Simulating heterogeneous populations using Boolean models

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Abstract

Certain biological processes such as the development of cancer are controlled by events that are very rare at the single-cell level, and which are therefore difficult to capture computationally through simulations of individual cells. Here we demonstrate a method for exactly simulating the dynamics of large populations directly, without having to model the individuals in the population. Our method can thus resolve even the rarest outcomes of the model. The only restriction is that the underlying model use only Boolean variables. The populations we consider are composed of non-interacting individuals that span an arbitrary mixture of states, but are described by the same underlying model, although by adding ‘mutation variables’ one can also simulate heterogeneity in model rules. Boolean models are thus ideal for exploring the dynamics of highly heterogeneous populations in which very rare subpopulations may play important roles.

Computer models are widely used to predict typical behaviors of biological systems, generally by simulating a number of instances of that model and enumerating the range of observed outcomes. In the most common random-sampling procedure called Monte Carlo [1], many individual instances of that model are initialized to different random starting states (determined by some predefined distribution) and then evolved over discrete time steps by applying the rules of the model. These simulations can be run very efficiently, allowing the evaluation of complex models, including whole-cell simulations [2]. By design, the basic random sampling procedure captures typical outcomes of these simulations, and only rarely finds unusual occurrences. Yet some biological processes are determined by the rare outliers [3]. For example, tumor initiation, progression, and drug resistance require either rare alterations to arise or selection of low frequency, pre-existing populations to clonally expand and eventually dominate the population in the long run [4, 6]. In situations where something is known about the circumstances leading to a rare outcome, one can bias Monte Carlo to oversample that outcome and then correct for the biased sampling (a strategy known as importance sampling [7]): however in practice the bias is usually limited to simple parameters such as a mutation rate, which can help narrow the search but may be insufficient to find an outcome that results only from very particular circumstances. Furthermore, oversampling can introduce sampling biases: for example, simulations using an unrealistically high mutation rate may discover extraordinary outcomes by virtue of having huge mutational loads, but such a mutational profile may be neither biologically-realistic nor evolutionarily-accessible.

The shortcomings of Monte Carlo have led to interest in exact, analytical solutions to biological models that are not limited by finite sampling; such exact results require very simple models. In particular, significant advances have been made in analyzing Boolean networks [8–16], a class of models built entirely from ON/OFF variables. With these models, the focus has been on extracting the possible long-term outcomes, or attractors. An attractor may be a stable state (steady state) or else a repeating pattern of states (limit cycle). Attractors have been found using network-reduction algorithms that find simple networks encoding the long-term behavior of more complex networks [9, 11, 17], methods that solve steady states as zeros of a polynomial equation [18], SAT methods [13, 14, 19], and binary decision diagrams [15, 16, 20]. See the introduction of Ref. [10] for a review of these techniques.

In this paper we show that Boolean networks can be used in another type of exact analysis: namely, direct simulation of population statistics, without having to simulate the individuals in that population. The simulations...
are exact, so they capture every subpopulation of the model and every event that occurs, no matter how rare. The output of our method is a time series of the frequency with which the characteristic(s) of interest appear in the population. The feasibility of building these simulations depends on the size and topology of the network (for complex or highly-recurrent networks it can be an exponential problem), though we can often simplify the difficult cases using an equation-reduction method that ignores the first few time steps of the simulation. In the limit where we focus only on the long-term behavior, the output describes the attractor dynamics of the characteristics of interest, thus relating our method to existing Boolean analyses.

Our method applies to deterministic [], probabilistic [], and continuous-time Boolean networks. Here, we only analyze synchronous Boolean network models (i.e. networks whose variables all update together in discrete time steps), but this is without loss of generality because an asynchronous network can be modeled as a probabilistic network with synchronous time steps [].

One major benefit to our method is its simplicity, as it follows from only two rules: 1) work in a linear basis whose variables are products of the Boolean state variables, and 2) ignore quickly-decaying modes if we are looking at late-time behavior. Because of rule 1, we refer to our approach as the product-basis method.

Results

Validation We tested the product-basis method using $10^4$ randomly generated 10-node deterministic networks, where each node’s input combined the output of 1-4 randomly-selected nodes using randomly-generated logic rules. First, we ran 100 Monte Carlo individual simulations of each network, using a random ensemble of initial states (equal probability for all states). Next, we used our product-basis method to generate exact time-evolution equations using a random amount of equation-reduction (which determines the earliest time point at which the simulation is valid). We then compared the predictions of our exact simulations to the population dynamics as predicted by Monte Carlo. In each case the product-basis simulation exactly reproduced the average of the Monte Carlo simulations. These networks were small, but still provided a stringent test of the numerics, as they could theoretically involve linear systems of equations up to size ($2^{10})^2$. During this test, our program also generated additional networks whose numerical error overstepped a threshold and were thus omitted from the Monte Carlo comparison; overall 18.5% of tests generated were discarded.

Next, we tested probabilistic networks (PBNs). Our main test consisted of full-time simulations (i.e. without equation-reduction) as those had lower numerical error and allowed for a tighter Monte Carlo comparison. In order to generate realistic PBNs, we augmented the original time-evolution functions $f_i^0$ with random rate parameters $r_i$, leading to the equations $f_i \leftarrow (1-r_i)x_i + r_i f_i^0$ for $0 \leq r_i \leq 1$. We generated $10^4$ PBNs using this technique, calculated the time-evolution equations for each, and again found that our method’s equations reproduced the results of Monte Carlo simulations to within statistical error. Finally, we tested $2.5 \times 10^3$ continuous-time networks generated using the same rule (which are slower to check by Monte Carlo), and again found complete agreement with Monte Carlo.

