space of \mathcal{N} . In addition, since $|S_{\mathcal{N}}[m]| = 1$, m is also a minimal trap space. To compute all fixed points of \mathcal{N} , we can add more constraints to the encoded ASP characterizing all conflict-free siphons (equivalently trap spaces). For every $v \in V$, we add to the encoded ASP the rule

$$p-v$$
; $n-v$.

that ensures that for every conflict-free siphon S, it contains either p-v or n-v for every $v \in V$. Equivalently, the trap space corresponding to S is always a fixed point. Now, the set of answer sets of the encoded ASP is equivalent to the set of fixed points of \mathcal{N} . In particular, when solving the encoded ASP using an ASP solver, we do not need to use the built-in option for computing set-inclusion maximal answer sets. Note that we can also build another ASP characterizing all fixed points of \mathcal{N} based on the equivalence between a fixed point of \mathcal{N} and a deadlock of its Petri net encoding [21]. This approach may give a more compact ASP.

Third, we consider the trap spaces intersecting a given subspace m^* of a Boolean network. A trap space m intersects m^* if and only if $S_{\mathcal{N}}[m] \cap S_{\mathcal{N}}[m^*] \neq \emptyset$. It follows that for every v, if $m^*(v) = 0$ then m(v) = 0 or $m(v) = \star$, if $m^*(v) = 1$ then m(v) = 1 or $m(v) = \star$. For the former case, we add to \mathcal{L} the ASP rule

$$:- n-v.$$

that ensures that m(v) cannot be 1. For the latter case, we add to \mathcal{L} the ASP rule

that ensures that m(v) cannot be 0. Now \mathcal{L} characterizes all trap spaces that intersect m^* .

Finally, we consider the trap spaces that are inside a given subspace m^* of a Boolean network. We first adjust \mathcal{L} to characterize all such trap spaces.

A trap space m is inside m^* if and only if $m(v) = m^*(v)$ for every $v \in D_{m^*}$.

If $m^*(v) = 0$, we add to \mathcal{L} the ASP rule

p-v.

that ensures that m(v) = 0. If $m^*(v) = 1$, we add to \mathcal{L} the ASP rule

n-v.

that ensures that m(v) = 1. It is noted that if we want to compute maximal trap spaces inside m^* , we need to exclude the conflict-free siphon corresponding m^* from the solution space. Specifically, we need to add to \mathcal{L} the ASP rule

$$p\hbox{-}v\hbox{-}\text{i1}; n\hbox{-}v\hbox{-}\text{i1}; \dots; p\hbox{-}v\hbox{-}\text{ik}; n\hbox{-}v\hbox{-}\text{ik}.$$

where $\{v_{i_1}, \ldots, v_{i_k}\}$ is the set of free nodes of m^* . This rule ensures that $m \neq m^*$. In the case that $m^* = \varepsilon$, we have all maximal trap spaces of the original Boolean network.

5. Inconsistent and incomplete data

632

634

636

638

640

641

642

So far we have assumed that we are always able to derive complete and consistent truth-tables (i.e., Boolean functions) that correctly capture the behavior of each node in a Boolean network. However, in practice it is rarely the case that a Boolean network of a biological system is fully understood and indeed, this is one important reason for modeling such networks. The data provided may be incomplete in the sense that information is missing about what happens in certain states, or it may be inconsistent in that we have conflicting information. The result is that the behavior of some nodes under certain conditions may be unknown [37].

Such incomplete and/or inconsistent behavioral information is problematic for Boolean networks with the synchronous update scheme, which are unable to represent the possibility of more than one next state. It can be addressed by using Boolean networks with asynchronous update schemes because of their non-determinism. However, multiple update functions may be

considered for each node, which is problematic for existing analysis methods in Boolean networks. For example, if there are k nodes with two possible Boolean functions, then the analysis methods such as PyBoolNet and mpbn need to consider 2^k possible Boolean networks due to they inputs are Boolean networks with complete Boolean functions. Moreover, one interesting question maybe raised from systems biologists is if there is a common behavioral property (e.g., minimal trap space, attractor) among all possible combinations. To answer this question, the existing analysis methods may consider 2^k possible Boolean networks as well.

648

650

651

652

653

654

655

656

657

658

650

661

663

665

667

669

670

675

We shall show how our Petri net approach presented the previous section can handle the problem of inconsistent and incomplete data. The idea is to build the Petri net encoding of the Boolean network from its partial truth tables. More specifically, for each unknown-output row in the truth table of a node, we create two conflicting transitions representing the two possible outputs (i.e., 0 or 1). For every remaining row, we create transitions as usual following the Petri net encoding [21]. Because of the non-deterministic choice mechanisms of Petri nets, the new Petri net is still equivalent in dynamics to the incomplete Boolean network. One notable advantage of the connection between trap spaces of a Boolean network and conflict-free siphons of its Petri net encoding is that it still holds if the Petri net encoding changes (e.g., more or less transitions) but still maintain the equivalence in dynamics to the Boolean network. Hence, the minimal trap spaces of the incomplete Boolean network can be computed by computing maximal conflict-free siphons of the resulting Petri net. As a consequence, we can easily answer the question on common behavioral property.

