New ideas for scalable attractor detection and control of asynchronous Boolean networks

Van-Giang Trinh¹, Samuel Pastva², Kyu Hyong Park³ and Jordan Rozum⁴

¹LIS, Aix-Marseille University, France
 ²Institute of Science and Technology, Austria
 ³Pennsylvania State University, United States of America
 ⁴Binghamton University, United States of America

June 6, 2023

Attractor detection and control of asynchronous Boolean networks

Two most important issues in Boolean networks research with many applications in various fields [Akutsu, 2018, Schwab et al., 2020].

Usually, the attractor detection issue is the preceding step of the control issue.

Unfortunately, both of them are very challenging, especially for large-scale networks.

Many methods have been proposed.

Attractor detection methods

```
BDD-based methods: genYsis [Garg et al., 2008], geneFatt [Zheng et al., 2013], CABEAN [Mizera et al., 2019, Su and Pang, 2021a].
```

Reduction-based methods: AEON [Benes et al., 2021].

```
Trap space-based methods: PyBoolNet <sup>1</sup> [Klarner et al., 2017], pystablemotifs [Rozum et al., 2021b, Rozum et al., 2021a].
```

Feedback vertex set-based methods: FVS-ABN [Trinh et al., 2020], iFVS-ABN [Giang and Hiraishi, 2021], mtsNFVS [Trinh et al., 2022b].

All of them have advantages and disadvantages. To our best knowledge, none of them can robustly handle complex networks with thousands of nodes.

¹It is not an exact method.

Control methods

BDD-based and decomposition-based methods: CABEAN [Su and Pang, 2021b, Su and Pang, 2021a].

Reduction-based methods: AEON [Brim et al., 2021].

Trap space-based methods:

PyBoolNet [Fontanals et al., 2020, Cifuentes-Fontanals et al., 2021], pystablemotifs [Zañudo and Albert, 2015, Rozum et al., 2021a].

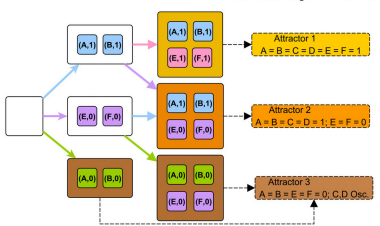
4 / 78

Objective

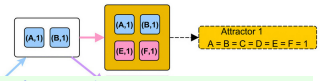
pystablemotifs and mtsNFVS have much more potential to reach the genome scale.

Both of them have their own advantages and disadvantages, but fortunately they can be complementary to each other.

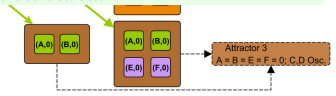
Develop a more efficient method that can benefit both the attractor detection and control issues, and in particular can handle networks at the genome scale.



E Succession diagram and attractors

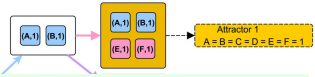


- stable motif = maximal trap space
- "quasi" attractor = minimal trap space
- motif-avoidant attractor = ?



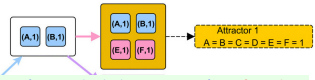
6/78

E Succession diagram and attractors

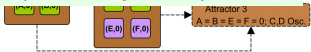


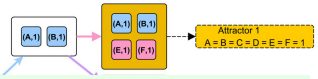
- pystablemotifs is beneficial to not only the attractor detection problem but also the control problem as it provides a succession diagram that can be efficiently used for control.
- If solely considering attractor identification, pystable-motifs seems to have disadvantages as compared to other methods such as AEON, iFVS-ABN, and mtsNFVS.

40 40 40 40 40 40 40 40

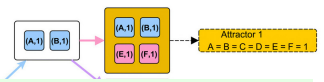


- pystablemotifs must check the existence of motif-avoidant attractors.
- Although it uses time reversal to prune the considered state space, the remaining part (called the terminal restriction space) may be still too large to be analyzed.





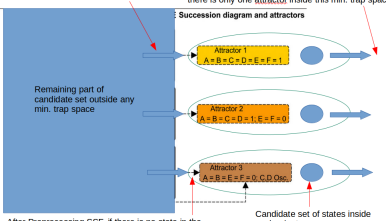
- pystablemotifs relies on PyBoolNet to compute stable motifs (i.e., maximal trap spaces). However, PyBoolNet does not work well with networks with Boolean functions having many input nodes.
- The reason for this is PyBoolNet mainly relies on primeimplicants computation, whereas the number of primeimplicants may be too huge if there are many input nodes.



- The size of the succession diagram may be too large.
- Apart from the theoretical worst case (n_{sm} ! where n_{sm} is the number of stable motifs), the number of stable motifs may be actually too large because for example, the network has two many source nodes (node A is a source node if $f_A = A$).

Preprocessing SSF (shrink the candidate set)

After Preprocessing SSF, if there is only one state, we do not need the reachability check. In this case, there is only one attractor inside this min. trap space.



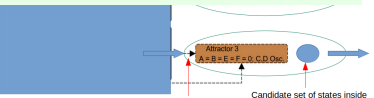
After Preprocessing SSF, if there is no state in the remaining part, we do not need the reachability check. In this case, there is no attractor outside any min. trap space. each min. trap space

Preprocessing SSF (shrink the candidate set)

After <u>Preprocessing SSF</u>, if there is only one state, we do not need the <u>reachability</u> check. In this case, there is only one attractor inside this min. trap space.



- The candidate set F (the blue part) is computed based on an negative feedback vertex set U^- of the ABN.

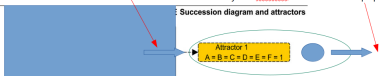


After Preprocessing SSE, if there is no state in the remaining part, we do not need the reachability check. In this case, there is no attractor outside any min. trap space.

Candidate set of states inside each min. trap space

Preprocessing SSF (shrink the candidate set)

After <u>Preprocessing SSF</u>, if there is only one state, we do not need the <u>reachability</u> check. In this case, there is only one <u>attractor</u> inside this min. trap space.



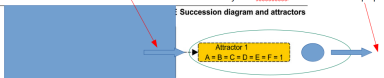
- If the size of the negative feedback vertex set is too large, there may be too many states in the candidate set F (i.e., the set of fixed points of the reduced STG), leading to extremely longer time for the computation of F, Preprocessing SSF, and maybe the reachability analysis.
- Note that real-world models usually have **small** minimum negative feedback vertex sets.

After Preprocessing SSE, if there is no state in the remaining part, we do not need the reachability check. In this case, there is no attractor outside any min. trap space.

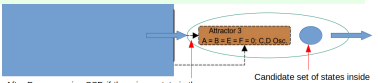
canuluate set of states inside each min. trap space

Preprocessing SSF (shrink the candidate set)

After Preprocessing SSF, if there is only one state. we do not need the reachability check. In this case, there is only one attractor inside this min. trap space.



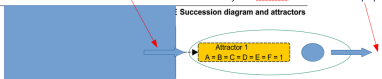
- mtsNFVS also uses PyBoolNet to compute the candidate set as well as the set of minimal trap spaces. Again, it will not work well with the case of many input nodes.



After Preprocessing SSF, if there is no state in the remaining part, we do not need the reachability check. In this case, there is no attractor outside any min. trap space. each min. trap space

Preprocessing SSF (shrink the candidate set)

After <u>Preprocessing SSF</u>, if there is only one state, we do not need the <u>reachability</u> check. In this case, there is only one <u>attractor</u> inside this min. trap space.



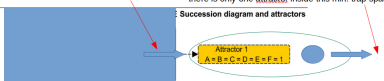
- If some motif-avoidant attractors exist or the results of Preprocessing SSF are not good enough, mtsNFVS still must check the reachability in asynchronous BNs, which is PSPACE-complete in general.
- Moreover, the target set for the reachability analysis is maybe small (i.e., the union of all minimal trap spaces).

After Preprocessing SSF, if there is no state in the remaining part, we do not need the reachability check. In this case, there is no attractor outside any min. trap space.

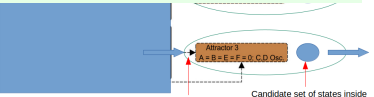
Candidate set of states inside each min. trap space

Preprocessing SSF (shrink the candidate set) we do not there is

After <u>Preprocessing SSF</u>, if there is only one state, we do not need the <u>reachability</u> check. In this case, there is only one attractor inside this min. trap space.



- Currently, mtsNFVS focuses solely on the attractor detection problem.



After Preprocessing SSF, if there is no state in the remaining part, we do not need the reachability check. In this case, there is no attractor outside any min. trap space.

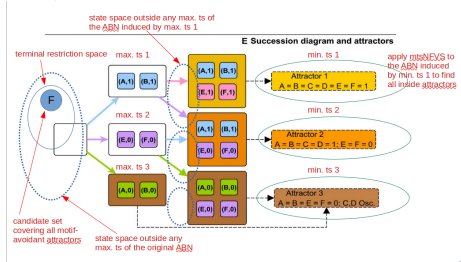
Candidate set of states insid each min. trap space

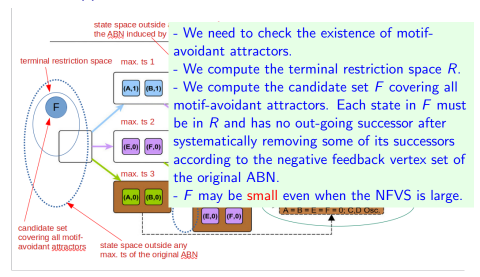
New idea

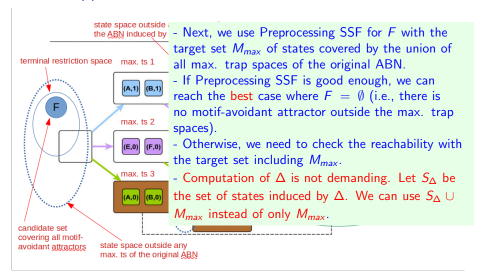
In principle, our new idea is to combine pystablemotifs and mtsNFVS to obtain a more efficient method that can benefit both the attractor detection and control problems.