Demonstration: T-cell network To demonstrate the product-basis method, we used the T-cell activation network described in Ref. [] (see Figure 10 and Table 2 of that paper). The T-cell network is a deterministic, 40-node network with fifty-four edges with multiple feedback loops, and whose attractors include both steady states and limit cycles. For demonstration purposes, we show a traditional Monte Carlo simulation of an individual T-cell network in Figure 1, obtained by choosing an initial Boolean state and straightforwardly applying the model rules over each successive time step. This particular simulation shows transient (nonrepeating) behavior for the first 10 time steps, leading to a limit cycle with a repeating period of 6 time steps.

Next, we performed a population-level simulation using our product-basis method. We first provided a target set of variables to follow in time, which the product-basis algorithm used to generate a set of time-evolution equations involving those variables (along with other variables that were added automatically to close the system of equations). We chose to track three variables: cyclic-AMP response element (CRE) mediated gene activation, the AP1 (Activating protein 1) transcription factor, and their co-occurrence which we label $AP1 \times CRE$. The co-occurrence variable was not redundant: although the CRE and AP1 states are anticorrelated, 0.5 if they are perfect correlated, or any other value on the interval [0, 0.5].

We first generated the product-basis time-evolution equations for our three variables of interest. Next, we set the model variables to an initial state representing a mixed population, and used the time-evolution equations to track the population-level average of each of the three variables for 50 time steps (Figure 1B). We stress that this is an exact result, with no sampling error. The start-
Figure 1. A) Time evolution of one instance of the T-cell network [24], starting from a random initial state. White/black rectangles signify OFF/ON Boolean states. B) Time evolution of the population fraction having activated CRE elements and/or expressing the transcription factor AP1 in a heterogeneous population of T-cell networks, computed using a product-basis calculation. The heterogeneous population begins at \( t = 0 \) as a uniform mixture of all possible \( 2^{33} \approx 10^{10} \) initial states of the upstream portion of the model. C) The effect of a \( 10^{-4} \) knock-out mutation rate per gene in the heterogeneous population. Monte Carlo, but not the product-basis calculation, required this high rate of mutations in order to detect persistent coactivation of CRE and AP1. D) The co-occurrence of CRE activation and AP1 expression in mutated networks shown on a log-10 scale (dotted red line), compared with the amount of this coexpression coincident with mutated cCbl (purple dots). The mutated fraction was computed by subtracting the time series of CRE and AP1 and WT-cCbl from the time series of CRE and AP1.

The co-occurrence of CRE activation and AP1 expression in mutated networks shown on a log-10 scale (dotted red line), compared with the amount of this coexpression coincident with mutated cCbl (purple dots). The mutated fraction was computed by subtracting the time series of CRE and AP1 and WT-cCbl from the time series of CRE and AP1.

Next we demonstrated the ability of the product-basis method to analyze mutations in the network by including the full set of possible gene knock-outs in the T-cell activation network. We did this by adding a set of ‘wild-type’ variables to the network, one for each original variable in the system, and included the wild-type variables in the update rules using an AND operation. For example, an update rule reading \([A \leftarrow B \text{ OR } C]\) became \([A \leftarrow (B \text{ OR } C) \text{ AND } A^{WT}]\). Note that the presence of mutations effectively doubles the size of the network and thus vastly increases the heterogeneity, which is determined by both the number of activation states of the original
variables and the number of mutational profiles, in total spanning the order of $10^{30}$ different subpopulations. Enumeration over the initial states is impossible at this level of diversity, and the traditional recourse is a sparse sampling method such as Monte Carlo, although that loses the ability to resolve very rare subpopulations. Despite this heterogeneity, our product-basis method was able to produce the exact time-evolution equations for CRE and AP1. We chose an initial population that contained each possible combination of knock-out mutations at a 0.01% mutation rate per variable (roughly the highest possible rate of homozygous knockouts given a 1%-per-gene-locus mutation rate [25]) superimposed upon a uniform mixture of each activation state (adjusted so that non-wild-type genes always began OFF). From this initial state, we again followed the exact time course of the CRE and AP1 population fraction, and compared it to our original wild-type result (Figure 1C). Notably, a small fraction of the population reached a steady state showing both CRE activation and AP1 expression. We also validated the result (to within statistical error) using Monte Carlo, although as shown in the figure, Monte Carlo was only useful for comparing the transient behavior, not the rare persistent subpopulation which fell far below its sampling resolution.

Finally, we used our method to infer the mutations leading to CRE and AP1 coexpression. We hypothesized that this was due to knockout of the cCbl gene in the recurrent core of this network, and tested this hypothesis by generating the time course of the three-way co-occurrence of CRE and AP1 and WT-cCbl, where WT denotes the respective wild-type variable. This final time series dropped to zero at steady state, indicating that mutations in this variable are necessary for persistent CRE and AP1 coexpression (see Fig. 1D).