Example 5.1. Consider an incomplete Boolean network $\mathcal{N} = (V, F)$ (mentioned in [37]), where $V = (g_1, g_2, g_3)$ and $F = (f_1, f_2, f_3)$ with $f_1 = g_2, f_3 = \neg g_1$. Herein, f_2 is an incomplete function and the partial truth table is as follows.

g_1	g_3	g_2
0	0	0
0	1	0
1	0	?
1	1	1

We have two possible update functions for g_2 : $f_2=g_1 \wedge g_3$ for the case that ?=0 and $f_2=g_1$ for the case that ?=1. Accordingly, we have

two possible Boolean networks. The first one has one minimal trap space: 001. The second one has two minimal trap spaces: 001 and 110. Both have a common minimal trap space: 001. The asynchronous dynamics of the incomplete Boolean network is shown in Figure 2a. The result Petri net following our idea is given in Figure 2b. Herein, $t_{g_2}^3$ and $t_{g_2}^4$ are the two conflicting transitions representing the unknown-output row. This Petri net has only one maximal conflict-free siphon $\{p_{g_1}, p_{g_2}, \overline{p}_{g_3}\}$ corresponding to the common minimal trap space 001.

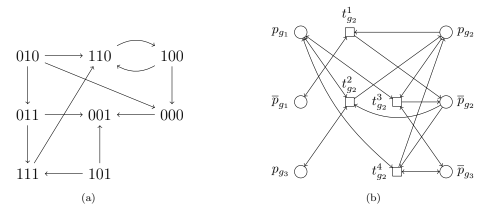


Figure 2: Asynchronous dynamics (self arcs are omitted for simplicity) and new Petri net encoding (the transitions corresponding to g_1 and g_3 are omitted for simplicity) of the incomplete Boolean network of Example 5.1

In summary, we can still perform meaningful analysis on the resulting Petri net. Note that the new Petri net may have more transitions, however the number of additional transitions is only O(2k) where k is the number of unknown-output rows in the truth tables. This is actually a significant advantage, since we do not need to consider 2^k possible Boolean networks. It is worth noting that the approach proposed in [37] only deals with the case that k = 1. Furthermore, as more data becomes available for the underlying biological system, the Petri net model can be refined to reduce the amount of non-determinism (i.e., the number of conflicting transitions) it contains. Hence, Petri nets provide an interesting means of documenting the development of knowledge about a biological system.

6. Motivating example

For a few years now we have been collaborating with biologists who build very large detailed and annotated maps and now wish to analyze the dynamics of the corresponding models. One of the main maps studied this way represents knowledge about the Rheumatoïd Arthritis [38], and was the main motivation for the development of a tool to automatically transform it into an executable Boolean network [6]. In the supplementary material of the paper, an excerpt of the map, focused around the apoptosis (cell death) module is transformed into a model of reasonable size, namely 180 Boolean variables (model F5_RA_apoptosis_executable_module.sbml of supplementary material S3, and model "RA-apoptosis" of Section 7). The study of such model, though, is a big hurdle. Indeed, as stated in the article about another model of the same size: "The size of the CaSQ-inferred MAPK model (181 nodes) made the calculation of stable states a non-realistic endeavour."

In practice, even if there is a huge number of attractors in such a model, obtaining a sample of those can reveal very useful to invalidate the model and lead to further refinement. In particular, it provides a feature-rich alternative to random simulations for this type of very non-deterministic model. Being able to detect that there are inconsistencies with published experimental data in some of the first 1000 attractors, for instance, can lead to a much quicker Systems Biology loop: model, invalidate, refine.

However, using a state-of-the-art tool like PyBoolNet [7] on that model actually fails at the phase of prime-implicant generation. mpbn [9] can return the first 1000 solution within 1.43s, but indeed, it limits the modeling range of the modelers as it does not permit using non-locally-monotonic Boolean functions. This is also true for the Alzheimer model also mentioned in that same article and originally from [39] (F4 file in the original supplementary material, and "Alzheimer" in Table 3), where PyBoolNet also fails at the prime-implicant computation and mpbn does not give any answer because this model is actually non-locally-monotonic. The current practice usually revolves then around fixing some source nodes to plausible values and reducing the model accordingly. While this approach makes sense, it relies on potentially arbitrary decisions, and hides away critical modelling choices that were actually not part of the original Boolean network or even of the starting map.

Using the ASP-based method presented in Section 4.3, it is possible to obtain the first 1000 minimal trap spaces (including ones that contain more

than one state) within 0.19s, which is much quicker than mpbn. Unfortunately since this was not available at the time, the analysis of the model remained very high-level and qualitative, instead of being able to use the rich information of computed minimal trap spaces.

7. Evaluation

To evaluate the performance of the newly proposed methods (implemented as a Python package named Trappist) and the state-of-the-art methods (bioLQM⁶, PyBoolNet [7, 19], and mpbn [9]), we compared them on both PyBoolNet's own model repository and many real-world models from various sources in the literature. It is worth noting that mpbn [9] only handles locally-monotonic models, whereas the other methods can handle general models. To obtain a more comprehensive comparison, we also used random models generated by a third-party software (i.e., BoolNet R package [29]). As explained in Section 6, in our benchmarks, we only searched for the first 1000 minimal trap spaces for each model. It is worth noting that unlike existing analysis shown in the literature, we did not fix specific values for source nodes in all the considered models.