Hereafter, we will present the general approach along with many new findings that can speedup the whole process.

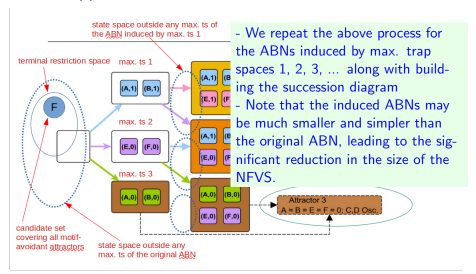
It follows the process of pystablemotifs (i.e., building the succession diagram).

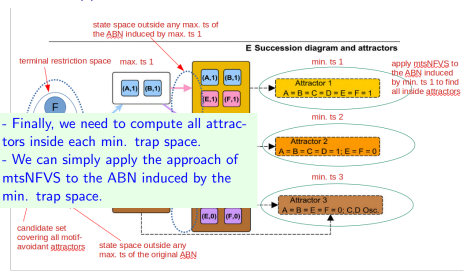


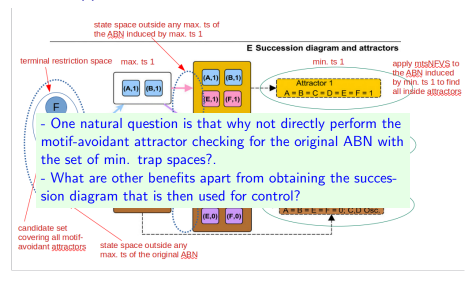


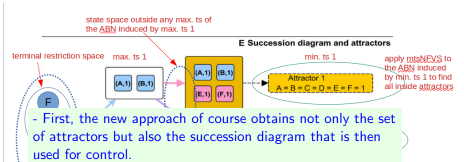


9/78

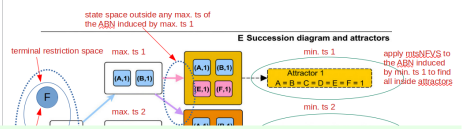








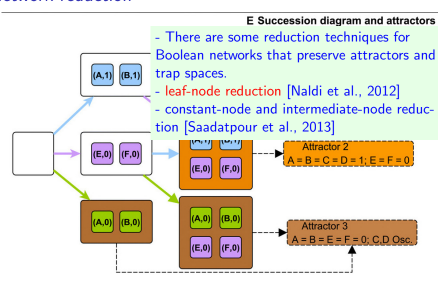
- Second, we observed that in this case the terminal restriction space may be too huge, leading to the candidate set F may be too large (hence, hard to be computed as well).
- Third, M_{max} is much broader than M_{min} in most cases, avoic hence using max. trap spaces can help Preprocessing SSF to reach good cases more easily than using min. trap spaces.

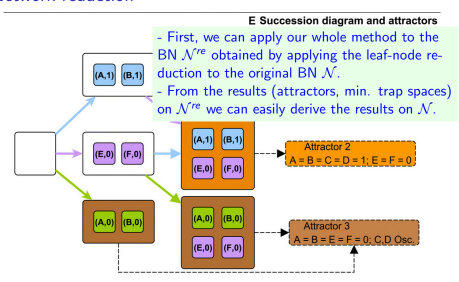


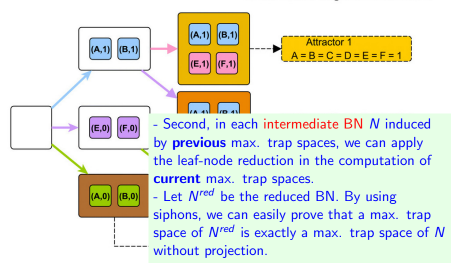
Jordan: We need to consider the percolation of a stable motif (LDOI) along with building the succession diagram. The percolation is based on PyBoolNet.

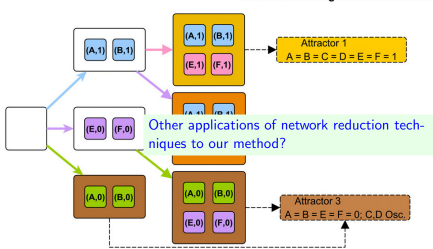
Sam: When we have already obtained the stable motif (i.e., max. trap space), we can perform the percolation easily by algebraically propagating the fixed values.

Giang: Sure. It is possible. Note that we can store Boolean functions as Boolean expressions or BDDs (not as prime-implicants).

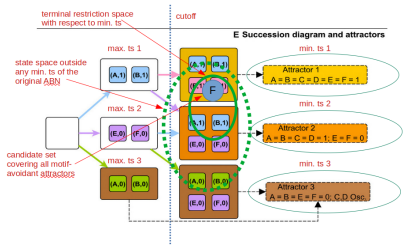


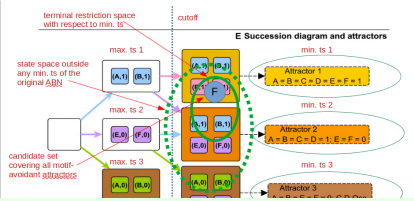




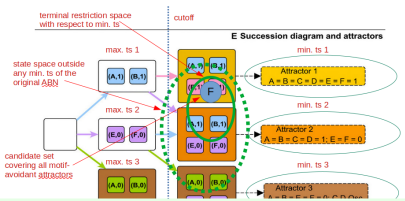


The above approach is promising. However, how to deal with the problem of large succession diagrams?.

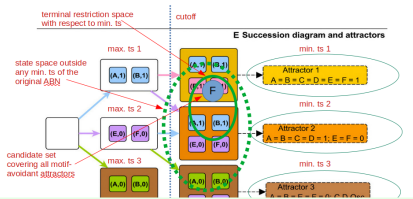




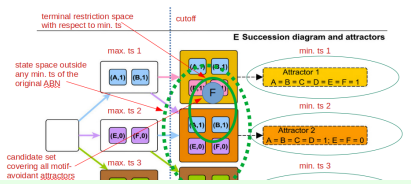
- When the size of the succession diagram becomes too large (maybe exceeding a threshold), we stop its construction.
- However, we still need to check the existence of motif-avoidant attractors.



- First, we compute all min. trap spaces of the original ABN.
- Second, we compute the terminal restriction space R with respect to these min. trap spaces.
- Next, we get the intersection between R and the state spaces induced by the already computed max. trap spaces and their LDOIs (e.g., max. ts 1, 2, 3).

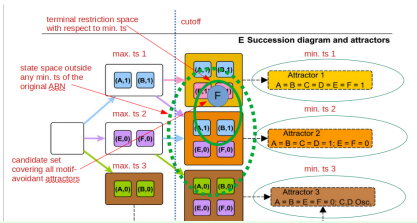


- Then, we compute the candidate set ${\it F}$ with respect to the NFVS of the original ABN.
- Each state in F must be in the intersected set obtained beforehand.
- F may be small even when the NFVS is large because the intersected set may be not too large.



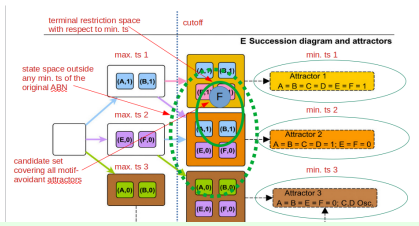
Next, we use Preprocessing SSF for F with the target set M_{min} of states covered by the union of all min. trap spaces of the original ABN.

- If Preprocessing SSF is good enough, we can reach the best case where $F=\emptyset$ (i.e., there is no motif-avoidant attractor).
- Otherwise, we need to check the reachability with the target set including M_{min} .



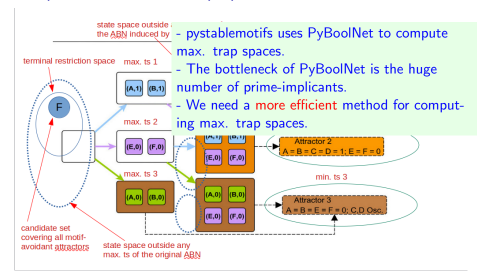
- Finally, we need to compute all attractors inside each min. trap space.
- We can simply apply the approach of mtsNFVS to the ABN induced by the min. trap space.

Deal with the problem of large succession diagrams

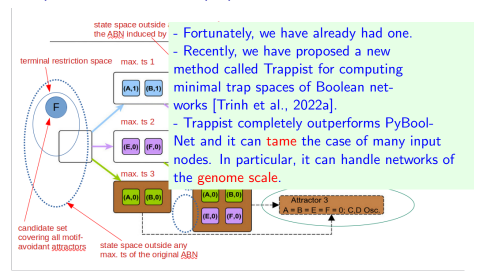


- After finishing the above process, we will obtain the whole attractor landscape and the partial succession diagram.