Our results from the T-cell network demonstrate several important aspects of our method. First, we are able to simulate extremely heterogeneous populations, involving far more subpopulations than could be analyzed individually. Second, although our method only deals with heterogeneity in the states of the Boolean variables, we can still simulate a genetically-heterogeneous population by augmenting the Boolean network with mutation variables. Third, we can exactly model subpopulations that are present at very low levels, which are difficult to resolve by random sampling (see the error bars in Figure 1B). For example, the contribution of each triple-mutant was factored in even though a given triple-mutant was present in only $10^{-12}$ of the population. While one might artificially raise the Monte Carlo mutation rate to oversample the mutations [7], this has the disadvantage of over-weighting the effect of multiple mutants, even though realistic evolutionary paths take one or very few mutational steps at a time [26]. In contrast, our exact result is dominated by the evolutionarily-accessible subpopulations that are closest to wild-type.

The code used to generate these results is named tCellActivationEx.m, and is available for download at https://github.com/CostelloLab/ProductBasis. The equation-generating process for Figures 1B and 1C took ~3 and ~300 seconds respectively using our code (written in MATLAB R2015b 8.6.0.267246, running on a 2.6GHZ Intel core i7 Mac with OS 10.9.5). The Monte Carlo comparison in Figure 1C ($n_{\text{runs}} = 10^4$) took ~140 seconds. Note that Monte Carlo error is proportional to $1/\sqrt{n_{\text{runs}}}$.

**Methods**

**Simulations of mixed populations**

The principle behind our method is to write the time evolution of each variable in our model using a linear equation. Doing so guarantees that the dynamical equations we derive for a single cell also fully describe the dynamics of a mixed population of cells, owing to the superposition property of linear systems.

The key to writing linear equations is to introduce a new variable to represent each nonlinear term in a naive update rule, acknowledging that we will have to solve for the dynamics of each new variable as well. In our case, each nonlinear term is a product of Boolean variables, so the update rule for its respective introduced variable will be a product of the constituent Boolean update rules. We demonstrate this procedure using Example 1.

![Figure 2](https://example.com/figure2.png)

**Figure 2.** The 3-Boolean network used in Example 1. Arrows indicate how each variable updates based on the values of its inputs at the previous time step. For example if either A or B is ON at time $t$ then C will be ON at time $t+1$; otherwise C will be OFF.

**Example 1: building equations**

Suppose we want to track the time evolution of vari-
able $A$ in the network shown in Figure 2. Since this network evolves by discrete time steps, we write $x_A(t+1) = f_A(x_B)$ where $f_A$ performs the NOT operation. A linear equation implementing the NOT gate is:

$$f_A(x_B) = 1 - x_B. \quad (1.1)$$

Evidently, in order to follow $x_A$ over all time we must also track the state of its input variable $B$ over time. $B$ implements an AND gate which is a nonlinear operation: $f_B = x_A \cdot x_C$. To make the equation linear, we introduce $x_{AC} = x_A \cdot x_C$, which is 1 if and only if both $A$ and $C$ are ON, and write $f_B$ in terms of this new variable.

$$f_B = x_{AC} \quad (1.2)$$

We still need to calculate $f_{AC}$ for our new variable $x_{AC}$, which is simply the product $f_A \cdot f_C$. (Proof: $f_{AC} = x_{AC}(t+1) = x_A(t+1) \cdot x_C(t+1) = f_A \cdot f_C$). $f_C$ implements an OR gate whose linear equation involves yet another product variable $x_{AB}$.

$$f_{AC} = f_A \cdot f_C$$

$$= (1 - x_B) \cdot (x_A + x_B - x_{AB})$$

$$= x_A - x_{AB} \quad (1.3)$$

The formula for $f_{AC}$ made use of the fact that any Boolean value squared equals itself: for example $x_B \cdot x_B = x_B$ and $x_B \cdot x_{AB} = x_B \cdot x_A \cdot x_B = x_{AB}$.

The process of replacing product terms with new variables, and then solving for the time evolution of those new variables, continues until the equations form a closed system: each variable’s time evolution is in terms of other variables in our system.

$$f_{AB} = (1 - x_B) \cdot (x_{AC})$$

$$= x_{AC} - x_{ABC} \quad (1.4)$$

$$f_{ABC} = (1 - x_B) \cdot (x_{AC}) \cdot (x_A + x_B - x_{AB})$$

$$= x_{AC} - x_{ABC} \quad (1.5)$$

This gives us a closed linear system. To avoid the constant term we can rewrite Eq. $f_A = x_{\emptyset} - x_B$, where $x_{\emptyset} = 1$ updates according to:

$$f_{\emptyset} = x_{\emptyset}. \quad (1.6)$$

Equations $1.1, 1.6$ together with an initial state in $(x_A, x_B, x_{AC}, x_{AB}, x_{ABC})$ describe the time evolution of these quantities in a single Boolean network as a sequence of 0s and 1s in each variable. The final step is to reinterpret these equations as describing the dynamics of a mixed population of networks, formally by taking the mean of both sides of each equation.

$$\langle f_A \rangle = 1 - \langle x_B \rangle \quad (1.1)$$

$$\langle f_B \rangle = \langle x_{AC} \rangle \quad (1.2)$$

$$\langle f_{AC} \rangle = \langle x_A \rangle - \langle x_{AB} \rangle \quad (1.3)$$

$$\langle f_{AB} \rangle = \langle x_{AC} \rangle - \langle x_{ABC} \rangle \quad (1.4)$$

$$\langle f_{ABC} \rangle = \langle x_{AC} \rangle - \langle x_{ABC} \rangle \quad (1.5)$$

Here angle-brackets denote an average, and we have used the notation $\langle f_i \rangle = \langle x_i(t+1) \rangle = f_i$. The linear equations are unaffected by the averaging process, showing that the same equations used to derive the dynamics of a single instance of a model also describe the mean values of those same variables in a mixed population. The only difference is that the equations were derived using binary variables, whereas the population-averaged variables are real-valued on the interval $[0,1]$: for example we would set $\langle x_A \rangle = 0.4$ if 40% of the population has gene $A$ set ON.