To solve the ASP problems, we used the same ASP solver Clingo [35] and the same configuration as that used in PyBoolNet [7, 19] and mpbn [9]. Specifically, we used the configuration -heuristic=Domain -enum-mod=domRec -dom-mod=3 (subset maximality, equivalent to the deprecated --dom-pref=32 --heuristic=domain --dom-mod=7 used by PyBoolNet). We ran all the benchmarks on a machine whose environment is CPU: Intel® CoreTM i9-11950H 2.60GHz \times 16, 16 GB DDR4 RAM, Ubuntu 20.04.5 LTS. Finally, we set a time limit of three minutes for each model.

All the models and a Jupyter notebook realizing the benchmarks can be found at https://github.com/soli/trap-spaces-as-siphons. These can be run on a Docker image in the cloud by clicking the "Binder" button.

7.1. PyBoolNet repository

Table 1 shows the experimental results on the models from the official PyBoolNet repository⁷. Column n denotes the number of nodes of each model. Column |M| denotes the number of minimal trap spaces and for each

⁶http://colomoto.org/biolgm/doc/tools-trapspace.html

⁷https://github.com/hklarner/pyboolnet/tree/master/pyboolnet/repository

Table 1: Timing comparisons (in seconds) between bioLQM (LQM), PyBoolNet (PBN), mpbn and the four variants of Trappist on the PyBoolNet repository.

							Trappist			
	model	n	M	LQM	PBN	mpbn	SAT	CLP	ILP	ASP
1	arellano_rootstem	9	4	0.13	0.01	0.00	0.00	-	-	0.01
2	$calzone_cellfate$	28	27	0.12	0.02	0.01	0.01	-	-	0.01
3	$dahlhaus_neuroplastoma$	23	32	0.11	0.03	0.01	0.01	-	-	0.01
4	davidich_yeast	10	12	0.11	0.02	0.01	0.01	-	-	0.01
5	dinwoodie_life	15	7	0.11	0.01	0.00	0.01	-	-	0.01
6	$dinwoodie_stomatal$	13	1	0.10	0.01	0.00	0.00	-	-	0.01
7	faure_cellcycle	10	2	0.11	0.02	0.01	0.01	-	-	0.01
8	grieco_mapk	53	18	0.19	0.03	0.02	0.03	-	-	0.02
9	irons_yeast	18	1	0.12	0.03	0.01	0.01	-	-	0.02
10	jaoude_thdiff	103	> 1000	N/A	0.85	0.45	0.56	-	-	0.09
11	klamt_tcr	40	8	0.11	0.01	0.01	0.01	-	-	0.02
12	krumsiek_myeloid	11	6	0.10	0.01	0.00	0.00	-	-	0.01
13	multivalued	13	4	0.10	0.01	0.00	0.00	-	-	0.01
14	n12c5	11	5	0.11	17.83	0.01	0.01	-	-	0.01
15	n3s1c1a	2	2	0.10	0.01	0.00	0.00	-	-	0.01
16	n3s1c1b	2	2	0.09	0.02	0.00	0.00	-	-	0.01
17	n5s3	4	3	0.10	0.02	NM	0.00	-	-	0.01
18	n6s1c2	5	3	0.10	0.02	0.00	0.00	-	-	0.01
19	n7s3	6	3	0.11	0.02	0.00	0.00	-	-	0.01
20	raf	3	2	0.10	0.01	0.00	0.00	-	-	0.01
21	$randomnet_n15k3$	15	3	0.10	0.02	NM	0.01	-	-	0.01
22	$randomnet_n7k3$	7	10	0.10	0.01	NM	0.00	-	-	0.01
23	$remy_tumorigenesis$	34	25	0.15	0.94	0.02	0.02	-	-	0.02
24	saadatpour_guardcell	13	1	0.10	0.06	0.00	0.00	-	-	0.02
25	$selvaggio_emt$	56	> 1000	N/A	0.48	0.28	0.28	-	-	0.09
26	$tournier_apoptosis$	12	3	0.10	0.01	0.00	0.00	-	-	0.01
27	$xiao_wnt5a$	7	4	0.10	0.01	0.00	0.00	-	-	0.01
28	zhang_tlgl	60	156	0.60	0.09	0.09	0.07	-	-	0.04
29	$zhang_tlgl_v2$	60	258	0.64	0.04	0.08	0.11	-	-	0.04

method is given the computation time in seconds, asking only for the first 1000 trap spaces. In the case of bioLQM, "N/A" means that the number of all minimal trap spaces of the model is larger than 1000 and we did not recorded the running time of bioLQM because it always requires to compute all minimal trap spaces. A number in bold indicates a ratio greater than three compared to the best result. "NM" indicates a non-locally-monotonic

model. There are four variants of Trappist: SAT (i.e., the MaxSAT-based method shown in Subsection 4.2), CLP (i.e., the CLP-based method shown in Subsection 4.2), ILP (i.e., the ILP-based method shown in Subsection 4.4), and ASP (i.e., the ASP-based method shown in Subsection 4.3).

As shown in Table 1, for most of the models of the PyBoolNet repository, the results are comparable with all minimal trap spaces found very fast. For 5 of the 29 models, mpbn did not give any answer because it recognized these models as not locally-monotonic. Note that on some very small models, Trappist is sometimes slower than PyBoolNet and/or mpbn, but still significantly under one second. On the contrary, on every model that was a bit challenging for PyBoolNet or mpbn, the new method is far more efficient with speedups between one and two orders of magnitude.