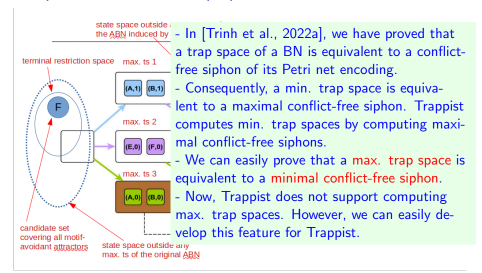
Computation of max. trap spaces

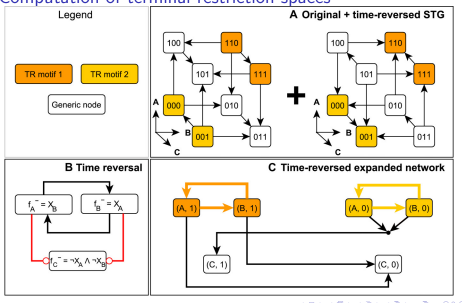


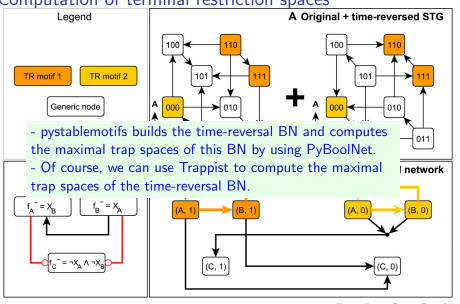
Computation of max. trap spaces

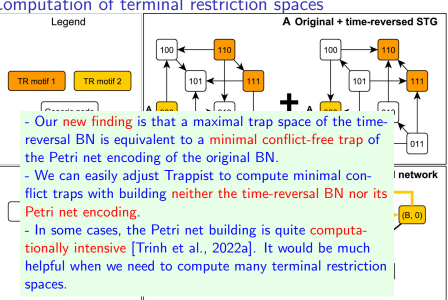


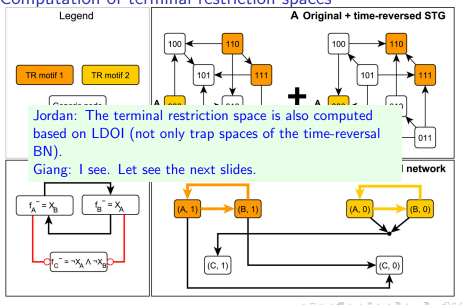
Computation of max. trap spaces

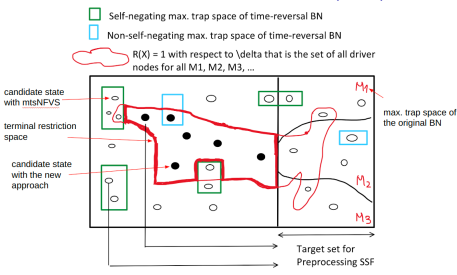




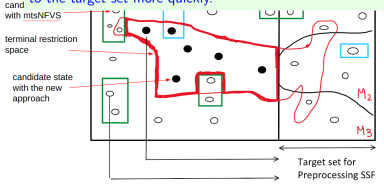








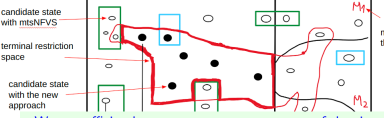
- I emphasize an advantage of the new approach.
- The new candidate set seems to have shorter distances to the target set.
- This is better for Preprocessing SSF as it may converge to the target set more quickly.



max. trap space of the original BN

- Self-negating max. trap space of time-reversal BN
- Non-self-negating max. trap space of time-reversal BN

R(X) = 1 with respect to \delta that is the set of all driver nodes for all M1, M2, M3, ...



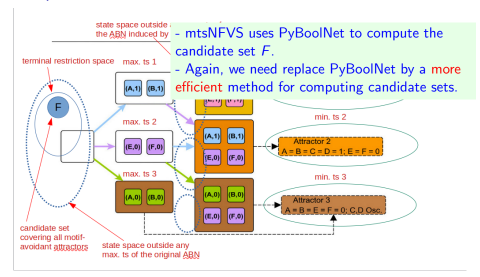
max. trap space of the original BN

- We can efficiently compute max. trap spaces of the time-reversal BN by using Trappist.
- The self-negation of a max. trap space of the timereversal BN can be checked by computing its LDOI in the original BN.

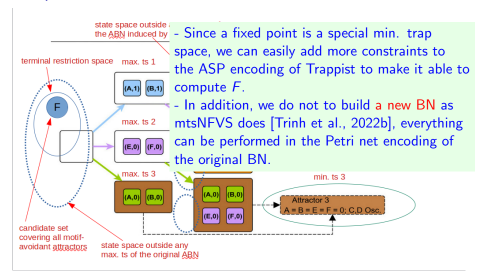
- Self-negating max. trap space of time-reversal BN Non-self-negating max. trap space of time-reversal BN R(X) = 1 with respect to \delta that is the set of all driver nodes for all M1, M2, M3, ... candidate state 0 0 with mtsNFVS 0 max, trap space of the original BN terminal restriction space candidate state with the new approach
- Δ is the set of all virtual nodes that individually drive any stable motif.
- A question is how to compute Δ efficiently? Note that we need to compute Δ many times in the whole process.
- If the computation is expensive, we can ignore R(X). In this case, I hope that self-negating max. trap spaces are good enough.

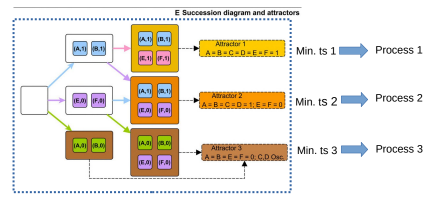
- Self-negating max. trap space of time-reversal BN Non-self-negating max. trap space of time-reversal BN R(X) = 1 with respect to \delta that is the set of all driver nodes for all M1, M2, M3, ... candidate state 0 \circ with mtsNFVS 0 max, trap space of the original BN terminal restriction space candidate state with the new approach
- The set of retained values B largely affects the size of the candidate set [Trinh et al., 2020].
- One interesting question is that how we can propose heuristics for setting B based on the information about the terminal restriction space to get a smaller candidate set?

Computation of candidate sets

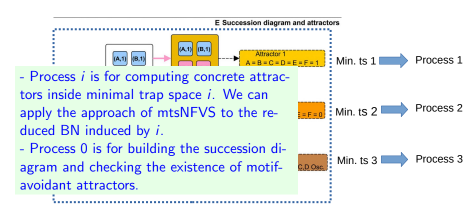


Computation of candidate sets

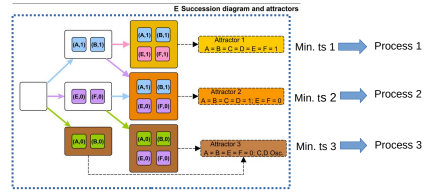




Process 0

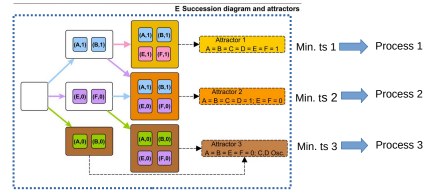


Process 0



Process 0

- Processes 0, 1, 2, ... can be completely paralleled.



Process 0

- Process 0 can even be divided into many paralleled subprocesses.

Target control

After finishing the attractor detection phase, we will obtain the whole attractor landscape as well as the partial/full succession diagram. If the succession diagram is full, its size should be moderate.

The next step is how to compute minimum control policies that drive the network dynamics into a target attractor/min. trap space from any initial state?

There are some existing methods for the target control of asynchronous Boolean networks: CABEAN [Su and Pang, 2021a], PyBoolNet [Fontanals et al., 2020, Cifuentes-Fontanals et al., 2021], pystablemotifs [Zañudo and Albert, 2015, Rozum et al., 2021a], AEON [Brim et al., 2021].

Target control: our observations I

CABEAN always ensures that the resulting control policies are minimum. It relies on the calculation of strong basin of attraction, which may be very computationally expensive.

PyBoolNet exploits the information on trap spaces to find control policies. However, it does not ensure that the resulting control policies are minimum. It uses model checking to improve the quality of the control policies, which may be very computationally expensive as well. Moreover, its scalability may be limited by the necessity to scan through the available perturbations.

pystablemotifs uses the succession diagram to find control policies. However, it does not ensure that the resulting control policies are minimum.

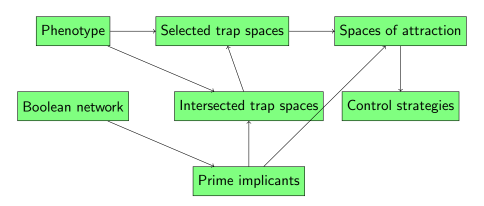
Target control: our observations II

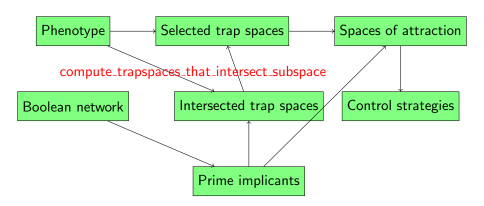
AEON uses methods based on the computation of the strong basin, but with symbolic representation to exhaustively analyse all perturbations simultaneously. A postprocessing step is then needed to extract the minimal perturbations from the final result. Alternatively, if the network has parameters, such postprocessing can also be used to extract perturbations with favourable robustness within the parameter space. The method is mainly limited by the complexity of the symbolic state space search for all available perturbations.

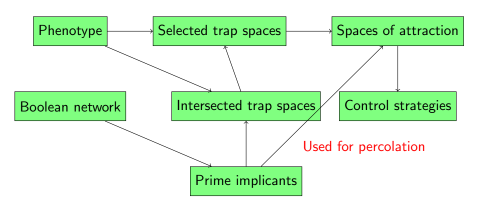
In the reported experiments [Su and Pang, 2021b, Su and Pang, 2021a], pystablemotifs is more time-efficient in most cases, though it did not return the minimum results in some cases.

Target control: our observations III

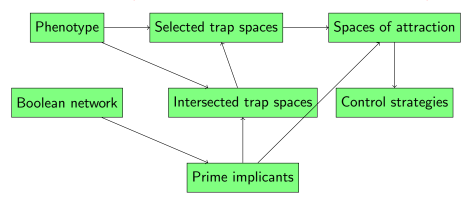
In our opinion, pystablemotifs has much more potential than the others. We should dive into it. Of course, we need to improve both its time-efficiency and accuracy.