Any state or mixture of states can be written as a linear combination of product-basis variables \{x\}, because these variables form a complete basis spanning the state space (see Appendix 1 for a proof; also Ref. \[18\] proves a similar result for a slightly different Boolean algebra). Since each time-evolution function $f$ is a sum over all states causing a ‘1’ in the output variable when written in the state basis, it follows that each $f$ is also a linear combination of our $x$ variables. Therefore our procedure for modeling a mixed population always works in principle, even if some networks require too many equations for this method to be practical.

We can extend mixed-population modeling to probabilistic \[21\] and continuous-time \[22\] Boolean networks. Probabilistic Boolean networks require no changes in the algorithm; the only difference is that the polynomial coefficients in our equations may not be integers. For example, if the NOT gate in Fig. 2 is leaky with $A$ turning ON with a probability of 0.9/0.4 depending on whether $B$ was OFF/ON at the last time step, then the transition rule for gene $A$ becomes $f_A = 0.9 - 0.5x_B$. As before, a linear equation can always be written, because a) a linear equation can still be written in the state space basis, and b) our $x$ variables are just a different basis covering the
state space (see Appendix 1). Probabilistic networks give one way to incorporate rate information into our model; another way is to work in continuous time using differential equations: $f_A = dx_A/dt$. The differential form does require one change in our method: the rate of change of a higher-order variable is found by using the product rule of derivatives. Whereas under a discrete update $f_{ABC}$ is the product $A \cdot B \cdot C \cdot \ldots$, for the differential case we compute:

$$
\frac{df_{ABC}}{dt} = \frac{d}{dt}(x_A x_B x_C \ldots)
= x_A x_B \ldots f_C + x_A x_C \ldots f_B + \ldots
$$

(1.7)

Also, under discrete updates the trivial function is $f_\emptyset = 1$ but with differential updates it is $f_\emptyset = 0$.

**Long-term behaviors**

A mixed-population simulation may or may not be practical, depending on whether the system of linear equations closes with a manageable number of variables $n$. In the worst case, the entire $2^n$-sized variable space could be involved. By counting subscripts we know that there are $n = 2^N$ product variables associated with an $N$-Boolean network, which is expected because our $x$-variables are simply a change of basis from the state space (see Appendix 1). Therefore the problem has potentially exponential complexity.

One way to make progress when the full closed system of equations is unmanageable is to focus on the attractors (steady states or limit cycles). The attractors are governed by a linear space whose size is determined by the number of attractor states, which for biological networks is usually much smaller than the full equation space. Mathematically, this means that our linear equations form a very degenerate system: if there are only $n^*$ attractor states, then there are only $n^*$ non-zero eigenvalues and $n^*$ linearly independent equations. So for a 50-node network with a single steady state attractor we might have $n = 2^{50} \approx 10^{15}$ equations in the worst case, but $n^* = 1$, which is a vastly smaller linear system. To only find the structure of the final-state space we select $n^*$ linearly independent variables, substitute them for the other variables in the time-evolution functions, and do an eigenvalue analysis on the much-smaller $n^* \times n^*$ system. A continuation of Example 1 gives a simple demonstration of this procedure.

**Example 1, continued**

Eqs. [1.1-1.6] in Example 1 contain a single linear dependency: $f_{AB} = f_{ABC}$. Since the network is deterministic, this dependency guarantees that after the first time step $x_{AB}$ will equal $x_{ABC}$, so we write $x_{AB} = x_{ABC}$. We use this fact to eliminate $x_{AB}$, giving a new set of steady state equations:

$$
\begin{align*}
    f_A &= 1 - x_B \\
    f_B &= x_{AC} \\
    f_{AC} &= x_A - x_{ABC} \\
    t \geq 2 \\
    f_{ABC} &= x_{AC} - x_{ABC} \\
    f_\emptyset &= 1.
\end{align*}
$$

(1.3)

Our new set of equations has the same non-zero eigenspace as the original set [1.1-1.6], except Eq. (1.3) is only valid from the second time step onwards. However, the new equations lack the null eigenspace because we removed the only linear dependency. States lying in the null eigenspace by definition decay and therefore correspond to transients in the time evolution, whereas eigenvectors whose eigenvalues have magnitude 1 do not decay and are part of the final attractor states. The 5 eigenvalues all have phases that are multiples of $2\pi/5$, indicating that the sole attractor is a limit cycle with a period of 5 time steps. The states are: $(000) \rightarrow (100) \rightarrow (110) \rightarrow (011) \rightarrow (000) \rightarrow \ldots$ at which point the sequence repeats.

Constraints are traditionally enforced by substitution; however, there are two possible issues with substituting constraint expressions in our case. First, there is no guarantee that when applying two constraints the second will not undo the work of the first, for example by reintroducing an index on a variable that was eliminated by the first constraint. The second issue is that two dependencies might constrain overlapping indices on the same variable. We have developed the following two constraint algorithms for dealing with these issues, designed for probabilistic and deterministic models respectively.