7.2. BBM repository

Currently, a research group has made a great effort for building a collection (called BBM) of real-world Boolean models from various sources used in systems biology. It aims to be a comprehensive collection suitable for benchmarking and testing new tools and methods. It is released and maintained at https://github.com/sybila/biodivine-boolean-models. We here tested all the compared methods on this model repository.

Table 2: Results on the real-world models from the BBM repository.

Method	# failures	avg-lqm (s)	avg-mono (s)	avg-all (s)
bioLQM	9 (134)	12.87	N/A	N/A
PyBoolNet	12	8.87	11.00	13.59
mpbn	2(187)	N/A	2.31	N/A
${\tt Trappist-MaxSAT}$	1	0.03	1.09	1.01
$ ext{Trappist-} ext{CLP}$	-	-	-	-
${ t Trappist-ILP}$	_	-	-	-
${ t Trappist-ASP}$	1	0.05	1.02	0.93

Table 2 shows the experimental results on the 211 real-world models from the BBM repository. Column 2 expresses the numbers of failures (i.e., did not finish the computation within a time limit of three minutes) of each method. For the case of bioLQM, we only considered the models that have at most 1000 minimal trap spaces. The number of such models is 134 (per all 211

models) and is denoted inside the parentheses. For the case of mpbn, we only considered the models that are locally-monotonic. The number of such models is 187 (per all 211 models) and is denoted inside the parentheses. Columns 3-5 express the average running time (in seconds) of each method for the models having at most 1000 minimal trap spaces, the locally-monotonic models, and all the models, respectively. Note that when computing the average running time, if the running time exceeds 180s, it is considered as 180s. From the results shown in Table 2, we reported several observations as follows.

7.3. Selected models

We used a set of real-world Boolean networks lying in various scales collected from numerous bibliographic sources. Most of these models are quite big (in size), complex (i.e., having high average in-degree, which is related to the number of prime-implicants) and have never been fully analyzed. Note that these models are not included in the PyBoolNet and BBM repositories. We then applied bioLQM, PyBoolNet, mpbn, and the four variants of Trappist to computing minimal trap spaces of these real-world models. Table 3 shows the obtained experimental results. "DNF" means that the method did not finish the computation (stopping at the first 1000 minimal trap spaces) within the timeout of two minutes. A number in bold indicates a ratio greater than or equal to 10 compared to the best result. The remaining notations are similar to those in Table 1. Hereafter, we analyze in detail the results with respect to minimal trap space computation.

The first observation is that for 26 of the 33 models (more than 78%), mpbn did not give any answer because it recognized that these models as not locally-monotonic. For 6 of the 33 models where mpbn returned the answers, mpbn and Trappist are comparable in computation time, though surprisingly mpbn appears a bit slower on average. Note however that mpbn was the only tool to provide a solution for the SN-5 model, thus confirming that if the activation function is in the right form, not having to compute the inactivation function's disjunctive normal form can render a difficult problem tractable. However, since mbpn can handle only locally-monotonic models and Trappist can handle general models, it is difficult to further compare between them. Hence, we focus on only comparisons between PyBoolNet and Trappist in the following observations.

The second observation is that the proposed method vastly outperforms PyBoolNet in computational time, on each and every model, and sometimes

Table 3: Timing comparisons (in seconds) between bioLQM (LQM), PyBoolNet (PBN), mpbn and the four variants of Trappist on selected models from the literature.

							Trappist			
	model	n	M	LQM	PBN	mpbn	SAT	CLP	ILP	ASP
1	metastatic [40]	10	4	0.10	0.04	NM	0.01	-	-	0.02
2	Arabidopsis_thaliana [40]	15	8	0.10	0.06	NM	0.01	-	-	0.02
3	p53_high_dna [40]	16	1	0.38	1.76	NM	0.08	-	-	0.14
4	p53_low_dna [40]	16	1	0.41	1.76	NM	0.07	-	-	0.14
5	FT-GRN [41]	23	32	\mathbf{DNF}	\mathbf{DNF}	NM	0.03	-	-	0.19
6	DNA_damage [40]	26	16	0.24	0.33	NM	0.02	-	-	0.05
7	Rho-GTPases [40]	33	2	0.17	0.57	40.39	0.07	-	-	0.11
8	Pluripotency [42]	36	440	\mathbf{DNF}	\mathbf{DNF}	NM	0.16	-	-	0.28
9	Pluripotent [40]	36	276	0.37	0.43	NM	0.07	-	-	0.06
10	Pancreatic_Cancer [40]	43	> 1000	N/A	0.11	0.36	0.17	-	-	0.06
11	Drosophila [43]	52	128	0.33	0.05	0.07	0.06	-	-	0.05
12	Cacace_TdevModel [44]	61	28	1.29	5.67	NM	0.06	-	-	0.08
13	hedgehog [40]	65	> 1000	N/A	\mathbf{DNF}	0.50	0.34	-	-	0.33
14	EMT [36]	69	268	39.22	1.01	0.20	0.12	-	-	0.05
15	Bcell [45]	73	72	0.23	0.04	0.08	0.06	-	-	0.05
16	mast_cell [6]	73	> 1000	N/A	0.09	0.55	0.37	-	-	0.15
17	Corral_ThIL17diff [46]	92	> 1000	N/A	107.57	0.76	0.56	-	-	0.16
18	Adhesion_CIP [47]	121	78	56.81	4.25	0.23	0.17	-	-	0.19
19	EMT_Mech [48]	136	82	\mathbf{DNF}	14.01	0.27	0.20	-	-	0.25
20	macrophage [40]	136	> 1000	N/A	0.54	1.09	0.84	-	-	0.27
21	angiogenesis [40]	141	> 1000	N/A	0.16	1.07	1.06	-	-	0.16
22	angiofull [49]	142	> 1000	N/A	0.17	1.06	0.88	-	-	0.23
23	EMT_Mech_TGFbeta [48]	150	492	\mathbf{DNF}	11.28	0.78	0.69	-	-	0.35
24	RA_apoptosis [6]	180	> 1000	N/A	\mathbf{DNF}	1.43	1.55	-	-	0.19
25	MAPK [6]	181	> 1000	N/A	13.58	1.76	1.51	-	-	0.27
26	Snf1-pathway [50]	202	> 1000	N/A	1.13	1.47	1.43	-	-	0.31
27	T-cell-co-receptor [40]	206	> 1000	N/A	\mathbf{DNF}	1.52	2.26	-	-	0.35
28	TcellCheckPoint [51]	218	> 1000	N/A	4.99	NM	1.96	-	-	0.28
29	Mycobacterium [40]	317	> 1000	N/A	0.42	2.36	4.91	-	-	0.44
30	Leishmania [40]	342	> 1000	N/A	\mathbf{DNF}	2.56	5.62	-	-	0.46
31	Cholocystokinin [6]	383	> 1000	N/A	0.36	2.99	4.81	-	-	0.37
32	Alzheimer [6]	762	> 1000	N/A	\mathbf{DNF}	NM	18.21	-	-	0.79

