

Combinatorial Representation of Parameter Space for Switching Networks*

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Abstract. We describe the theoretical and computational framework for the Dynamic Signatures Generated by Regulatory Networks (DSGRN) database. The motivation stems from an urgent need to understand the global dynamics of biologically relevant signal transduction/gene regulatory networks that have at least 5 to 10 nodes, involve multiple interactions, and have decades of parameters. The input to the database computations is a regulatory network, i.e., a directed graph with edges indicating up or down regulation. A computational model based on switching networks is generated from the regulatory network. The phase space dimension of this model equals the number of nodes and the associated parameter space consists of one parameter for each node (a decay rate) and three parameters for each edge (low level of expression, high level of expression, and threshold at which expression levels change). Since the nonlinearities of switching systems are piecewise constant, there is a natural decomposition of phase space into cells from which the dynamics can be described combinatorially in terms of a state transition graph. This in turn leads to a compact representation of the global dynamics called an annotated Morse graph that identifies recurrent and nonrecurrent dynamics. The focus of this paper is on the construction of a natural computable finite decomposition of parameter space into domains where the annotated Morse graph description of dynamics is constant. We use this decomposition to construct an SQL database that can be effectively searched for dynamical signatures such as bistability, stable or unstable oscillations, and stable equilibria. We include two simple 3-node networks to provide small explicit examples of the type of information stored in the DSGRN database. To demonstrate the computational capabilities of this system we consider a simple network associated with p53 that involves 5 nodes and a 29-dimensional parameter space.

Key words. regulatory networks, global dynamics, switching systems

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1. Introduction. Though the method presented in this paper is general, our primary motivation arises from the need to understand the global dynamics of signal transduction/gene regulatory networks, e.g., [35]. Our mathematical abstraction of a regulatory network \mathbf{RN} is a directed graph where the nodes (vertices) $V = \{1, \dots, N\}$ indicate the species and the edges E indicate the activation or repression of production of one species by another (this

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is made precise in Definition 2.1). There are at least three fundamental challenges in determining if a given regulatory network provides a biologically relevant model: determining completeness, authentication, and nonlinear interaction. As discussed in [36] completeness and authentication are concerned with whether the relevant species are included in the regulatory network and whether the proposed interactions are correct. Genomic sequencing data informs completeness, while biochemical knowledge is required for authentication. While the quantity of genomic data is rapidly increasing, detailed biochemical information is still sparse. Since the interaction between species is typically governed by multiscale processes determining appropriate explicit nonlinearities let alone realistic physical parameters is extremely difficult.

We propose to address these challenges by employing a crude, compact, robust, mathematically rigorous, finitely presented description of dynamics that allows for a combinatorial representation of parameter space. As a consequence, given a regulatory network \mathbf{RN} , we are able to build the associated *database of dynamic signatures* which codifies the global dynamics over all of parameter space. The underlying theoretical ideas have already been exploited to study a variety of mathematical and biological models [4, 8, 9, 7]. However, to effectively use these ideas in the context of moderately sized gene regulatory networks where completeness and authentication are in question, we require orders of magnitude greater efficiency for approximating the dynamics and the ability to work with much higher dimensional parameter spaces. This paper describes and justifies a revised approach that achieves the desired efficiency and provides a natural decomposition of parameter space.

Our starting point has much in common with an approach often referred to as logical networks [2]. The difficulties, alluded to above, in determining and then parameterizing appropriate interactions and nonlinearities has led to the widespread use of relatively simple models that aim to capture qualitative features of the dynamics. The simplest ones are the Boolean models, where each node is represented as either on or off; the dynamics of the i th node consist of evaluation of a logical function defined by this binary representation of the state of the system. The evolution of the network proceeds in discrete steps in time which can either synchronously evaluate all functions [3, 13] or do so asynchronously [10, 45, 38].

Our approach is most closely associated with asynchronous logical networks. Given a regulatory network \mathbf{RN} with N nodes, the associated *switching system* [27, 28, 29, 30, 17, 18, 19, 16] is an N -dimensional system of ordinary differential equations of the form

$$(1) \quad \dot{x} = -\Gamma x + \Lambda(x),$$

where Γ is a diagonal matrix with positive entries and Λ is a piecewise constant function (see Definition 2.4).

There are a variety of parameters associated with switching networks. To each variable x_i there is a *decay rate* $\gamma_i > 0$ (the diagonal terms in Γ). To each edge in \mathbf{RN} , corresponding to the impact of species i on j , we associate two positive expression levels, low $l_{j,i}$ and high $u_{j,i}$, and a threshold $\theta_{j,i}$ for x_i at which the expression levels switch. The values of the function Λ are determined by sums and products of the parameters l and u where the particular formula is generated by logical rules that indicate how different nodes interact. Thus given a regulatory network the set of parameters lies in $[0, \infty)^D$, where $D = N + 3 \cdot \#(E)$, where N is the number

of nodes, and $\#(E)$ is the number of edges. Our goal is to characterize the global dynamics for every point in parameter space.