**Equation-reduction 1** The most straightforward way to simplify the equations is to use a variable-substitution strategy that prevents a variable from reappearing in a calculation after it has been eliminated. Our strategy solves each dependency for its ‘lowest’ variable based on a binary representation of its indices and eliminates that variable from the rest of the calculation by simple substitution. Each constraint thus replaces its substituted variable exclusively with ‘higher’-ordered variables, thus guaranteeing that the application of constraints will never cause an infinite loop. We retain each constraint in memory and reapply it any time the substituted variable

\[ [\text{Example 1, continued}] \]
reappears in another time-evolution equation. Note that equation-reduction method 1 may only apply one constraint if two constraints apply to a given variable. This reduction method is the only one that can be used by PBNs or asynchronous networks.

**Equation-reduction 2** There will be no time savings in the calculation if we only eliminate a variable after we have already computed its time-evolution function. Our second equation-reduction method simplifies a calculation while solving for a model’s long-term behavior, by taking advantage of the fact that each linear dependency involving Boolean variables constrains not only those variables, but also any variables that are factorized by them (i.e. they have all of the dependent variable’s subscripts). Thus a dependency involving a low-order variable with few indices can exclude a significant fraction of the variable space. We find these constraints concurrently with the process of adding equations, and thereby potentially avoid having to evaluate a significant fraction of our variable space. This is the default reduction method for deterministic networks.

For example, in the network of Figure 3, variable \( x_{ABC} \) is subject to constraints from both \( x_{AB} \) and \( x_{BC} \), and substituting one of these constraints could potentially eliminate the subscript \( B \) that the other constraint requires. We avoid both these problems by multiplying constraints rather than substituting them, using the fact that we can freely duplicate (or remove duplicates of) constrained indices on Boolean variables (because a Boolean raised to any positive power equals itself). This process forces the removal of all variables containing certain indices and lacking others, of which there is always at least one.

**Example 2: applying constraints**

Suppose we want to find the long-time behavior of Boolean variables A, C, and E in the network of Figure 3. After two iterations of solving for time-evolution functions we have:

\[
\begin{align*}
    f_A &= x_{BC} \\
    f_C &= x_A + x_B - x_{AB} \\
    f_E &= x_D \\
    f_B &= x_A \\
    f_D &= x_{ABC} \\
    f_{AB} &= x_{ABC} \\
    f_{BC} &= x_A
\end{align*}
\]

At this point there are two linear dependencies: \( f_{AB} = f_D \) and \( f_{BC} = f_B \), implying that

\[
\begin{align*}
    x_{AB} &= t \geq 2 x_D \\
    x_D &= t \geq 2 x_{AB} \\
    x_B &= t \geq 2 x_{BC}
\end{align*}
\]

Since we are only interested in the long-time behavior, we will use these constraints to simplify our equations. For example, if we were to retain \( x_{ABC} \) it would be affected by the relationships involving \( x_{AB}, x_B, \) and \( x_{BC} \), and it would not be possible to enforce all of these by substitution because there is only one \( B \)-index on \( x_{ABC} \). But our method enforces constraints by multiplying them:

\[
\begin{align*}
    x_{ABC} &= x_{AB} \cdot x_B \cdot x_{BC} \cdot x_{ABC} \\
              &= x_D \cdot x_{BC} \cdot x_B \cdot x_{ABC} \\
              &= x_{ABCD}.
\end{align*}
\]

More generally, the first constraint attaches an \( AB \) index to every variable containing a \( D \), and a \( D \) index to each variable with \( AB \) indices, and the second constraint adds a \( C \) index to every variable with a \( B \) index.

Constraining our system and eliminating disused variables gives us

![Figure 3. The network used in Example 2.](image-url)
This produces new dependencies:

\[ f_{AE} \equiv x_{ABCD} \quad (2.2) \]
\[ x_{ABCD} \equiv x_{AE} \quad (C7) \]
\[ x_{CE} \equiv x_{ABCD} \quad (C8) \]

With these constraints our example ends with the following closed system of dynamical equations:

\[ f_{AC} = x_{BC} \quad (2.7') \]
\[ f_{ABCDE} = x_{ABCDE} \quad (2.13) \]
\[ f_{AC} = x_{BC}. \]

The attractor is always reached at or before time step 3. The constraint equations \((C1 \ldots C9)\) map our variables of interest \((x_A, x_C, \text{and } x_E)\) to linear combinations of variables in the final system: for example, \(x_A\) is mapped by constraint \(C4\) to \(x_{AC} + x_{AE} - x_{ABC}\), then mapped using constraints \(C1, C7\) and \(C8\) to \(x_{AC}\), whose dynamics are given by the final time-evolution equations. The eigenvalues of this final system are \((-1, 1, 1)\), implying a steady state along with a period-2 cycle.

Note that multiplication of variables becomes a union operation on the variable indices in the product basis.

Each dependency produces at least one constraint that permanently eliminates at least one of the dependent variables, and often many other variables as well. \textit{Proof}: if the dependency contains only one term, then that variable is zero, eliminating it and all variables it factorizes. If there is more than one term in the dependency, then each lowest-index variable (which may be \(x_3 = 1\)) accumulates at least one index from another variable in the dependency. Therefore each lowest-index variable is always eliminated. \textit{Corollary}: the calculation eventually terminates because there are a finite number of variables, and each new linear dependency removes at least one further variable.