with orders of magnitude of difference (e.g., for most models in the 100–1000 nodes size range). Note that for all the cases where PyBoolNet did not manage to finish before the timeout, as marked by "DNF" in Table 3, the timeout occurred during the computation of the prime-implicants. Hence, not even a single minimal trap space was output by that method. The computational

advantage is therefore immediately a practical advantage since on the one hand the state-of-the-art method did not allow any analysis whatsoever of the models, and on the other hand the proposed method could provide, very often under one second, the first thousand minimal trap spaces. For modellers having a critical look at a model and in a *model*, *invalidate*, *refine* loop this means a huge difference in the models that are amenable to study.

Note that even with a very restricted time-limit of two minutes, it was possible with the proposed technique to find all minimal trap spaces of small models (roughly under 130 nodes, i.e., considered as quite big up to now). Though it might seem impractical to handle tens of thousands of such possible complex attractors in a manual way, i.e., to compare them to specific experimental conditions and corresponding data, we hope that an automatic analysis of such attractors might become possible with systematic verification methods, not unlike that described in [51]. Since the ASP code is declarative by nature, it is also possible to add to it supplementary constraints coming from the modeler in case one is looking for specific attractors. Finally, sampling from the ASP-generated solutions as is done in [52] would allow for a different type of exploration.

The third observation is that for all the models where PyBoolNet finished before the timeout, once PyBoolNet went through the prime-implicant phase, its ASP solving phase quickly returned the first 1000 minimal trap spaces, all under one second. For these models, the ASP solving phase of the proposed method also took very short time, all under one second. Hence, with the experimental results shown in this paper, the practical differences between our ASP encoding and that of PyBoolNet are not distinctly exposed. The fact that our new ASP encoding is guaranteed to be linear in the number of nodes of the original model does not seem to be crucial here, however a much deeper analysis of those cases remains to be done.

Note that though enumerating the extremal siphons of a Petri net is exponential (see [17] for instance) this is apparently not the bottleneck of the proposed method, showing once again that networks obtained from biochemical models do have a specific structure.

7.4. Randomly generated models

We randomly generated a set of N-K models [1] with network size n in the set $\{100, 150, 200, 250, 300, 350, 400\}$ and K = 3 (i.e., each node has exactly three input nodes). We chose N-K models because they are a useful tool for studying the dynamics of Boolean networks [1, 7]). For each network size, 50

instances were generated using the generateRandomNKNetwork function. In total, we have 350 random models. We then applied the compared methods to these models and recorded the numbers of failures (i.e., failed to obtain the result within a time limit of three minutes) as well as the average running time (inside the parentheses) in each method for each network size n. It is worth noting that N-K models usually have small numbers of minimal trap spaces [7]. Hence, we searched for all solutions in each model, which makes the comparison to bioLQM more comprehensive. In addition, each node has only three input nodes, i.e., the number of prime-implicants of the associated Boolean function is small. Hence, PyBoolNet always passed the phase of computing prime-implicants in every model even within 1s, which enables us to compare the ASP encoding of PyBoolNet and that of Trappist.