We hasten to add (this is made clear in the sections that follow) that we do not view (1) as a mathematical model for the biological process; rather (1) is used only to motivate the combinatorial computations that are the focus of this paper. To emphasize this point we make two comments. First, Λ is piecewise constant (and not even defined when $x_i = \theta_{j,i}$); thus classical trajectories to (1) do not exist. Second, the main result of [25] is that applying the methods described in this paper to two-dimensional switching systems results in a representation of the global dynamics that is valid for a much wider family of nonlinearities, i.e., for a system of the form

$$(2) \quad \dot{x} = -\Gamma x + f(x)$$

where f is a Lipschitz continuous function. Algebraic topological tools such as the Conley index [39] can be applied to this representation to extract information concerning the structure of invariant sets for dynamics of (2). We also note that for a typical parameter value one can obtain explicit a priori estimates in terms of parameters of the system (see [25]) for how much the nonlinearity f can differ from the switching nonlinearity Λ . It is in this sense that we view (1) as a computational tool, rather than the mathematical model of biological reality. An important implication is that we can obtain rigorous results about the dynamics without explicit/detailed knowledge of the appropriate nonlinearity for the biological problem of interest. It is worth contrasting our approach to more classical methods for relating the dynamics of smoothed systems (2) with the discontinuous switching systems (1) [21, 33, 47].

To approximate the dynamics of the switching system at a fixed parameter value we use the thresholds θ to decompose phase space. These decompositions are then used to determine state transition diagrams (see section 3). In this format the dynamics is represented by a sparse directed graph \mathcal{F} with roughly $\prod_{n=1}^N [O(n) + 1]$ vertices, where $O(n)$ is the number of out-edges at node n . Given the size of the regulatory networks that we are interested in analyzing, storing all the state transition graphs that arise as one sweeps through parameter space is impractical. With this in mind we focus on the essential dynamical structures: the *recurrent dynamics*, i.e., the nontrivial strongly connected components of \mathcal{F} ; and the *gradient-like dynamics*, i.e., the reachability, defined by paths in \mathcal{F} , between the recurrent components. There are efficient (both in time and memory) graph algorithms that allow one to identify strongly connected components (see [4, 8] and references therein, and see [6] for an application of these techniques in the context of regulatory networks). For many networks of interest this step can be performed rapidly enough so that it is tractable to repeat it over many parameter values. We encode this information in the form of an annotated *Morse graph* $\text{MG}(\mathcal{F})$. The Morse graph is the minimal directed acyclic graph such that each nontrivial strongly connected component is represented by a distinct node and the edges indicate the reachability information inherited from \mathcal{F} between the nodes (see section 3.3). The annotations consist of optional information, typically problem specific, that allow for easier identification between the dynamics captured by our approach and the dynamics of biological interest.

The computational steps described above are valid for individual parameter values. A fundamental contribution of this paper is the identification of a natural decomposition of

parameter space into regions, called *parameter cells*, over which the transition graphs and hence Morse graphs are constant. The cells are given in terms of explicit polynomial inequalities in the parameter values and hence take the form of semialgebraic sets. Since the global dynamics of switching networks is parameter dependent and we are working with high dimensional parameter spaces it should come as no surprise that understanding the geometry and organizing the structure of all parameter cells is nontrivial. With this in mind we introduce the *parameter graph* (see section 4), an undirected graph where each node is identified with a parameter cell and the edges provide information about how the parameter cells are related. In fact, we make use of two parameter graphs. The first, called the *geometric parameter graph* (GPG), is based on the topology of parameter space as a subset of $[0, \infty)^D$. This provides a description of the decomposition of parameter space in a language familiar to researchers in the field of dynamical systems. The second, called the *combinatorial parameter graph* (CPG), is what is actually computed. We prove that there is a graph homomorphism $h: \text{GPG} \rightarrow \text{CPG}$ and conjecture that the geometric and combinatorial parameter graphs are equivalent but only have a proof (see Theorem 4.18) for regulatory networks whose nodes have at most three in-edges and out-edges.