Applying the equation-reduction method to each dependency in our linear system \(F = \{f_i\}\) is guaranteed to eliminate the entire null space of \(F\), simply because variables will be removed from the system until there are no more dependencies. On the other hand, our equation-reduction method does not affect the non-null eigenspace involving the variables of interest, because the constraints \(C\) map those variables to variables in the final equations: \(X_{final} = C \cdot X_{interest}\). Therefore the long-time behavior is accurately modeled by \(F_{interest} = C^T F_{final} C\), which therefore contains all the persistent eigenmodes.
In a deterministic Boolean network all eigenvalues of $F$ have modulus either 0 or 1, owing to the fact that the state space (and therefore by Appendix 1 our product-basis space) is finite. Therefore every eigenmode must either decay to zero in finite time (implying that $\lambda = 0$), or else repeat in $\Delta t$ time steps ($|\lambda| = 1$ and $\arg(\lambda) = 2\pi k/\Delta t$ for integer $k$).

**Probabilistic and asynchronous Boolean networks**

Our method can be used to model large populations of probabilistic Boolean networks (PBNs) \[21, 27\], in which several state transitions are possible at each time step, and the various transitions may have different probabilities. The time evolution of an individual PBN is stochastic, so from the beginning we work with the population-averaged $f_{ij}$, etc., which can be thought of as the unconditional probabilities of an individual PBN evolving by $x_i \leftarrow 1$, $x_{ij} \leftarrow 1$, etc. over a time step. In the population-averaged variables, the time evolution of each product variable $f_{ij}(x)$ is again a product of single-variable time-evolution functions $f_i : f_j$, because the constituent single-variable functions $(f_i, f_j)$ are drawn independently for each individual in the population. Thus we can build time-evolution equations for PBNs that are valid for all time using the same equation-multiplying process we used for deterministic networks. Since PBN equations are solved using population-averaged variables that are not Boolean (so $f_{ij} \neq f_{ij}^2$) we are only able to constrain these equations using equation-reduction method 1: a constraint on a given averaged variable $x_{ij}$ cannot be used to factorize higher-order averaged variables such as $x_{ij}^2$; thus reduction method 2 cannot work.

The full system of linear time-evolution equations $F$ of a PBN contains eigenmodes having real-valued eigenvalues whose modulus is on the interval $[0, 1]$. (A modulus larger than 1 would represent a mode growing without bound, which is impossible viewed from the state-space basis because the population fraction in each state $b_i$ is restricted to the interval $[0, 1]$). Therefore, unlike a deterministic network, a PBN can have slowly-decaying modes with eigenvalues between 0 and 1. For PBNs we generalize our equation-reduction method to identify decaying modes before all $f_i$ have been solved (i.e. at a stage where we have only computed some $F_{\text{partial}}$ which is in general a rectangular matrix). These modes $m$ have the property $m \cdot F = \lambda [m 0]$. We discard these modes after they have become sufficiently small, defined by $e^{-\lambda(t_{\text{max}}(f_{m}))} < \epsilon$ where $t_{\text{max}}$ are the ‘starting times’ of the involved equations and $\epsilon$ is a user-defined threshold.

Large populations of asynchronous networks behave identically to large populations of PBNs \[23\] if we define a uniform time step: the likelihoods of the various possible updates give the state-transition weights in the corresponding synchronous PBN. If this time step is small enough, then the likelihood of two causally-connected asynchronous updates happening in the same step is small, and in this limit the local update rules for a PBN accurately model the asynchronous network. Therefore our analysis also applies to large populations of asynchronous networks for small time steps. The conversion of an asynchronous updating scheme to a PBN lowers some of the eigenvalues that would otherwise be of magnitude 1 in a synchronous version of the network, and therefore our equation-reduction method can identify more degeneracies in asynchronous deterministic networks than in their synchronous counterparts, partially compensating for the fact that each degeneracy allows for the removal of only one variable.

**Calculational notes** In order to be useful for simplifying calculations, the two equation-reduction methods must identify linear dependencies in our update rules before the linear system closes – in other words, when our partially-computed update function $F_{\text{partial}}$ is still a rectangular matrix. Our implementation of the reduction method finds these dependencies by using QR decomposition to find a set of dependent variables, then solves the update rules for these variables in terms of the independent variables using matrix division. For PBNs, the complication is that decaying eigenvalues of the full system are generally nonzero, so the decaying modes are not simple linear dependencies. Note that each decaying eigenvalue is part of the spectrum of $F_\square$ (the square component of $F_{\text{partial}}$), although the reverse is not true. Our implementation uses the spectrum of $F_\square$ as a set of candidate eigenvalues, and for each candidate $\lambda$ finds the linearly dependent subspace using a QR decomposition of $F_{\text{partial}} - \lambda I_{\text{partial}}$, where $I$ is rectangular with ones on the diagonal and zeros elsewhere.

When using equation-reduction method 2, the order in which we calculate the time-evolution of new variables and remove dependencies greatly influences the complexity of the calculation. One general rule we have found is that constraints on the lowest-index variables are most helpful for deterministic networks, because they factorize the largest part of the variable space. Following this rule of thumb, our implementation solves the time-evolution equations for only those variables having the fewest indices between each search for new dependencies. Additionally, our code solves for low-index factors of new variables even if they were not directly involved in earlier equations in hopes that they will produce helpful constraints. In most tests, this prioritization method speeds up the calculation.