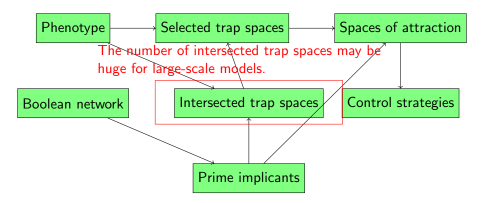


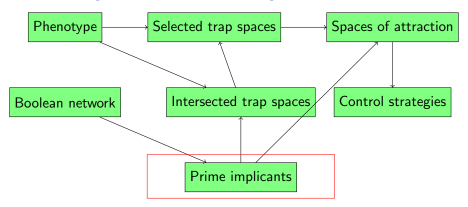




Exploration of all possible sub-spaces whose numbers of fixed variables are up to min(m, n) (m is the maximum size of the control strategies)

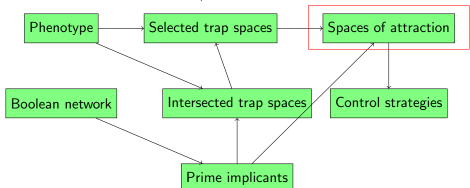




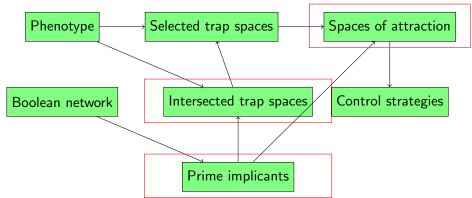


The number of prime implicants may be huge for models with high average connectivity.

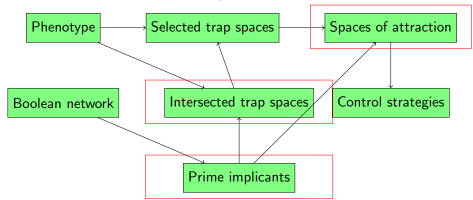
The number of explored sub-spaces is exponential in n in the worst case.



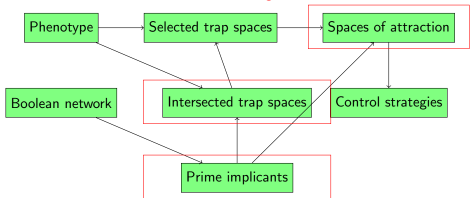
The number of explored sub-spaces is exponential in n in the worst case.



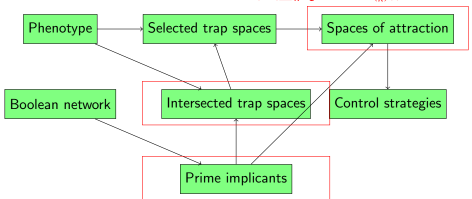
Propose a new implementation for this step based on ASP.



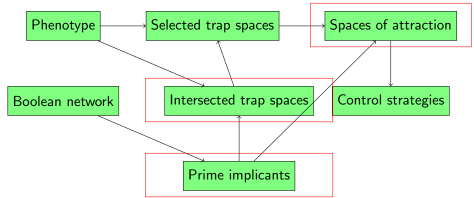
It is possible to optimize the ASP encoding.



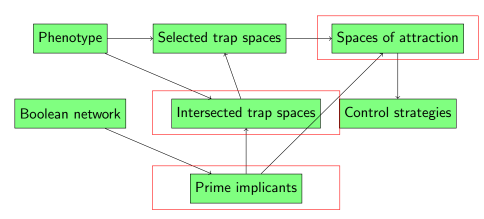
However, even with an optimal ASP encoding, the running time of the ASP solver may still large because the search space of the problem itself is intrinsically huge $(\mathcal{O}(\sum_{k=1}^{\min(m,n)} 2^i \times \binom{n}{k}))$.

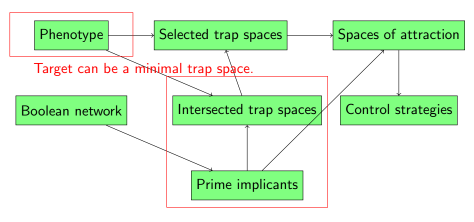


Hence, we retain the opinion that pystablemotifs has much more potential than the others.



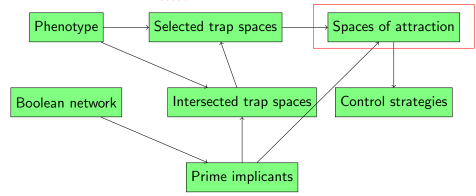
Some potential improvements to PyBoolNet [Laura et al., 2022]

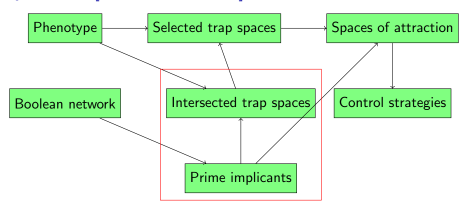




Use Trappist.

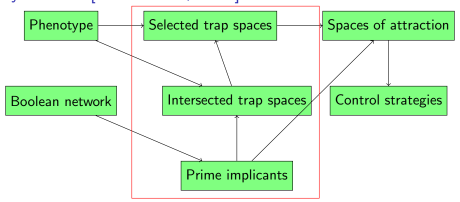
However, we need to adjust the ASP encoding because it relies on prime implicants. The new ASP encoding should rely on Boolean functions instead.





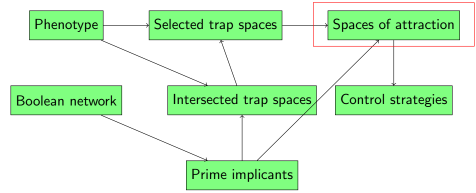
The bottleneck on the number of intersected trap spaces still remains.



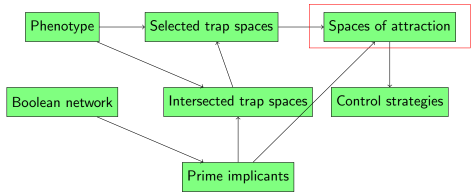


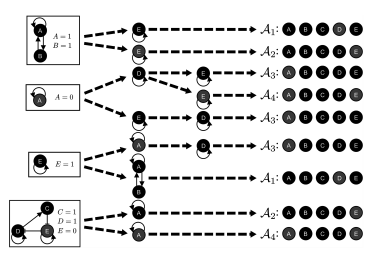
We can use BDDs to compute and represent the set of intersected trap spaces (also the set of selected trap spaces).

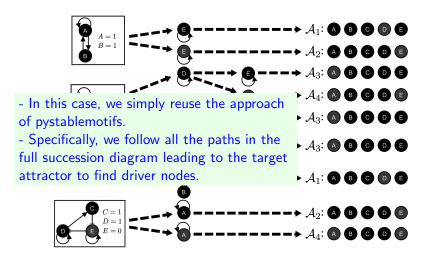
However, we need to adjust the ASP encoding because it relies on the set of selected trap spaces. We need to encode the BDD as ASP rules.

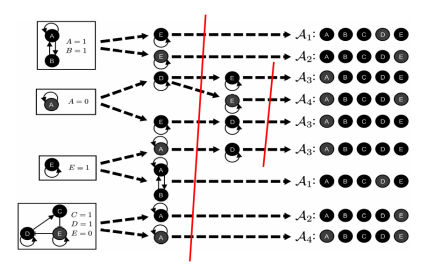


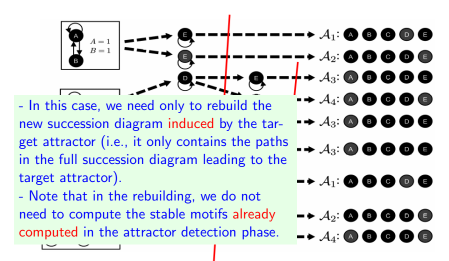
Along with optimizing the ASP encoding, I wonder if we can write a conference paper based on these new ideas. Of course, the first thing we need to do is to develop a concrete method, implement it, and then test it on large real-world models.

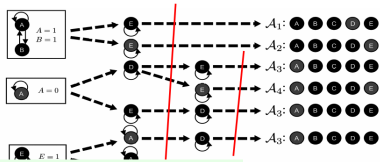








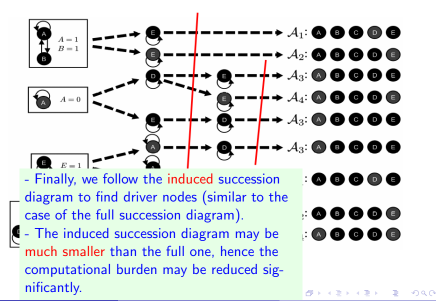


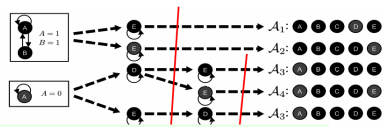


- One question may be whether we need to compute both $\{E=1\}$ and $\{E=0\}$. No, we do not need. We have already known the target attractor, hence we can add constraints to the ASP encoding to guide Trappist for searching only stable motifs that are compatible with the target attractor \mathcal{A}_3 .



$$- \rightarrow \mathcal{A}_4$$
: A B C D E





Jordan: The control approach of pystablemotifs is independent to the update scheme. CABEAN is designed for asynchronous Boolean networks. Hence, the problem of not obtaining the minimum control polices is not really mattered with pystablemotifs.

Giang: I understand. Let me recall the control example.