Our construction of the CPG for **RN** is based on two facts. First, the graph structure is actually a canonical graph product over factor graphs CPG_n which depend only on the local structure of **RN** around each network node n . Second, each CPG_n is a connected subgraph of a larger graph of combinatorial parameters and may be constructed via a graph traversal search for *realizable* combinatorial parameters, i.e., those that are realized by some geometric parameter of **RN**. In particular, for each node n in **RN** we compute the set of possible combinatorial parameters, which is determined by the number of out-edges at n , the number of in-edges at n , and the logic that determines how the information from the in-edges is processed. To identify whether a particular combinatorial parameter is realizable we make use of cylindrical algebraic decompositions (CAD) [15] to determine if there exists a solution to an associated set of inequalities. Recall that a CAD of a semialgebraic set is a recursive set of inequalities that defines the elements of the set. As discussed in Remark 4.16, CAD computations are expensive. However, once done they can be reused at a constant cost. Thus our strategy is to perform the CAD computations and store them. A list of the node structures for which CAD computations have been performed is given in Table 1.

Our inability to prove that $h: \text{GPG} \rightarrow \text{CPG}$ is always an isomorphism stems from our lack of general understanding of the geometry of individual and pairs of regions on which transition graphs are constant. However, the CAD computations, once performed, provide sufficient information to check the conjecture. This is the essence of the proof of Theorem 4.18.

The concepts and techniques introduced in this paper have allowed us to develop the DSGRN (*Dynamic Signatures Generated by Regulatory Networks*) software [32] that has the following features and capabilities:

- It can compute the size of parameter graphs (number of parameter nodes) for any regulatory network constructed using components found in Table 1. In particular we supply a web-based program to design such networks which automatically tabulates the size of the parameter graph.

Table 1
Network node components.

| $\#\mathbf{S}(i)$ | $\#\mathbf{T}(i)$ | M_i | $\#\mathbf{PG}_i/\#\mathbf{T}(i)!$ |
|-------------------|-------------------|-------------|------------------------------------|
| 1 | 1 | x | 3 |
| 1 | 2 | x | 6 |
| 1 | 3 | x | 10 |
| 2 | 1 | $x + y$ | 6 |
| 2 | 1 | xy | 6 |
| 2 | 2 | $x + y$ | 20 |
| 2 | 2 | xy | 20 |
| 2 | 3 | $x + y$ | 50 |
| 2 | 3 | xy | 50 |
| 3 | 1 | $x + y + z$ | 20 |
| 3 | 1 | xyz | 20 |
| 3 | 1 | $x(y + z)$ | 20 |
| 3 | 2 | $x + y + z$ | 150 |
| 3 | 2 | xyz | 150 |
| 3 | 2 | $x(y + z)$ | 155 |
| 3 | 3 | $x + y + z$ | 707 |
| 3 | 3 | xyz | 707 |
| 3 | 3 | $x(y + z)$ | 756 |

- It can access a database of CADs of parameter cells corresponding to parameter nodes for regulatory networks constructed using components found in Table 1.
- It can compute annotated Morse graphs given a parameter (cell) and a regulatory network.
- It can compute databases of annotated Morse graphs over an entire (combinatorial) parameter graph given a regulatory network. These databases are designed using SQL and support a range of queries over Morse graph attributes and annotations.
- We supply a web interface to interact with databases that can filter parameter graphs to show only nodes which satisfy certain queries.
- We supply a command line interface which allows access to phase space information for the associated switching system of a regulatory network given a particular parameter of interest.
- We have supplied documentation of the program along with tutorial materials.

To provide the reader with intuition concerning the output in section 5 we consider three regulatory networks. The first two, the repressilator and the bistable repressilator, consist of three nodes and three and four edges, respectively. For these examples the parameter graphs are sufficiently small that they can be easily visualized. In general, the output of DSGRN grows rapidly as a function of the size of the regulatory network and thus can be accessed efficiently only through queries. To give a sense of the computational capabilities of DSGRN for problems of biological interest we consider a subnetwork associated with p53 and report the computational times and costs. The parameter graph information can be accessed elsewhere [32].

Before concluding this introduction we return to the challenge of determining completeness and authentication where we believe DSGRN can be a useful tool. These challenges imply that

in early stages of modeling one cannot necessarily assume that a proposed regulatory network is “correct.” By allowing one to compare the output of the model dynamics against experimental data, Boolean models provide a computationally tractable means to attempt to identify the existence of missing species and interaction and/or to exclude nonexistent interactions and unnecessary species (see, for example, [12, 11, 31, 40]). Because the computations that we perform to identify the dynamics is purely combinatorial, the cost of our computations is similar to that of pure Boolean models. However, because we model using real numbers and ordinary differential equations, DSGRN can capture finer dynamical structures that presumably can be more readily identified with experimental values. More significantly, the fact that DSGRN provides a complete description of the dynamics over all of parameter space opens up new opportunities for deciding upon the plausibility of or comparison of different models, e.g., how stable is the desired dynamic phenotype to changes in parameters, and for control of the dynamics, e.g., which changes in parameters result in a desired dynamic phenotype. We leave the implementation of these ideas to future works.