Table 4: Results on N-K models.

n	LQM	mpbn	PBN	SAT	CLP	ILP	ASP
100	50 (> 180)	50 (N/A)	0 (0.07)	0 (0.05)	- ()	- ()	0 (0.09)
150	50 (> 180)	50 (N/A)	0(0.14)	0 (0.10)	- ()	- ()	0(0.14)
200	50 (> 180)	50 (N/A)	0(0.43)	0(0.25)	- ()	- ()	0(0.24)
250	50 (> 180)	50 (N/A)	0(1.92)	0(1.04)	- ()	- ()	0 (0.56)
300	50 (> 180)	50 (N/A)	0(9.68)	0(4.46)	- ()	- ()	0(1.83)
350	50 (> 180)	50 (N/A)	1(46.54)	0(20.09)	- ()	- ()	0(6.10)
400	50 (> 180)	50 (N/A)	29 (144.09)	12 (90.36)	- ()	- ()	1 (33.01)

Table 4 shows the experimental results on N-K models. Column n denotes the network size. Columns LQM and PBN show the results of bioLQM and PyBoolNet, respectively. For each method, the number outside the parentheses indicates the number of failures, whereas the number inside the parentheses indicates the average running time (in seconds). Note that when computing the average running time, if the running time exceeds 180s, it is considered as 180s. From these results, we obtained several observations consistent with those obtained for real-world models.

TODO: ...
First, ...
Second, ...
Third, ...

Finally, ...

8. Conclusion

899

900

901

903

905

907

908

909

910

911

912

914

916

918

920

921

922

924

926

927

928

929

930

931

932

In this article we have explored and proved for the first time the equivalence between (minimal) trap spaces of a general Boolean network and (maximal) conflict-free siphons of its Petri net encoding. We have shown several important applications of this finding to studying properties of trap spaces in Boolean networks. As an important practical application of the equivalence, we have proposed a new approach for the computation of minimal trap spaces of Boolean networks, based on the enumeration of maximal conflictfree siphons of Petri nets. We have also proposed the four possible methods using MaxSAT, CLP, ILP, and ASP for implementing the new approach. The proposed methods have been evaluated on many real-world models from the literature and randomly generated models. The experimental results show that the new approach vastly outperforms all the state-of-the-art methods in terms of general Boolean networks and is comparable to the mpbn method even better in average in term of locally-monotonic Boolean networks. We believe that this opens up the way to a much better analysis of large Boolean networks, which is needed with the advent of automatic model-generation pipelines [53].

Although the experimental results show the superiority of our approach to mpbn in general, we however note that there is a model in the BBM repository (with identifier 122) where all the four proposed methods for the new approach did not manage to finish the Petri net conversion before the timeout, whereas mpbn can still handle this model. The model is not very large but its Boolean functions are rather complicated. This points to the fact that our current choice of using a BDD-based translation to obtain that Petri net encoding, though it provides a small/efficient ASP might be too costly to handle the complex models. In such a case, a more naive encoding might provide a much larger ASP program, with many redundant rules, but easier/faster to obtain. The evaluation of the feasibility of such strategy, and of its impact on smaller instances, remains to be done. Recognizing that a model is locally-monotonic and applying in that specific case dedicated strategies as those of mpbn might also be a partial solution.

It is worth noting that there may be possibly other methods for computing minimal/maximal conflict-free siphons in Petri nets, like the methods for generic siphon computation in the field of Petri nets (see [33] for a survey

about these methods). Although these approaches do not directly support the minimal/maximal conflict-free siphon computation now, we plan to investigate them in the future. They could replace our proposed methods if they give significantly better performance. However, the current methods appear to already perform very well even on the biggest models we have considered.

Finally, we think that the links between Petri nets and Boolean networks that we stumbled upon in this method might have deeper roots. Exploring those connections might lead both to interesting topics of research for Petri nets, like a notion of trap-spaces, and for Boolean networks.

15 References

941

942

- [1] L. Glass, S. A. Kauffman, The logical analysis of continuous, non-linear
 biochemical control networks, J. Theor. Biol. 39 (1973) 103–129.
- [2] R. Thomas, Boolean formalisation of genetic control circuits, J. Theor.
 Biol. 42 (1973) 565–583.
- 950 [3] R. Thomas, R. d'Ari, Biological feedback, CRC press, 1990.
- 951 [4] R. Thomas, Regulatory networks seen as asynchronous automata: a logical description, J. Theor. Biol. 153 (1991) 1–23.
- [5] R.-S. Wang, A. Saadatpour, R. Albert, Boolean modeling in systems
 biology: an overview of methodology and applications, Phys. Biol. 9
 (2012) 055001.
- [6] S. S. Aghamiri, V. Singh, A. Naldi, T. Helikar, S. Soliman, A. Niarakis,
 J. Xu, Automated inference of Boolean models from molecular interaction maps using CaSQ, Bioinform. 36 (2020) 4473–4482.
- 959 [7] H. Klarner, A. Bockmayr, H. Siebert, Computing maximal and minimal 960 trap spaces of Boolean networks, Nat. Comput. 14 (2015) 535–544.
- [8] S. Chevalier, C. Froidevaux, L. Paulevé, A. Y. Zinovyev, Synthesis of Boolean networks from biological dynamical constraints using answerset programming, in: International Conference on Tools with Artificial Intelligence, IEEE, 2019, pp. 34–41.