(C) (D) (E)

Control features

					Control target	ntrol target			Update		
Method		Tool	Steady state	Trap space attractor	Complex attractor	Subspace	Arbitrary subset	Async	Sync	Gen. async	
Basins	ВА	CABEAN [14]	✓	✓	✓	-	-	1	_	_	
Stable motifs	SM	StableMotifs [7]	✓	✓	-	-	-,	✓		-	
Percolation-only	PO	Caspo [21]	✓	✓	-	✓	-	✓	✓	✓	
Trap spaces	TS	PyBoolNet [9]	✓	✓	-	✓	-	✓	✓	✓	
Completeness	CN	PyBoolNet	✓	✓	-	✓	-	✓	✓	✓	
Model checking	MC	PyBoolNet	✓	✓	✓	✓	✓	✓	✓	✓	

Model checking-based method [Cifuentes-Fontanals et al., 2022]²

 $^{^2}$ Cifuentes-Fontanals L, Tonello E and Siebert H (2022) Control in Boolean Networks With Model Checking. Front. Appl. Math. Stat. 8:838546. doi: 10.3389/fams.2022.838546 $\,\,$ $\,\,$

Control features

					Control target				Update		
Method		Tool	Steady state	Trap space attractor	Complex attractor	Subspace	Arbitrary subset	Async	Sync	Gen. async	
Basins	ВА	CABEAN [14]	✓	✓	✓	-	-	✓	-	_	
Stable motifs	SM	StableMotifs [7]	✓	✓	-	-	-	✓	-	-	
Percolation-only	PO	Caspo [21]	✓	✓	-	✓	-	✓	✓	✓	
Trap spaces	TS	PyBoolNet [9]	✓	✓	-	✓	-	✓	✓	✓	
Completeness	CN	PyBoolNet	✓	✓	-	✓	-	✓	✓	✓	
Model checking	MC	PyBoolNet	✓	✓	✓	✓	✓	<	✓	✓	

I hope to develop more control features for nfvs-motifs: complex attractor, sub-space, arbitrary subset. I am thinking about this.

Model checking-based method [Cifuentes-Fontanals et al., 2022]

Network		Size	Inputs	Outputs	Attractors	
					Steady	Cyclic
Cell-Fate	[2]	28	3	3	27	0
MAPK	[3]	53	4	3	12	6
T-LGL	[4]	60	6	3	0	3

Input components are fixed in the T-LGL network and free in the Cell-Fate and MAPK networks, unless specified otherwise.

The running time of the model checking-based method (MC) is large (several hours).

Model checking-based method [Cifuentes-Fontanals et al., 2022]

Network		Size	Inputs	Outputs	Attractors	
					Steady	Cyclic
Cell-Fate	[2]	28	3	3	27	0
MAPK	[3]	53	4	3	12	6
T-LGL	[4]	60	6	3	0	3

Input components are fixed in the T-LGL network and free in the Cell-Fate and MAPK networks, unless specified otherwise.

We can see that MC seems to be not time-efficient for large-scale models.

Model checking-based method [Cifuentes-Fontanals et al., 2022]

Network		Size	Inputs	Outputs	Attractors	
					Steady	Cyclic
Cell-Fate	[2]	28	3	3	27	0
MAPK	[3]	53	4	3	12	6
T-LGL	[4]	60	6	3	0	3

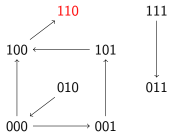
Input components are fixed in the T-LGL network and free in the Cell-Fate and MAPK networks, unless specified otherwise.

It would be great if nfvs-motifs can provide all features as MC does but run much rapidly.

Improve the accuracy

Consider the example asynchronous Boolean network (Figure 2 in [Cifuentes-Fontanals et al., 2021]).

$$\begin{cases} f_1 = \neg x_2 \lor (x_1 \land \neg x_3) \\ f_2 = (x_1 \land \neg x_3) \lor (x_2 \land x_3) \\ f_3 = (\neg x_1 \land \neg x_2) \lor (x_2 \land x_3) \end{cases}$$

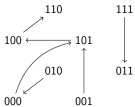


The result of pystablemotifs is $\{x_1 = 1, x_3 = 0\}$. The result of CABEAN (also of PyBoolNet with model checking) is $\{x_3 = 0\}$.

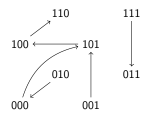
We have obtained a preliminary idea for improving the accuracy of pystablemotifs. However, we need to make it more specific as well as investigate its efficiency. In most cases, smaller control policies, longer computational time. We will present it in the next slides.

Consider the synchronous counterpart of the previous asynchronous Boolean network.

$$\begin{cases} f_1 = \neg x_2 \lor (x_1 \land \neg x_3) \\ f_2 = (x_1 \land \neg x_3) \lor (x_2 \land x_3) \\ f_3 = (\neg x_1 \land \neg x_2) \lor (x_2 \land x_3) \end{cases}$$



$$\begin{cases} f_1 = \neg x_2 \lor (x_1 \land \neg x_3) \\ f_2 = (x_1 \land \neg x_3) \lor (x_2 \land x_3) \\ f_3 = (\neg x_1 \land \neg x_2) \lor (x_2 \land x_3) \end{cases}$$



Assume that the target attractor is $\{110\}$. The result of pystablemotifs is $\{x_1 = 1, x_3 = 0\}$. The minimum result still should be $\{x_3 = 0\}$.

$$\begin{cases} f_1 = \neg x_2 \lor x_1 \\ f_2 = x_1 \end{cases}$$



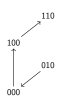
 $\{x_1=1,x_3=0\}$ is a stable motif obtained by pystablemotifs. The function minimal_drivers (in the file drivers.py) finds the minimum subsets of stable motif's states that, when fixed in the logical model, are enough to force the state of every node in the motif into the stable motif M. This function considers all subsets of size 1, 2, ..., |M|-1 and relies on LDOI (implied, contradicted =

logical_domain_of_influence(driver_dict,primes)).

In the example, $\{x_3 = 0\}$ cannot be obtained by using this method.

4□ > 4□ > 4□ > 4□ > 4□ > 4□ >

$$\begin{cases} f_1 = \neg x_2 \lor x_1 \\ f_2 = x_1 \end{cases}$$



I guess if we use DOI instead of LDOI, we can get $\{x_3 = 0\}$ instead of $\{x_1 = 1, x_3 = 0\}$. However, calculating DOI can be difficult in general.

Research question: How to develop an efficient algorithm for driver node identification in a stable motif regardless of update schemes?

The efficient word means that the algorithm is fast and can return smaller (maybe not optimal) sets of drive nodes.

Ideas for driver node identification

From the previous example, we observed that:

- If fixing $x_1 = 1$, the resulting BN has two max. trap spaces $\{x_3 = 0\}$ and $\{x_2 = 1, x_3 = 1\}$. The latter is not compatible with the stable motif $\{x_1 = 1, x_3 = 0\}$.
- If fixing $x_3 = 0$, the resulting BN has only one max. trap space $\{x_1 = 1\}$. All max. trap spaces are compatible with the stable motif $\{x_1 = 1, x_3 = 0\}$.

We formulate a problem as follows.

Driver node identification

Let $\mathcal N$ be a BN and $M=\{x_{i_1}=a_{i_1},...,x_{i_k}=a_{i_k}\}$ be a max. trap space of $\mathcal N$. Find all minimum subsets S of M such that of max. trap spaces of the new BN induced by S are compatible with M.

This problem is related to the fixed point control problem considered in [Biane and Delaplace, 2019, Moon et al., 2022].

Ideas for driver node identification

We can solve this problem by using answer set programming. The main idea is as follows.

- Consider the nodes in M as controllable nodes (x_1, x_3) and the remaining nodes as uncontrollable nodes (x_2) .
- Introduce an auxiliary Boolean variable d_i for each controllable node x_i (d_1 , d_3). If $d_i = 1$, node x_i is fixed to a_i . If $d_i = 0$, node x_i is updated by using its original Boolean function.
- Let \mathcal{L} be the ASP characterizing the trap spaces of \mathcal{N} considering all d_i . Let C be the constraint characterizing the max. trap space M $(x_1 = 1 \land x_3 = 0)$.
- Compute the set A of minimal answer sets (projecting only d_i not x_i) of $\mathcal{L} + C$. $A = \{\{d_1\}, \{d_3\}\}$
- Compute the set A' of minimal answer sets (projecting only d_i not x_i) of $\mathcal{L} + \neg \mathcal{C} + A$. $A' = \{\{d_1\}\}$
- Return $A \setminus A'$. Return $\{\{d_3\}\}$ where $\{d_3\}$ corresponds to $\{x_3 = 0\}$

Ideas for driver node identification

This ASP-based method can avoid considering explicitly all subsets of size 1, 2, ..., |M| - 1. This exploits the power of ASP solvers (e.g., clingo).

However, it is still needed to test its efficiency in practice. Moreover, we also need to investigate its correctness thoroughly.

Bottlenecks of both pystablemotifs and mtsNFVS

The new approach still encounters the two bottlenecks of both pystablemotifs and mtsNFVS.

Bottlenecks of both pystablemotifs and mtsNFVS

The case of many source nodes where there are actually too many attractors (at least 2^k where k is the number of source nodes). mtsNFVS is less affected than pystablemotifs, but this is still problem because mtsNFVS uses the set of explicit (not symbolic) min. trap spaces.

We have found a wise way to symbolically represent the maximal answer sets (corresponding to min. trap spaces) of the encoded ASP characterizing all trap spaces.

For max. trap spaces, we need to adjust the current method. We are thinking about this.

Bottlenecks of both pystablemotifs and mtsNFVS

There exist some very huge attractors inside min. trap spaces. In this case, the number free variables of a min. trap space is still too large (even all the nodes of the original network). pystablemotifs only ends with min. trap spaces. It is very hard for mtsNFVS to reach the best case (i.e., |F|=1) to avoid the reachability analysis.

One example is the model shown in https://github.com/jcrozum/pystablemotifs/issues/83. This network has no stable motifs, and we have exactly one attractor for each combination of source node values.

Our general idea is as follows.

We have the information about the min. trap space. We can use this
information to guide Preprocessing SSF to reach the easy cases where
we can avoid or at least reduce the reachability analysis times.