2. Switching networks. We review the essential concepts of switching networks. In section 2.1 we define regulatory networks and their associated switching systems. In section 2.2 we discuss the interpretation of the nonlinearities in the switching system as performing logical operations on the inputs. In section 2.3 we use the discontinuities of the nonlinearity of (1) to impose a natural decomposition of phase space.

2.1. Regulatory networks.

Definition 2.1. A regulatory network $\mathbf{RN} = (V, E)$ is an annotated finite directed graph with vertices $V = \{1, \dots, N\}$ called network nodes and annotated directed edges $E \subset V \times V \times \{\rightarrow, \vdash\}$ called interactions. An \rightarrow annotated edge is referred to as an activation and an \vdash annotated edge is called a repression. We indicate that either $i \rightarrow j$ or $i \vdash j$ without specifying which by writing $(i, j) \in E$. We allow for self-edges but admit at most one edge between any two nodes, e.g., we cannot have both $i \rightarrow j$ and $i \vdash j$ simultaneously. The set of sources and targets of a node n are denoted by

$$\mathbf{S}(n) := \{i \mid (i, n) \in E\} \quad \text{and} \quad \mathbf{T}(n) := \{j \mid (n, j) \in E\}.$$

The cardinality of $\mathbf{S}(n)$ and $\mathbf{T}(n)$ are denoted by $\#\mathbf{S}(n)$ and $\#\mathbf{T}(n)$. Each node is equipped with a multilinear function $M_i : \mathbb{R}^{\mathbf{S}(i)} \rightarrow \mathbb{R}$, called the logic of node i (a detailed description of its derivation is presented in section 2.2).

For convenience we abuse notation and occasionally write a network node as x_i instead of i . For instance, we may write $(i, j, \rightarrow) \in E$ and $(i, j, \vdash) \in E$, respectively, as $x_i \rightarrow x_j$ and $x_i \vdash x_j$.

Remark 2.2. Throughout this paper we assume that the regulatory network \mathbf{RN} does not have any direct negative self-regulation $i \vdash i$ for any i . This is done for technical reasons related to the code (see Remark 3.9). This is not a serious restriction. In biological systems negative self-regulation is often mediated by an intermediary, e.g., $x_i \rightarrow X_i \vdash x_i$ [20, 23]. Furthermore, future planned developments of the code will allow the user to remove this restriction.

Definition 2.3. Given a regulatory network $\mathbf{RN} = (V, E)$, for each edge $(i, j) \in E$ (i.e., $i \rightarrow j$ or $i \dashv j$) we associate three parameters: $l_{j,i}$, $u_{j,i}$, and $\theta_{j,i}$. (Note the matrix-style subscript order convention.) Additionally, to each node $i \in V$ we associate a decay rate γ_i . Each of these parameters is a real number, so we may regard the collection of all these parameters as a tuple $(l, u, \theta, \gamma) \in \mathbb{R}^D$. We call this collection of numbers a parameter for \mathbf{RN} .

Definition 2.4. Given a regulatory network \mathbf{RN} the associated switching system at parameter $(l, u, \theta, \gamma) \in \mathbb{R}^D$, where $D = N + 3 \cdot \#(E)$, is given by

$$(3) \quad \dot{x}_j = -\gamma_j x_j + \Lambda_j(x), \quad j = 1, \dots, N,$$

where

$$(4) \quad \Lambda_j := M_j \circ \sigma_j.$$

Here $\sigma_j : \mathbb{R}^N \rightarrow \mathbb{R}^{\mathbf{S}(j)}$ is a multidimensional step function defined componentwise (i.e., by its coordinate projections $\pi_i(\sigma_j)$) for each $i \in \mathbf{S}(j)$ as

$$(5) \quad \sigma_{j,i} = \pi_i(\sigma_j(x)) := \begin{cases} l_{j,i} & \text{if } i \rightarrow j \text{ and } x_i < \theta_{j,i} \text{ or if } i \dashv j \text{ and } x_i > \theta_{j,i}, \\ u_{j,i} & \text{if } i \rightarrow j \text{ and } x_i > \theta_{j,i} \text{ or if } i \dashv j \text{ and } x_i < \theta_{j,i}, \\ \text{undefined} & \text{otherwise.} \end{cases}$$

Remark 2.5. Switching systems written in the form (3) are equivalent to (1) where Γ is the diagonal matrix with diagonal entries $\Gamma_{ii} := \gamma_i$ and the i th coordinate of Λ is Λ_i .