The product-basis method, particularly the equation-reduction component, is susceptible to numerical problems that fall into two categories: 1) polynomial coefficients can become very large, and 2) gradual erosion of
numerical precision can blur the distinction between zero and nonzero terms. Our implementation of the product-basis method is usually able to prevent (1) from becoming a problem in deterministic networks, due to an extra filter in the calculation (see below). In addition our code monitors both sources of error; in the latter case by tracking the floating-point error in each step of the calculation. When either type of calculation error exceeds defined bounds, our method exits with an error message.

The reason polynomial coefficients can become large is that multiplication by constraints can produce several identical terms that add together. For example, the constraint \(x_1 = x_2 + x_3\) takes \(x_{123} \rightarrow 2x_{123}\). Typically the variable being multiplied would be found to be zero later in the calculation (so having a coefficient of 2 is not an error in the math), but if many such constraints are applied and the coefficient grows exponentially then the numerics will eventually fail. We have included a filter that can usually identify these outlier coefficients and remove them for deterministic networks using the rule that all individual (not population-averaged) product variables are either 0 or 1 at all time steps. For example, if \(f_4 = 2x_{123}\) then we can reason that if \(f_4\) is either 0 or 1, then \(x_{123}\) cannot be 1 and therefore we have a new constraint \(x_{123} = 0\) which simplifies \(f_4 \rightarrow 0\). In general, we identify these removable terms using the heuristic that if a large coefficient multiplying a combination of integer-weighted variables is greater than the sum of all other coefficients plus one, then that combination of variables (which is an integer for a homogeneous population) must be zero.

**Discussion**

Our product-basis method allows the direct simulation of highly heterogeneous populations, including the transient processes that are ignored by Boolean-attractor analyses. We can also use equation-reduction to find the late-time behavior at lower computational cost, which in the limit gives the long-term attractor dynamics of the target variables. This is a slightly different output than that produced by most existing Boolean analytic methods which generate the attractors of the entire network state. Our approach can be used to follow single variables of the system over time, as well as the correlations between these variables that are both necessary and sufficient to fully describe the dynamics of the population. We also showed that our method, when applied to a network augmented by mutation nodes, can effectively explore heterogeneity in the network rules in addition to heterogeneity in network state. In each of our simulations, all subpopulations are exactly accounted for in the output time series, no matter how rare. The only requirement for use of our approach is that the underlying model be built using Boolean variables.

The key to our method is to write the time-evolution equations as a linear system, but in a different basis than the usual state space basis. Our variables have several advantages over state space variables. First, descriptors of a mixed population naturally use words that correspond more closely to our variables than to individual states. For example, we might specify that half the population starts with both genes \(A\) and \(B\) on, which implies that \(x_{AB} = 0.5\), but is agnostic about the state of other variables. Another advantage is that our equations often close using relatively few of our product variables for any mixed population, whereas the number of equations required in the state space basis scales with the heterogeneity of the population: the simulations we showed in Figure 1 would require all \(2^N\) state space variables. Thus our choice of variables is better for modeling very heterogeneous populations. Finally, our basis allows some variables to factorize others, allowing us to vastly simplify the calculation in many cases where we only care about the long-term behavior.

We acknowledge that our method can become intractable for complex networks due to the fact that the construction of these simulations is potentially an exponential problem. A full simulation can require up to \(2^N\) equations to model; even the attractor analysis is known to be NP-hard [25]. Large size, complex logic rules, and certain feedback loops seem to cause the equation set to be large. These are fundamental limitations. However, the equation-reduction method for finding attractors is somewhat of an art, and our future work will aim to improve this part of the calculation for typical network models. Notably, our current implementation of the two equation-reduction methods often aborts due to numerical error exceeding bounds; improving the numerical stability will be a priority for future work.

The product-basis method can be applied to any system involving heterogeneous populations, as long as the individuals in a population can be modeled using Boolean logic. Heterogeneity plays a major role in such varied systems as healthy and cancerous tissues, evolution at the organism scale, and the social dynamics of unique individuals [29]. In all of these cases, rare and unexpected dynamics are difficult to capture by simulations of individuals, while pure attractor analyses may miss important aspects of the dynamics. We have demonstrated that the methodology outlined here can help to capture these important but elusive events.
Appendices

Appendix 1: correlation variables form a complete and independent basis

For $N$ Boolean variables there are $2^N$ variables in the state space basis (labeled $b$): this is just the number of states of the system. Likewise there are $2^N$ variables in the product space basis (labeled $x$) because there are $2^N$ combinations of subscripts on these variables: each of $N$ subscripts may be present or absent. Therefore the two spaces have the same number of variables, but this does not prove that the product-basis spans the entire Boolean space. In order to show that the $x$-variables form a complete basis in $b$-space, we imagine explicitly writing the transformation matrix from $b$-space to $x$-space. For example, for the case of three Boolean variables this matrix is:

$$
T_{xb} = \begin{bmatrix}
1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
1 & 0 & 1 & 0 & 1 & 0 & 1 & 1 \\
0 & 1 & 1 & 0 & 0 & 1 & 1 & 1 \\
0 & 0 & 1 & 0 & 0 & 0 & 1 & 1 \\
0 & 0 & 0 & 1 & 1 & 1 & 1 & 1 \\
0 & 0 & 0 & 0 & 1 & 1 & 1 & 1 \\
0 & 0 & 0 & 0 & 1 & 1 & 1 & 1 \\
0 & 0 & 0 & 0 & 0 & 1 & 1 & 1 \\
0 & 0 & 0 & 0 & 0 & 0 & 1 & 1 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 \\
\end{bmatrix}
$$

The transformation matrix is upper-triangular, with ones on the diagonal. The reason is that each $x$-variable is turned on by the $b$-variable having the same indices (hence the ones along the diagonal), and by any $b$-variables containing additional indices implying a higher position in the matrix (since the indices are arranged in binary order of their subscripts); however an $x$-variable is never turned on by a $b$-variable that is missing one of its indices, which is why the lower triangular block is empty. Since the matrix is triangular the eigenvalues are the ones on the diagonal, so the determinant is one, the transformation is non-degenerate (and volume-preserving) and therefore the $x$-space spans the $b$-space.