- [9] L. Paulevé, J. Kolčák, T. Chatain, S. Haar, Reconciling qualitative,
 abstract, and scalable modeling of biological networks, Nat. Commun.
 11 (2020) 1–7.
- ⁹⁶⁸ [10] M. Noual, D. Regnault, S. Sené, About non-monotony in Boolean automata networks, Theor. Comput. Sci. 504 (2013) 12–25.
- [11] J. L. Peterson, Petri net theory and the modeling of systems, Prentice
 Hall PTR, 1981.
- 972 [12] T. Murata, Petri nets: Properties, analysis and applications, Proc. IEEE 77 (1989) 541–580.
- V. N. Reddy, M. L. Mavrovouniotis, M. N. Liebman, Petri net representations in metabolic pathways, in: International Conference on Intelligent Systems for Molecular Biology, AAAI, 1993, pp. 328–336.
- 977 [14] I. Zevedei-Oancea, S. Schuster, Topological analysis of metabolic net-978 works based on Petri net theory, Silico Biol. 3 (2003) 323–345.
- 979 [15] M. A. Blätke, M. Heiner, W. Marwan, Biomodel engineering with Petri 980 nets, in: Algebraic and Discrete Mathematical Methods for Modern 981 Biology, Elsevier, 2015, pp. 141–192.
- [16] O. Oanea, H. Wimmel, K. Wolf, New algorithms for deciding the siphon trap property, in: International Conference on Applications and Theory
 of Petri Nets, Springer, 2010, pp. 267–286.
- [17] F. Nabli, T. Martinez, F. Fages, S. Soliman, On enumerating minimal siphons in Petri nets using CLP and SAT solvers: theoretical and practical complexity, Constraints An Int. J. 21 (2016) 251–276.
- [18] V. Trinh, B. Benhamou, K. Hiraishi, S. Soliman, Minimal trap spaces of logical models are maximal siphons of their Petri net encoding, in: International Conference on Computational Methods in Systems Biology, Springer, 2022, pp. 158–176.
- 992 [19] H. Klarner, A. Streck, H. Siebert, PyBoolNet: a python package for the 993 generation, analysis and visualization of Boolean networks, Bioinform. 994 33 (2017) 770–772.

- [20] L. C. Fontanals, E. Tonello, H. Siebert, Control strategy identification
 via trap spaces in Boolean networks, in: International Conference on
 Computational Methods in Systems Biology, Springer, 2020, pp. 159–
 175.
- ⁹⁹⁹ [21] T. Chatain, S. Haar, L. Jezequel, L. Paulevé, S. Schwoon, Characterization of reachable attractors using Petri net unfoldings, in: International Conference on Computational Methods in Systems Biology, Springer, 2014, pp. 129–142.
- [22] C. Chaouiya, E. Remy, P. Ruet, D. Thieffry, Qualitative modelling of genetic networks: From logical regulatory graphs to standard Petri nets,
 in: International Conference on Applications and Theory of Petri Nets,
 Springer, 2004, pp. 137–156.
- ¹⁰⁰⁷ [23] C. Chaouiya, A. Naldi, E. Remy, D. Thieffry, Petri net representation of multi-valued logical regulatory graphs, Nat. Comput. 10 (2011) 727–750.
- 1009 [24] T. Chatain, S. Haar, J. Kolcák, L. Paulevé, A. Thakkar, Concurrency 1010 in Boolean networks, Nat. Comput. 19 (2020) 91–109.
- [25] C. Chaouiya, D. Bérenguier, S. M. Keating, A. Naldi, et al., SBML qualitative models: a model representation format and infrastructure to foster interactions between qualitative modelling formalisms and tools, BMC Syst. Biol. 7 (2013) 1–15.
- [26] S. M. Keating, D. Waltemath, M. König, F. Zhang, et al., SBML Level
 3: an extensible format for the exchange and reuse of biological models,
 Mol. Syst. Biol. 16 (2020) e9110.
- [27] C. Chaouiya, A. Naldi, D. Thieffry, Logical modelling of gene regulatory
 networks with GINsim, in: Bacterial Molecular Networks, Springer,
 2012, pp. 463–479.
- [28] A. Naldi, P. T. Monteiro, C. Müssel, C. for Logical Models, Tools,
 H. A. Kestler, D. Thieffry, I. Xenarios, J. Saez-Rodriguez, T. Helikar,
 C. Chaouiya, Cooperative development of logical modelling standards
 and tools with CoLoMoTo, Bioinform. 31 (2015) 1154–1159.

- 1025 [29] C. Müssel, M. Hopfensitz, H. A. Kestler, BoolNet an R package for generation, reconstruction and analysis of Boolean networks, Bioinform. 26 (2010) 1378–1380.
- [30] D. Angeli, P. D. Leenheer, E. Sontag, A Petri net approach to persistence
 analysis in chemical reaction networks, in: Biology and Control Theory:
 Current Challenges, Springer, 2007, pp. 181–216.
- [31] D. Angeli, P. D. Leenheer, E. D. Sontag, Persistence results for chemical reaction networks with time-dependent kinetics and no global conservation laws, SIAM J. Appl. Math. 71 (2011) 128–146.
- [32] E. Degrand, F. Fages, S. Soliman, Graphical conditions for rate independence in chemical reaction networks, in: International Conference on Computational Methods in Systems Biology, Springer, 2020, pp. 61–78.
- 1037 [33] G. Liu, K. Barkaoui, A survey of siphons in Petri nets, Inf. Sci. 363 (2016) 198–220.
- 1039 [34] A. Ignatiev, A. Morgado, J. Marques-Silva, RC2: an efficient MaxSAT solver, J. Satisf. Boolean Model. Comput. 11 (2019) 53–64.
- [35] M. Gebser, B. Kaufmann, R. Kaminski, M. Ostrowski, T. Schaub,
 M. Schneider, Potassco: The Potsdam answer set solving collection,
 AI Commun. 24 (2011) 107–124.
- 1044 [36] J. C. Rozum, J. G. T. Zañudo, X. Gan, D. Deritei, R. Albert, Parity 1045 and time reversal elucidate both decision-making in empirical models 1046 and attractor scaling in critical Boolean networks, Sci. Adv. 7 (2021) 1047 eabf8124.
- 1048 [37] L. J. Steggles, R. Banks, O. Shaw, A. Wipat, Qualitatively modelling and analysing genetic regulatory networks: a Petri net approach, Bioinform. 23 (2006) 336–343.
- [38] V. Singh, M. Ostaszewski, G. D. Kalliolias, G. Chiocchia, R. Olaso,
 E. Petit-Teixeira, T. Helikar, A. Niarakis, Computational systems biology approach for the study of rheumatoid arthritis: from a molecular
 map to a dynamical model, Genom. Comput. Biol. 4 (2018) 100050.