The dependence on x of the right-hand side of (3) involves the expression $\sigma_j(x)$. Since σ_j is a multidimensional step function which compares variables to thresholds, a grid-like structure is imposed upon phase space.

Definition 2.6. Let $z = (u, l, \theta, \gamma)$ be a parameter for the regulatory network \mathbf{RN} . For each $i \in V$, we adopt the convention that $\theta_{-\infty,i} = 0$ and $\theta_{\infty,i} = \infty$. For all $i \in V$, $j_1, j_2 \in V \cup \{-\infty, \infty\}$, we say $\theta_{j_1,i}$ and $\theta_{j_2,i}$ are consecutive thresholds if $\theta_{j_1,i} < \theta_{j_2,i}$ and there does not exist $\theta_{j,i}$ such that $\theta_{j_1,i} < \theta_{j,i} < \theta_{j_2,i}$. For each $i = 1, 2, \dots, N$, suppose $\theta_{a_i,i}$ and $\theta_{b_i,i}$ are consecutive thresholds. Then we say that the product of open intervals

$$\kappa := \prod_{i=1}^N (\theta_{a_i,i}, \theta_{b_i,i})$$

is a cell of the regulatory network. We denote the collection of cells as $\mathcal{K}(z)$. If $a_j \in V$, then we say that the bounded hyperplane

$$\kappa_j^- := \prod_{i=1}^{j-1} (\theta_{a_i,i}, \theta_{b_i,i}) \times \{\theta_{a_j,j}\} \times \prod_{i=j+1}^N (\theta_{a_i,i}, \theta_{b_i,i})$$

is a left face of κ with projection index j and switching index a_j . Similarly, if $b_j \in V$, then we say the bounded hyperplane

$$\kappa_j^+ := \prod_{i=1}^{j-1} (\theta_{a_i,i}, \theta_{b_i,i}) \times \{\theta_{b_j,j}\} \times \prod_{i=j+1}^N (\theta_{a_i,i}, \theta_{b_i,i})$$

is a right face of κ with projection index j and switching index b_j . A face of a cell κ is either a left or a right face of κ .

We restrict our focus to nonnegative parameters $\bar{Z} \subset [0, \infty)^D \subset \mathbb{R}^D$ for which we interpret the l and u values as the *lower* and *upper* values that may be taken, respectively. This imposes the additional requirement that $l_{j,i} \leq u_{j,i}$.

Definition 2.7. Given a switching network **RN** the associated parameter space \bar{Z} is defined to be the collection of all parameters $(l, u, \theta, \gamma) \in [0, \infty)^D$ for which $l_{j,i} \leq u_{j,i}$ for all $(i, j) \in E$. A parameter $z = (l, u, \theta, \gamma) \in \bar{Z}$ is regular if

1. the inequalities are satisfied strictly, i.e., $0 < l_{j,i} < u_{j,i}$, $0 < \gamma_i$, and $0 < \theta_{j,i}$,
2. for each fixed i the threshold values $\theta_{j,i}$ are distinct, and
3. for each $\kappa \in \mathcal{K}(z)$ the value $\Lambda_i(\kappa) \neq \gamma_i \theta_{j,i}$ for each threshold $\theta_{j,i}$ that defines κ .

We denote the collection of regular parameters by Z . Notice that \bar{Z} is (as the notation suggests) the topological closure of Z in \mathbb{R}^D and the set Z is generic in \bar{Z} . For a regular parameter $z \in Z$ the thresholds $\{\theta_{j,i} : j \in \mathbf{T}(i)\}$ occur in some definite (total) order $O_i(z)$ for each $i \in V$. We denote by $O(z) := (O_1(z), \dots, O_N(z))$ the collection of these orders over all nodes and the collection of all orderings $O(z)$ over all of parameter space as

$$O(Z) := \bigcup_{z \in Z} O(z).$$

2.2. Network node logics. Definition 2.4 does not specify the nonlinear functions M_j . As indicated in the introduction our approach is associated with the interpretation of regulatory networks as logical networks. To be more precise, a logical expression involving truth variables v_i , logical conjunctives \wedge (i.e. ANDs), and logical disjunctives \vee (i.e., ORs) leads to an analogous arithmetic expression by replacing \wedge with \cdot and \vee with $+$. For example, $(a \vee b) \wedge c$ becomes $(a + b)c$. Observe that given truth variables v_i , a logical expression $L(v_1, v_2, \dots, v_n)$ (without negations) leads unambiguously to the multilinear arithmetic expression $M(x_1, x_2, \dots, x_n)$ given by

$$M(x_1, x_2, \dots, x_n) := \sum_{L(v_1, \dots, v_n) = T} \prod_{v_i = F} (1 - x_i) \prod_{v_i = T} x_i,$$

where $x_i \in \mathbb{R}$.