This is a very rare case. We can anticipate it in advance via the information about the number of free nodes of the min. trap space.

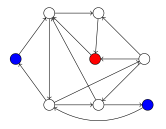
In this case, it is harder for Preprocessing SSF to converge into one state (i.e., the best case where we do not need to check the reachability).

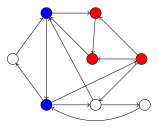
However, we know that it is likely that there is only one huge attractor and all candidate states are in this attractor.

My idea is as follows.

- First, choose a pivot state s^p from the candidate set F_m .
- Second, from s^p create and gradually enlarge a "trap" to attract other states in F_m . The trap is the set of states that can be reach s^p .
- Third, perform Preprocessing SSF and the enlargement in parallel. When a state converges into the trap, we can exclude it from F_m .
- Finally, we have F_m with the expectation that it contains only the pivot state.

The principle is that larger trap, more possibility to converge it.



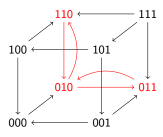


How to efficiently enlarge the trap (i.e., computing predecessors)? Using BDDs as in [Garg et al., 2008, Benes et al., 2021]?

Thanks to time-reversal, we can compute successors (instead of predecessors in the original ABN) in the time-reversal ABN.

This process can purely rely on the function evaluation (not BDDs or unfoldings). Hence, it is expected to be viable for very large models.

$$\begin{cases} v_1, \neg v_1 \land v_2 \land \neg v_3 \\ v_2, \neg v_1 \lor \neg v_3 \\ v_3, \neg v_1 \land v_2 \land \neg v_3 \end{cases}$$

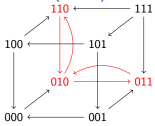


This BN has one min. trap space $m = \{v_1 = \star, v_2 = \star, v_3 = \star\}$. There is one attractor inside m, $\{010, 011, 110\}$.

The time-reversal BN has one max. trap space $\{v_1=1, v_2=\star, v_3=1\}$, which is self-negating. Herein, $\{101, 111\}$ is not in the attractor.

Applying the approach of mtsNFVS with $U_{min}^- = \{v_1, v_3\}$ and $b_1 = 1, b_3 = 1$. Then, $F = \{011, 110, 101\}$. We can exclude 101 from F before any further analysis.

$$\begin{cases} v_1, \neg v_1 \land v_2 \land \neg v_3 \\ v_2, \neg v_1 \lor \neg v_3 \\ v_3, \neg v_1 \land v_2 \land \neg v_3 \end{cases}$$



Theorem

Let m be a trap space of a BN N. Let N_m be the BN induced by m and N' be the time-reversal BN of N_m . If s is an attractor state inside m, then s cannot lie in any self-negating max. trap space of N'.

Proof

Assume s is in a self-negating max. trap space M of N'. Then, all states of the attractor containing s are also in M. The states satisfy M but following the transitions of N, they should reach states not satisfying M because M is a self-negating max. trap space of N'. This is a contradiction.

Reachability analysis

We may still need to check the reachability in some cases, for example, there is a motif-avoidant attractor, the Preprocessing SSF is not good enough.

The reachability analysis is the last resort ensuring the correctness of our approach.

The reachability in ABNs is PSPACE-complete. Its research is far from our focus now. Hence, we should use the existing methods for the reachability analysis in a wise manner relying on the information we have already known.

Reachability analysis (cont.)

For example, in the motif-avoidant attractor checking,

• If |F|=1, it is likely that there is a motif-avoidant attractor, the result of the reachability analysis is likely unreachable. In this case, we should first use a static analyzer (e.g., PINT [Paulevé, 2017]) that is efficient for very large models in the case of unreachable.

For example, in the process of each min. trap space m,

• If $|F_m| = 2$, it is likely that there is only one attractor and the result of the reachability analysis is always reachable. In this case, we should first use a bounded model checking-based approach (e.g., SAT, ASP) that is efficient for very large models in the case of reachable. Goal-driven Petri net unfolding [Chatain and Paulevé, 2016] is also a possible option in this case, but I do not think it can handle very large models.

We need to consider this problem thoroughly.

Implementation

Trappist (in Python): .bnet file \xrightarrow{pyeda} BDDs of Boolean functions \rightarrow Petri net (digraph) \rightarrow ASP (string) \xrightarrow{clingo} min./max. trap spaces (json)

We think that we should start from scratch, i.e., we remove completely the dependency on PyBoolNet because it uses a heavy data structure (i.e, prime-implicants).

We propose to design a completely new system for attractor detection and control. We will gradually adapt the functions from pystablemotifs, mtsNFVS, and Trappist to the new system. Of course, we need a good system design first.

Note that our proposed approach contains many constituent tasks (e.g., computation of min./max. trap spaces, computation of LDOI, driver node identification). The new system should be easy to replace new implementations for each constituent task.

Trappist: new features

Now, Trappist supports computing

- max. trap spaces (original/time-reversal BN)
- min. trap spaces (original/time-reversal BN)
- fixed points (original/time-reversal BN)

Introduction ...

System design

Design a new system for attractor detection and control (input/output formats, components, functions, etc.).

Discuss ...

Computing negative feedback vertex sets

Regarding the NFVS size, the result of AEON is better than that of mtsNFVS is most cases for real-world models. This is opposite for the case of random models.

Regarding the running time, AEON is much faster than mtsNFVS for both real-world and random models.

There is a matter needed to be considered in our whole algorithm. If we only compute the NFVS (for the original ABN) one time, we can try both AEON and mtsNFVS. But, if we compute an NFVS for each intermediate ABN (induced in a maximal trap space or stable motif), we should try only AEON because of its time efficiency.

Of course, we can try all options and see which is the best.

Computing negative feedback vertex sets (cont.)

"I think I have discovered the cause of the check failure. On line 62 of SignedGraph.py, there is the line ..."

Indeed, SignedGraph.py is my own code (not of the FVSpython code). Note that the FVSpython code only computes an FVS, not an NFVS. Hence, I think the FVSpython code was properly converted from python 2 to python 3.

Now, the easiest option is that I will fix my own code so that it is properly compliant with python 3.

Compute the candidate set of states I

I first recall that the candidate set of states is the set of fixed points of the reduced STG with respect to an NFVS and a set of Boolean values assigned to nodes in this NFVS.

In our new algorithm, we restrict it to the terminal restriction space. Specifically, we can ignore states belonging:

- Stable motifs of G: already in form of sub-spaces.
- Self-negating stable motifs of TR(G): already in form of sub-spaces.
- States for which R(X) is zero: needed to convert to a list of sub-spaces.

Actually, we only need to find the DNF of $\neg R(X)$. The computation of prime implicants is not necessary. There are two cases:

Compute the candidate set of states II

- $\neg R(X)$ is represented by a BDD: the union of all paths from the root to the one node. Each path is represented as a sub-space.
- ¬R(X) is represented by an expression (e.g., pyeda expr): each
 conjunction of the DNF is represented as a sub-space. As I know, the
 DNF conversion of pyeda does not require the computation of prime
 implicants.

Samuel: Since R(X) depends on the Boolean update functions, can we encode the avoidance of states in $\neg R(X)$ directly in the encoded ASP (i.e., no need a DNF)?

Giang: It is possible. Let consider for example $\neg R(X) = ((A \lor B) \land \neg C) \lor D$. Then the encoded ASP using the characterization of deadlocks should be:



Compute the candidate set of states III

```
: \neg_{\sqcup} pD.

: \neg_{\sqcup} temp1_{\sqcup}; _{\sqcup} nC.

temp1_{\sqcup} : \neg_{\sqcup} pA.

temp2_{\sqcup} : \neg_{\sqcup} pB.
```

This ASP has the same set of answer sets (maybe not equivalent to) as the ASP using the DNF of $\neg R(X)$ (i.e., $(A \land \neg C) \lor (B \land \neg C) \lor D$) as follows:

```
:-upD.
:-upAu;unC.
:-upBu;unC.
```

Compute the candidate set of states IV

Note that in general obtaining a single DNF may be exponential. The above approach does not required the DNF of $\neg R(X)$, it only relies on the syntax of $\neg R(X)$.

In addition, the compute_fixed_point_reduced_STG function should be included a new parameter induced_sub_space, which ensures that all fixed points are inside a given maximal trap space. Note however that this is needed if we only use the NFVS of the original ABN.

Motif-avoidant checking

"I highlighted a few minor style issues, plus a question about one of the functions in state_utils.py ..."

I assume that for a state, the value of every node is specified. But of course, for the case that we do not specify all the values of the regulators of f, an exception should be thrown.

I will fix it and other minor style issues.

Next steps

I propose to start writing the manuscript in parallel with the implementation. We have four people and I think this strategy will help us to save a lot of time.

<u>Publication plan</u>: is it similar to the case of the stable motifs-based method?

- A main journal paper: do you have any target in mind?
- A tool paper: Application notes in Oxford Bioinformatics?

What do you think?

A new related paper

I have just found a related paper.

Sugyun An, So-Yeong Jang, Sang-Min Park, Chun-Kyung Lee, Hoon-Min Kim, Kwang-Hyun Cho, Global stabilizing control of large-scale biomolecular regulatory networks, Bioinformatics, 2023;, btad045

It is compared to pystablemotifs for the target control.

I have not read the details yet. But for me, again the results reported in this paper seem not to be reliable because:

- "Among the biological Boolean networks in the Cell Collective (Helikar, et al., 2012), we selected 39 networks that have more than two attractors (including at least one point attractor) when all input nodes are fixed with the Off state."
- From Fig 6.C, I found that it omits the inflammatory-bowel model (in the Cell Collective). This model has 47 nodes, and it is hard for PyBoolnet.

Experimentation

We plan to test the new method on models without constant nodes.