- [39] S. Ogishima, S. Mizuno, M. Kikuchi, A. Miyashita, R. Kuwano,
 H. Tanaka, J. Nakaya, AlzPathway, an updated map of curated signaling pathways: towards deciphering Alzheimer's disease pathogenesis,
 in: Systems Biology of Alzheimer's Disease, Springer, 2016, pp. 423–432.
- [40] C. Kadelka, T.-M. Butrie, E. Hilton, J. Kinseth, H. Serdarevic, A metaanalysis of Boolean network models reveals design principles of gene regulatory networks, arXiv preprint arXiv:2009.01216 (2020).
- [41] E. C. Chávez-Hernández, S. Quiroz, B. García-Ponce, E. R. Álvarez-Buylla, The flowering transition pathways converge into a complex gene regulatory network that underlies the phase changes of the shoot apical meristem in Arabidopsis thaliana, Front. Plant Sci. 13 (2022) 852047.
- [42] A. Yachie-Kinoshita, K. Onishi, J. Ostblom, M. A. Langley, E. Posfai,
 J. Rossant, P. W. Zandstra, Modeling signaling-dependent pluripotency
 with Boolean logic to predict cell fate transitions, Mol. Syst. Biol. 14
 (2018) e7952.
- 1070 [43] M. R. Vega, Analyzing toys models of Arabidopsis and Drosphila us-1071 ing Z3 SMT-LIB, in: Independent Component Analyses, Compressive 1072 Sampling, Wavelets, Neural Net, Biosystems, and Nanoengineering XII, 1073 volume 9118, SPIE, 2014, pp. 240–254.
- [44] E. Cacace, S. Collombet, D. Thieffry, Logical modeling of cell fate specification—Application to T cell commitment, in: Current Topics in Developmental Biology, Elsevier, 2020, pp. 205–238.
- ¹⁰⁷⁷ [45] P. Dutta, L. Ma, Y. Ali, P. M. Sloot, J. Zheng, Boolean network model-¹⁰⁷⁸ ing of B-cell apoptosis and insulin resistance in type 2 diabetes mellitus, ¹⁰⁷⁹ BMC Syst. Biol. 13 (2019) 1–12.
- 1080 [46] K. F. Corral-Jara, C. Chauvin, W. Abou-Jaoudé, M. Grandclaudon,
 1081 A. Naldi, V. Soumelis, D. Thieffry, Interplay between SMAD2 and
 1082 STAT5A is a critical determinant of IL-17A/IL-17F differential expres1083 sion, Mol. Biomed. 2 (2021) 1–16.
- [47] E. Guberman, H. Sherief, E. R. Regan, Boolean model of anchorage dependence and contact inhibition points to coordinated inhibition but semi-independent induction of proliferation and migration, Comput. Struct. Biotechnol. J. 18 (2020) 2145–2165.

- [48] E. Sullivan, M. Harris, A. Bhatnagar, E. Guberman, I. Zonfa, E. R. Regan, Boolean modeling of mechanosensitive Epithelial to Mesenchymal Transition and its reversal, bioRxiv (2022).
- 1091 [49] N. Weinstein, L. Mendoza, I. Gitler, J. Klapp, A network model to
 explore the effect of the micro-environment on endothelial cell behavior
 during angiogenesis, Front. Physiol. 8 (2017) 960.
- [50] T. Lubitz, N. Welkenhuysen, S. Shashkova, L. Bendrioua, S. Hohmann, E. Klipp, M. Krantz, Network reconstruction and validation of the Snf1/AMPK pathway in baker's yeast based on a comprehensive literature review, npj Syst. Biol. Appl. 1 (2015) 1–10.
- [51] C. Hernandez, M. Thomas-Chollier, A. Naldi, D. Thieffry, Computational verification of large logical models—Application to the prediction of T cell response to checkpoint inhibitors, Front. Physiol. 11 (2020) 558606.
- [52] S. Chevalier, V. Noël, L. Calzone, A. Y. Zinovyev, L. Paulevé, Synthesis and simulation of ensembles of Boolean networks for cell fate decision, in: International Conference on Computational Methods in Systems Biology, Springer, 2020, pp. 193–209.
- [53] M. Ostaszewski, A. Niarakis, A. Mazein, I. Kuperstein, R. Phair,
 A. Orta-Resendiz, V. Singh, S. S. Aghamiri, M. L. Acencio, E. Glaab,
 et al., COVID19 Disease Map, a computational knowledge repository of
 virus-host interaction mechanisms, Mol. Syst. Biol. 17 (2021) e10387.