Note that for every logical expression where each variable occurs *at most once* the recipe of replacing \wedge with \cdot and \vee with $+$ produces an arithmetic expression which is equivalent to this multilinear expression.

For the purposes of this paper our focus is on regulatory networks where one considers a *logic* for each network node j consisting of a logical expression involving each of the variables $x_i \in \mathbf{S}(j)$ precisely once. The multilinear functions M_j appearing in (4) are obtained from these logical expressions.

Remark 2.8. These assumptions on the structure of the terms in M_j imply that Λ_j can always be expressed in the form of a sum of monomials involving the step functions $\sigma_{j,i}(x)$ where the degree of $\sigma_{j,i}(x)$ is either zero or one (see also [5]), and thus this is a proper subset of the set of multilinear functions.

2.3. Cells and vector fields. If we restrict to a cell, then (3) reduces to a much simpler linear form. In particular, observe that given a cell κ , if $x, x' \in \kappa$, then $\Lambda(x) = \Lambda(x')$, and therefore $\Lambda(\kappa)$ is well-defined. In accordance with this observation we make the following definitions.

Definition 2.9. *A regulatory network \mathbf{RN} and a choice of parameter values $z \in Z$ leads to a uniquely defined switching system (1) and set of cells $\mathcal{K}(z)$. The κ -cell vector field for a cell $\kappa \in \mathcal{K}(z)$ is given by*

$$(6) \quad f^\kappa(x) := -\Gamma x + \Lambda(\kappa).$$

Observe that

$$\Phi(\kappa) := \Gamma^{-1}\Lambda(\kappa)$$

is a global attracting fixed point for (6). Accordingly we say that a cell κ is an attracting cell if

$$\Gamma^{-1}\Lambda(\kappa) \in \kappa.$$

3. State transition diagrams. As indicated in the introduction we capture the dynamics of (3) via *state transition diagrams*, directed graphs where the vertices correspond to regions of phase space, and the edges indicate how regions are related by the dynamics. To aid the reader the concepts of this section are illustrated in Figure 1 using a simple example of an \mathbf{RN} with two nodes and three edges (see Figure 1(a)). The associated switching system is shown in Figure 1(b) based on the logical expression $L(x_1, x_2) = x_1 \wedge x_2$. We begin in section 3.1 by defining wall-labelings that encapsulate combinatorial information derived from the κ -cell vector fields (6). In section 3.2 we give three different constructions of state transition diagrams. In section 3.3 we show that these three constructions are equivalent in the sense that they lead to equivalent dynamical information.

3.1. Wall-labelings. Faces of cells play a key role in our combinatorial representation of the dynamical system (3). However, each such face has two adjacent cells and the κ -cell vector fields on either side may differ. Accordingly we refine our concept of face to make reference to one of the adjacent cells.

Definition 3.1. *A wall is a pair (τ, κ) where κ is a cell and τ is a face of κ . Each wall inherits the projection and switching indexes from the corresponding face τ of κ . We say the sign of the wall (τ, κ) is 1 (and write $\text{sgn}(\tau, \kappa) = 1$) if τ is a left face of κ and we say the sign of the wall is -1 if τ is a right face of κ (and write $\text{sgn}(\tau, \kappa) = -1$). For a fixed parameter value z we denote the collection of walls by $\mathcal{W}(z)$.*

Observe that given a wall (τ, κ) there are three possibilities with respect to the κ -cell vector field f^κ : it is everywhere tangential to τ ; it points out of κ everywhere on τ ; or it points into κ everywhere on τ . If the projection index of τ is i , then these three cases are determined by the sign of the expression $f_i^\kappa(\tau)$ and whether the wall corresponds to a left or a right face (i.e., the sign of the wall), which in turn can be determined as a function of parameters. We summarize this in the following definition.