Appendix 2: algorithm and code

Pseudocode is given in Algorithm 1. The full code is available at: [https://github.com/CostelloLab/ProductBasis](https://github.com/CostelloLab/ProductBasis).
Algorithm 1 build closed system of equations $F$

1: Initialize set of unsolved variables with variables of interest: $X \leftarrow \{x_1, x_2, ..., x_n\}$
2: Initialize set of variable update rules: $F \leftarrow \emptyset$
3: Initialize set of constraints: $C \leftarrow \emptyset$
4: Initialize simulation start time: $t_{sim} \leftarrow 1$
5: Initialize set of equation start times of $F$: $T_F \leftarrow \emptyset$
6: Initialize set of constraint start times: $T_C \leftarrow \emptyset$

7: while $X$ is not empty do

8:  % Reduce equations if necessary
9:  if $\text{size}(F) > \text{equation_reduction_threshold}$ then
10:     $F_{\Box} \leftarrow$ square-matrix component of $F$ (i.e. only terms in solved variables for each $f_i$)
11:     $\lambda \leftarrow$ set of eigenvalues of $F_{\Box}^T$
12:     for each $\lambda_i \in \lambda$ do
13:         $G \leftarrow F - \lambda_i[I \ 0]$
14:         $R \leftarrow$ upper-triangular matrix from $QR(G)$
15:         $V_j \leftarrow$ set of dependent variables $j$ found from $|R_{jj}| < \epsilon$
16:         $D \leftarrow$ set of linear-dependencies found by $V_j = G(V_j)/G(V_j)$
17:         if $V_j \neq \emptyset$ then
18:             $t_{sim} \leftarrow \min(t_{sim}, \min(T_F(x_k \in D)) + \max(1, \ceil(\log \epsilon_{\text{decay}}/ \log |\lambda_i|)))$
19:             break
20:         end if
21:     end for
22:     $D \leftarrow$ sort $\{d_i \text{ for which } n_{\text{terms}}(d_i) \leq 2 \min n_{\text{terms}}(d)\}$ by number of terms
23:     for each $d_i \in D$ do
24:         NewConstraints($d_i, t_{sim}$)
25:     end for
26:     Sort $C$ by number of terms
27:     end if
28:  % Add new equations
29:  $X \leftarrow X \cup$ lowest-index factors of $X$
30:  Sort $X$ by number of terms
31:  for each $x_i \in X$ do
32:      $f_i \leftarrow 1$
33:      for each Boolean factor $x_b$ of $x_i$ do
34:          if $f$ is a discrete update rule then
35:              $f_i \leftarrow f_i \cdot f_b$
36:          else
37:              $f_i \leftarrow f_i + f_b \cdot \prod_{x_j \neq x_i \in X} x_j$
38:          end if
39:      end for
40:  $F \leftarrow F \cup \text{Constrain}(f_i, 1)$
41:  end for
42: end while
function NewConstraints(d, tc)  
  if d ← RemoveOversizeCoefficients(d/minCoeff(d)) adds no new constraint then  
    for each xj ∈ d that does not factor another xk≠j ∈ d do  
      RHSj ← solve (d = 0) for xj  
      if {xj} ≠ Constrain({RHSj}, tc)[1] then  
        for each prior constraint (xp = RHSp) do  
          if xp is factorized by xj and xp·RHSj = RHSp then  
            C ← C \ {cp}  
          end if  
        end for  
      end if  
    end for  
    C ← RemoveOversizeCoefficients(C \ {xj = RHSj})  
    for each xk = multiples of xj do  
      Xnew ← new terms in Constrain(xk)  
      X ← X ∪ Xnew  
      A ← {al = coefficients of xk in F}  
      F ← F + A·(RHSk − xk)  
      (tf for all l such that al ≠ 0) ← tc  
    end for  
  end if  
end function

function Constrain(poly, tc)  
  while ∃ unconstrained terms in poly do  
    for each constraint xi = RHSi do  
      % Multiply all factored variables by the constraint  
      for each unconstrained variable xj with coefficient aj in poly do  
        if xj is factorized by xi then  
          polynew ← xj·RHSi  
          if xj ⊈ polynew then  
            poly ← (poly − aj·xj) ∪ polynew  
            tfi ← tc  
          end if  
        end if  
      end for  
    end for  
    poly ← RemoveOversizeCoefficients(poly)  
  end while  
return Polys  
end function

% Check for overlarge coefficients  
function RemoveOversizeCoefficients(poly)  
  while ∃ unchecked coefficients in poly do  
    for each coefficient ai in poly do  
      nj = round(ai/aj)  
      if |ai| > ∑ j |aj − njaj| then  
        NewConstraints(nj, tsim)  
        aj ← aj − njaj for all j  
        break  
      end if  
    end for  
  end while  
return Polys  
end function
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