N-K models: expected to handle networks with K=2 and up to 500 nodes.

scale-free models: expected to handle networks with K=10 and up to 5000 nodes.

real-world models: expected to handle all the largest and most complex networks that we can find in the literature. The repository ² maintained by Samuel Pastva is a good source.

Conclusion

The new approach exploits advantages of both pystablemotifs and mtsNFVS.

It benefits not only the attractor detection issue but also the control issue of asynchronous Boolean networks.

We believe that it will be the most promising approach that can reach the genome scale.

Collaboration

We hope we will have a nice collaboration on this work. Hence, we should make clear something at the beginning.

Participants:

- Our side: Van-Giang Trinh and Samuel Pastva.
- Your side: Jordan Rozum and Kyu Hyong Park.

Authorship (if we can publish some papers :)):

 Two first authors, one from our side and one from your side with the equal contribution?

What do you think?

Next steps

Task 1: Complete our new approach and make it as detailed as possible.

- Find solutions for some hard problems (e.g., the case of very huge attractors).
- Prepare more specific slides (or a report) presenting technical details of the new approach.
- Add more concrete examples for illustration.

Task 2: Start the implementation of our new approach.

- Design a new system for attractor detection and control (input/output formats, components, functions, etc.).
- Programming

We should create a Github repository first. Everything will be updated there.

Our proposal

I and Samuel will focus on Task 1. Of course, Jordan and Kyu will also involve in this task to make the new approach as powerful as possible.

Jordan and Kyu will lead Task 2 because you already had the holistic view about pystablemotifs. You can divide and assign to all of them implementation sub-tasks. I and Samuel will contribute to this process.

Should keep regular exchanges between us.

Your suggestions?

New preprint for enumerating attractors in asynchronous BNs based on reduction³.

Let \mathcal{A} be an ABN and w be its non-auto-regulated node. Let \mathcal{A}^w be the ABN obtained by removing w from \mathcal{A} and replacing w by its Boolean function in all Boolean functions of its inputs. Then if att is an attractor of \mathcal{A} , then there exists at least one attractor for \mathcal{A}^w in $\pi^w(att)$ and for each $x \in \pi^w(att)$ contained in an attractor of \mathcal{A}^w , $\gamma^w(x) \in att$.

$$\pi^w \colon \{0,1\}^n \mapsto \{0,1\}^{n-1}$$
 is the projection on $V \setminus \{w\}$.

$$\gamma^w : \{0,1\}^{n-1} \mapsto \{0,1\}^n \text{ is the mapping where } \gamma^w(x^w) = (x^w, f_w(x^w)).$$

Can be used to efficiently enumerating all attractors of the original ABN.

Some issues:

- \mathcal{A}^w may have motif-avoidant attractors, whereas \mathcal{A} have no ones. See the case of h in Figure 1 of the paper.
- \mathcal{A}^w may have non-univocal attractors, whereas \mathcal{A} may have no ones. See the case of g in Figure 1 of the paper.
- \mathcal{A}^w has less nodes than \mathcal{A} but its Boolean functions may be much more complex.
 - ▶ The minimum NFVS of A^w may be larger than that of A.
 - Reduce the efficiency of trappist because the density of \mathcal{A}^w may be much larger than that of \mathcal{A} .
 - ► For AEON, if there are many dependencies, the BDDs will also have many internal dependencies which increase size.

The new method is very efficient for real-world models, but not for N-K models. Why?

- By applying the mediator-node reduction to the set of real-world models, we found that the reduced models are very easy to analyze and their number of attractors is equal to that of the original models.
- In addition, real-world models are quite sparse, leading to the Boolean functions of the reduced models are not very complex.
- This is not the case for N-K models.

What we can explore in this direction?

- Of course, we can get a much smaller candidate set for nfvs-motifs by computing attractors of the reduced model. If lucky, we can avoid some bottlenecks of nfvs-motifs, for example, a minimal trap space m has too many free variables.
- Try to handle the problems of introducing motif-avoidant and non-univocal attractors to the reduced model.
- Heuristics for stop conditions and the order of removed nodes (based on not only the number of nodes but also the function complexity). I guess it might be related to FVS or NFVS.
- Investigate how does the reduction change the structure of the succession diagram. For example, I guess \mathcal{A}^w might have a deeper SD than \mathcal{A} has.
- In case we must check reachability, how the reduction can help the reachability analysis more efficient?
- Investigate how the reduction can help the control algorithms.
- Theoretical findings



Conjecture

Let $\mathcal A$ be an ABN and w be its mediator node. Let $\mathcal A^w$ be the ABN obtained by removing w from $\mathcal A$ and replacing w by its Boolean function in all Boolean functions of its inputs. Then the set of attractors of $\mathcal A^w$ one-to-one corresponds to that of $\mathcal A$.

w is a mediator node if $in(w) \cap out(w) \neq \emptyset$.

The case $in(w) \cap out(w) \neq \emptyset \wedge |in(w)| = |out(w)| = 1$ has been proven in³

Not sure whether the conjecture has been proven?

³Saadatpour, A., Albert, R.,& Reluga, T. C. (2013). A reduction method for Boolean network models proven to conserve attractors. SIAM Journal on Applied Dynamical Systems, 12(4), 1997-2011.

Conjecture

Let \mathcal{A} be an ABN and w be its non-auto-regulated node. Let \mathcal{A}^w be the ABN obtained by removing w from \mathcal{A} and replacing w by its Boolean function in all Boolean functions of its inputs. Then if m is a minimal trap space of \mathcal{A} , then $\pi^w(m)$ is also a minimal trap space of \mathcal{A}^w .

More generally, we can investigate the relations regarding trap spaces (minimal/maximal) between the reduced model and the original one.

I and Sam have discussed about this new direction. We also have some promising preliminary ideas.

We propose us to consider this direction after finishing our first paper.

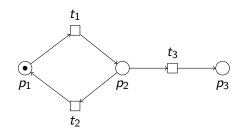
What do you think?

Thank you for your attention!

Petri nets

Bipartite graph

Places PTransitions TWeighted arcs W



A <u>marking</u> 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. For example, p_1 is marked, p_2 and p_3 are unmarked.

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. For example, $pred(p_2) = \{t_1\}$, $succ(p_2) = \{t_2, t_3\}$, $pred(t_2) = \{p_2\}$, and $succ(t_2) = \{p_1\}$.

Siphons

Siphon

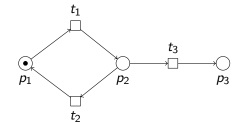
A siphon of a Petri net (P, T, W) is a set of places S such that:

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

Once a siphon is unmarked, it remains unmarked.

$$S = \{p_1, p_3\}$$
 is not a siphon because $S \cap succ(t_3) = \{p_1, p_3\} \cap \{p_3\} = \{p_3\} \neq \emptyset$ but $S \cap pred(t_3) = \{p_1, p_3\} \cap \{p_2\} = \emptyset$.

Here: \emptyset , $\{p_1, p_2\}$, $\{p_1, p_2, p_3\}$



Traps

Trap

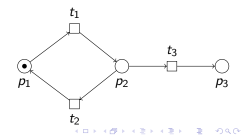
A <u>trap</u> of a Petri net (P, T, W) is a set of places S such that:

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

Once a trap is marked (i.e., at least one of its places is marked), it remains marked.

$$S = \{p_1, p_3\}$$
 is not a trap because $S \cap pred(t_1) = \{p_1, p_3\} \cap \{p_1\} = \{p_1\} \neq \emptyset$ but $S \cap succ(t_1) = \{p_1, p_3\} \cap \{p_2\} = \emptyset$.

Here: \emptyset , $\{p_1, p_2, p_3\}$



Petri net of a Boolean model

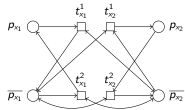
The original encoding was established in [Chaouiya et al., 2004].

Two places for each gene: $v \rightsquigarrow p_v, \overline{p_v}$

Solutions of $f_{\nu} \not\leftrightarrow \nu \rightsquigarrow$ transitions from p_{ν} to $\overline{p_{\nu}}$ (and back)

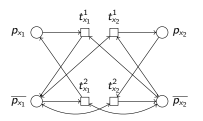
At any marking m of the Petri net encoding a Boolean model, it always holds that $m(p_v) + m(\overline{p}_v) = 1$.

$$\begin{cases} f_1 = (x_1 \wedge x_2) \vee (\neg x_1 \wedge \neg x_2) \\ f_2 = (x_1 \wedge x_2) \vee (\neg x_1 \wedge \neg x_2) \end{cases}$$



Conflict-free siphons/traps

A siphon/trap is called **conflict-free** if it does not contain both p_v and $\overline{p_v}$ for all $v \in V$.



Siphon	Conflict-free?
Ø	yes
$\{\overline{p_{x_1}},\overline{p_{x_2}}\}$	yes
$\{p_{x_1},\overline{p_{x_1}}\}$	no
$\{p_{x_2},\overline{p_{x_2}}\}$	no

A conflict-free siphon (resp. trap) is $\underline{\text{maximal}}$ if it is not a subset of any other conflict-free siphon (resp. trap).