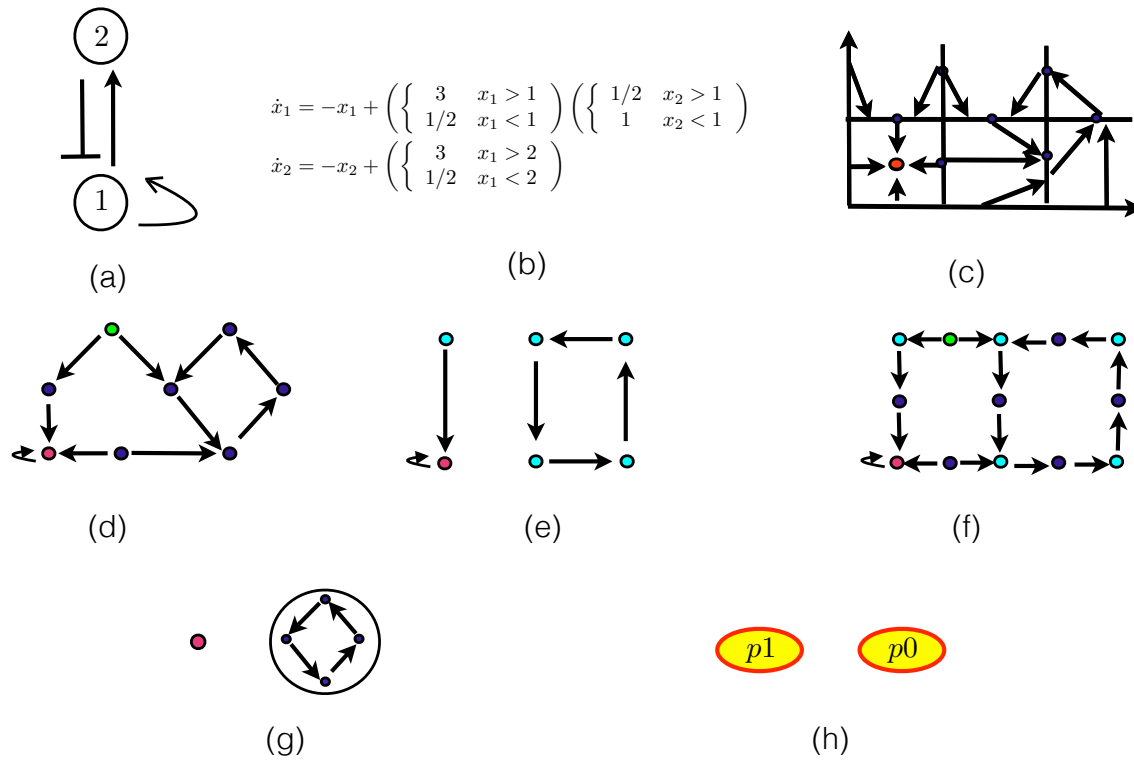


Figure 1. (a) Regulatory network RN; (b) a set of equations with a particular choice of parameters for RN; (c) phase space; (d) wall graph; (e) domain graph; (f) wall-domain graph; (g) a set of strongly connected components of the wall, domain, or wall-domain graph; (h) Morse graph representing strongly connected components.

Definition 3.2. Consider a switching network at a parameter value $z \in Z$. The wall-labeling of $\mathcal{W}(z)$ is the function $\ell : \mathcal{W}(z) \rightarrow \{-1, 0, 1\}$ defined as follows. Let $(\tau, \kappa) \in \mathcal{W}(z)$ have projection index i and switching index j . Then define

$$(7) \quad \ell((\tau, \kappa)) := \text{sgn}(\tau, \kappa) \cdot \text{sgn}(f_i^\kappa(\tau)) = \text{sgn}(\tau, \kappa) \cdot \text{sgn}(-\gamma_i \theta_{j,i} + \Lambda_i(\kappa)).$$

Note the last equality follows since if $x \in \tau$, then $x_i = \theta_{j,i}$.

Remark 3.3. As (7) makes clear the wall-labeling function depends explicitly on the choice of parameters for the switching network. However, given any two parameter values for which the ordering of the thresholds is the same there is an obvious identification of the cells and walls. Using this identification, the wall-labeling is completely determined by the values of $\text{sgn}(f_i^\kappa(\tau))$ over the collection of cells. As such we can define equivalence classes of parameter values over which wall-labelings are constant.

Definition 3.4. A wall (τ, κ) is an outgoing wall if $\ell(\tau, \kappa) = -1$, an incoming wall if $\ell(\tau, \kappa) = 1$, and a tangential wall if $\ell(\tau, \kappa) = 0$.

For parameters in the set Z (i.e., regular parameters) we will have only outgoing and incoming walls.

Proposition 3.5. *Given a switching system with parameter $z \in Z$, there are no tangential walls. That is, the wall-labeling function ℓ satisfies*

$$\ell(\tau, \kappa) \neq 0$$

for all $(\tau, \kappa) \in \mathcal{W}(z)$.

The classification of walls according to the value of the wall-labeling function arises from geometric considerations of the flows on the cells. We leave the proof of the following to the reader.