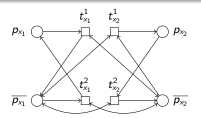
Conflict-free siphons are trap spaces

Theorem 1

Let \mathcal{M} be a Boolean model and \mathcal{P} be its Petri net encoding. There is a one-to-one correspondence between the set of **trap spaces** of \mathcal{M} and the set of **conflict-free siphons** of \mathcal{P} .

$$\begin{cases} f_1 = (x_1 \wedge x_2) \vee (\neg x_1 \wedge \neg x_2) \\ f_2 = (x_1 \wedge x_2) \vee (\neg x_1 \wedge \neg x_2) \end{cases}$$

Trap space	Conflict-free siphon
**	Ø
11	$\{\overline{p_{x_1}},\overline{p_{x_2}}\}$



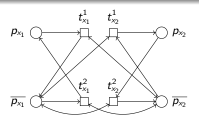
Maximal conflict-free siphons are minimal trap spaces

Theorem 2

Let $\mathcal M$ be a Boolean model and $\mathcal P$ be its Petri net encoding. There is a one-to-one correspondence between the set of **minimal trap spaces** of $\mathcal M$ and the set of **maximal conflict-free siphons** of $\mathcal P$.

$$\begin{cases} f_1 = (x_1 \wedge x_2) \vee (\neg x_1 \wedge \neg x_2) \\ f_2 = (x_1 \wedge x_2) \vee (\neg x_1 \wedge \neg x_2) \end{cases}$$

Trap space	Conflict-free siphon
**	Ø
11	$\{\overline{p_{x_1}},\overline{p_{x_2}}\}$



Proposed method for minimal trap space computation

From Theorem 2, we propose a new method for computing minimal trap spaces of a Boolean model \mathcal{M} .

- Build the Petri net encoding \mathcal{P} of \mathcal{M} .
- Compute all maximal conflict-free siphons of P.
- Convert the obtained maximal conflict-free siphons into the corresponding minimal trap spaces.

Petri net transformation

Transforming a Boolean model into its Petri net encoding can be done via computing Disjunctive Normal Forms (DNF) of each Boolean function [Chatain et al., 2014]. This transformation is implemented in the bioLQM³ library using BDDs.

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.

We use the above transformation in our proposed method.

³http://www.colomoto.org/biolqm/

Maximal conflict-free siphon computation

Characterize all siphons of the encoded Petri net as a system of Boolean rules.

$$p \in S \Rightarrow \bigvee_{p' \in pred(t)} p' \in S, p \in P, t \in T, t \in pred(p)$$

Add to the system the Boolean rules representing the conflict-freeness.

$$p_v \in S \Rightarrow \overline{p_v} \not \in S \land \overline{p_v} \in S \Rightarrow p_v \not \in S, v \in V$$

Encode the system as an ASP.

Use an ASP solver (e.g., clingo [Gebser et al., 2011]) to compute all set-inclusion maximal answer sets of the ASP.

Set-maximality through "heuristics" clingo --heuristic=Domain --enum-mod=domRec --dom-mod=3

References I

Akutsu, T. (2018).

Algorithms for analysis, inference, and control of Boolean networks. World Scientific, Singapore.

Benes, N., Brim, L., Pastva, S., and Safránek, D. (2021). Computing bottom SCCs symbolically using transition guided reduction.

In <u>International Conference on Computer Aided Verification</u>, pages 505–528. Springer.

Biane, C. and Delaplace, F. (2019).

Causal reasoning on Boolean control networks based on abduction:

Theory and application to cancer drug discovery.

IEEE/ACM Transactions on Computational Biology and Bioinformatics, 16(5):1574–1585.

References II

- Brim, L., Pastva, S., Šafránek, D., and Šmijáková, E. (2021). Parallel one-step control of parametrised boolean networks. Mathematics, 9(5):560.
 - Chaouiya, C., Remy, E., Ruet, P., and Thieffry, D. (2004).

 Qualitative modelling of genetic networks: From logical regulatory graphs to standard Petri nets.
 - In Cortadella, J. and Reisig, W., editors, <u>Applications and Theory of Petri Nets 2004</u>, 25th International Conference, ICATPN 2004, <u>Bologna</u>, Italy, June 21-25, 2004, <u>Proceedings</u>, volume 3099 of <u>Lecture Notes in Computer Science</u>, pages 137–156. Springer.

References III

Chatain, T., Haar, S., Jezequel, L., Paulevé, L., and Schwoon, S. (2014).

Characterization of reachable attractors using Petri net unfoldings. In Mendes, P., Dada, J. O., and Smallbone, K., editors, Computational Methods in Systems Biology - 12th International Conference, CMSB 2014, Manchester, UK, November 17-19, 2014, Proceedings, volume 8859 of Lecture Notes in Computer Science, pages 129–142. Springer.

- Chatain, T. and Paulevé, L. (2016). Goal-driven unfolding of Petri nets.
- Cifuentes-Fontanals, L., Tonello, E., and Siebert, H. (2021). Control in Boolean networks with model checking. arXiv preprint arXiv:2112.10477.

References IV

Cifuentes-Fontanals, L., Tonello, E., and Siebert, H. (2022). Control in Boolean networks with model checking. Frontiers in Applied Mathematics and Statistics, 8.

Fontanals, L. C., Tonello, E., and Siebert, H. (2020).

Control strategy identification via trap spaces in Boolean networks.

In Abate, A., Petrov, T., and Wolf, V., editors, Computational

Methods in Systems Biology - 18th International Conference, CMSB

2020, Konstanz, Germany, September 23-25, 2020, Proceedings, volume 12314 of Lecture Notes in Computer Science, pages 159–175.

Springer.

References V

Garg, A., Cara, A. D., Xenarios, I., Mendoza, L., and Micheli, G. D. (2008).

Synchronous versus asynchronous modeling of gene regulatory networks.

Bioinform., 24(17):1917-1925.

Gebser, M., Kaufmann, B., Kaminski, R., Ostrowski, M., Schaub, T., and Schneider, M. (2011).

Potassco: The Potsdam answer set solving collection.

Al Commun., 24(2):107-124.

Giang, T. V. and Hiraishi, K. (2021).

An improved method for finding attractors of large-scale asynchronous Boolean networks.

In 2021 IEEE Conference on Computational Intelligence in Bioinformatics and Computational Biology (CIBCB), pages 1–9. IEEE.

References VI

Klarner, H., Streck, A., and Siebert, H. (2017).

PyBoolNet: a python package for the generation

PyBoolNet: a python package for the generation, analysis and visualization of Boolean networks.

Bioinform., 33(5):770–772.

Laura, C. F., Tonello, E., and Siebert, H. (2022).

Computing trap space-based control strategies for Boolean networks using answer set programming.

In INTERNATIONAL CONFERENCE OF COMPUTATIONAL METHODS IN SCIENCES AND ENGINEERING ICCMSE 2021. AIP Publishing.

Mizera, A., Pang, J., Qu, H., and Yuan, Q. (2019). Taming asynchrony for attractor detection in large Boolean networks. IEEE ACM Trans. Comput. Biol. Bioinform., 16(1):31–42.

References VII

- Moon, K., Lee, K., Chopra, S., and Kwon, S. (2022).

 Bilevel integer programming on a Boolean network for discovering critical genetic alterations in cancer development and therapy.

 European Journal of Operational Research, 300(2):743–754.
- Naldi, A., Monteiro, P. T., and Chaouiya, C. (2012). Efficient handling of large signalling-regulatory networks by focusing on their core control.
 - In Gilbert, D. R. and Heiner, M., editors, Computational Methods in Systems Biology 10th International Conference, CMSB 2012, London, UK, October 3-5, 2012. Proceedings, volume 7605 of Lecture Notes in Computer Science, pages 288–306. Springer.

References VIII



Paulevé, L. (2017).

Pint: A Static Analyzer for Transient Dynamics of Qualitative Networks with IPython Interface.

ln

CMSB 2017 - 15th conference on Computational Methods for Systems E volume 10545 of Lecture Notes in Computer Science, pages 370 – 316. Springer.



Rozum, J. C., Deritei, D., Park, K. H., Gómez Tejeda Zañudo, J., and Albert, R. (2021a).

pystablemotifs: Python library for attractor identification and control in Boolean networks.

Bioinform.

References IX



Rozum, J. C., Zañudo, J. G. T., Gan, X., Deritei, D., and Albert, R. (2021b).

Parity and time reversal elucidate both decision-making in empirical models and attractor scaling in critical Boolean networks. Sci. Adv., 7(29).



Saadatpour, A., Albert, R., and Reluga, T. C. (2013).

A reduction method for Boolean network models proven to conserve attractors.

SIAM J. Appl. Dyn. Syst., 12(4):1997-2011.



Schwab, J. D., Kühlwein, S. D., Ikonomi, N., Kühl, M., and Kestler, H. A. (2020).

Concepts in Boolean network modeling: What do they all mean? Comput. Struct. Biotechnol. J., 18:571–582.



References X



Su, C. and Pang, J. (2021a).

Cabean 2.0: Efficient and efficacious control of asynchronous Boolean networks

In International Symposium on Formal Methods, pages 581–598. Springer.



Su, C. and Pang, J. (2021b).

CABEAN: a software for the control of asynchronous boolean networks.

Bioinform., 37(6):879-881.



Trinh, G. V., Akutsu, T., and Hiraishi, K. (2020).

An FVS-based approach to attractor detection in asynchronous random Boolean networks.

IEEE/ACM Trans. Comput. Biol. Bioinf. in press.



References XI



In Petre, I. and Paun, A., editors, Computational Methods in Systems Biology - 20th International Conference, CMSB 2022, Bucharest, Romania, September 14-16, 2022, Proceedings, volume 13447 of Lecture Notes in Computer Science, pages 158–176. Springer.

Trinh, V., Hiraishi, K., and Benhamou, B. (2022b).

Computing attractors of large-scale asynchronous boolean networks using minimal trap spaces.

In BCB '22: 13th ACM International Conference on Bioinformatics, Computational Biology and Health Informatics, Northbrook, Illinois, USA, August 7 - 10, 2022, pages 13:1–13:10. ACM.

References XII



Zañudo, J. G. T. and Albert, R. (2015). Cell fate reprogramming by control of intracellular network dynamics. PLOS Computational Biology, 11(4):e1004193.



Zheng, D., Yang, G., Li, X., Wang, Z., Liu, F., and He, L. (2013). An efficient algorithm for computing attractors of synchronous and asynchronous Boolean networks. PloS One, 8(4):e60593.