Proposition 3.6. *A cell κ is attracting if and only if every wall $(\tau, \kappa) \in \mathcal{W}$ is an incoming wall.*

In Figure 1(c) there are six cells, and a schematic representation of the direction of the flow in each cell is shown. The only attracting cell is the leftmost cell on the bottom, marked with a red dot in center.

A stronger relation between the labeling of walls and the dynamics of the κ -equation is as follows. Again the proof is left to the reader.

Proposition 3.7. *Let $x \in \kappa \in \mathcal{K}(z)$ where $z \in Z$ is a regular parameter value and let ψ_κ denote the flow generated by (6). If κ is an attracting cell, then there exists a unique time $t_x^- < 0$ such that $\psi_\kappa((t_x^-, \infty), x) \subset \kappa$ and $\psi_\kappa(t_x^-, x) \in \bar{\tau}$ where (τ, κ) is an incoming wall and $\bar{\tau}$ denotes closure of τ . If κ is not an attracting cell, then there exist unique times $t_x^- < 0 < t_x^+$ such that $\psi_\kappa((t_x^-, t_x^+), x) \subset \kappa$, $\psi_\kappa(t_x^-, x) \in \bar{\tau}$ where (τ, κ) is an incoming wall, and $\psi_\kappa(t_x^+, x) \in \bar{\tau}'$ where (τ', κ) is an outgoing wall.*

3.2. State transition diagram constructions. Recall that a state transition diagram is a directed graph. To emphasize that we employ this as a means of representing information about dynamics we adopt an equivalent perspective: a state transition diagram is a combinatorial multivalued map $\mathcal{F}: \mathcal{V} \rightrightarrows \mathcal{V}$ (where \mathcal{V} is the collection of vertices) such that $\nu' \in \mathcal{F}(\nu)$ if and only if there is a directed edge $\nu \rightarrow \nu'$ in the state transition diagram. Using the multivalued map notation the existence of a path from ν to ν' is expressed by $\nu' \in \mathcal{F}^k(\nu)$ for some positive integer k .

Let ℓ be the wall-labeling for a switching network at a fixed parameter. We give three constructions for state transition diagrams. In each case, note that the state transition diagrams depends only on ℓ , and hence it is through the wall-labelling that the state transition diagrams inherit their dependence on parameters. In fact, as indicated by Remark 3.3, this inheritance remains constant on equivalence classes of parameter values.

Definition 3.8. *The wall graph $\mathcal{F}: \mathcal{V} \rightrightarrows \mathcal{V}$ induced by a fixed wall-labeling is defined as follows. The set of vertices \mathcal{V} is in 1-1 correspondence with the set of attracting cells and faces of all cells. For each pair of faces τ, τ' admitting a cell κ such that (τ, κ) is an incoming wall and (τ', κ) is an outgoing wall, there is an edge $\tau \rightarrow \tau'$, or equivalently $\tau' \in \mathcal{F}(\tau)$. For each attracting domain κ , $\kappa \in \mathcal{F}(\kappa)$ and $\kappa \in \mathcal{F}(\tau)$ for each wall (τ, κ) .*

Remark 3.9. A consequence of the assumption that the regulatory network does not have any direct negative self-regulation (see Remark 2.2) is that every node in the wall graph has an out-edge.

The wall graph for the system in Figure 1(b) is shown in Figure 1(d). Each dark vertex corresponds to a transparent wall, while the green vertex corresponds to a white wall. The red vertex with a self-edge represents an attracting cell.

Definition 3.10. The domain graph $\mathcal{F}: \mathcal{V} \rightrightarrows \mathcal{V}$ induced by a fixed wall-labeling ℓ is defined as follows. The set of vertices \mathcal{V} is in 1-1 correspondence with the collection of fundamental domains $\mathcal{K}(\kappa)$. If some cell κ is an attracting domain, then $\kappa \in \mathcal{F}(\kappa)$. Furthermore, $\kappa' \in \mathcal{F}(\kappa)$ whenever there exists a face τ such that (τ, κ) and (τ, κ') are walls such that $\ell((\tau, \kappa)) = -1$ (indicating an outgoing wall of κ) and $\ell((\tau, \kappa')) = 1$ (indicating an incoming wall of κ').

The domain graph for the system in Figure 1(b) is shown in Figure 1(e). Again, the red vertex with a self-edge represents an attracting cell and teal vertices are nonattracting cells.

Definition 3.11. The wall-domain graph $\mathcal{F}: \mathcal{V} \rightrightarrows \mathcal{V}$ induced by a given wall-labeling is defined as follows. The set of vertices \mathcal{V} is in 1-1 correspondence with the collection of all cells and the faces of all cells. There are three types of edges. If (τ, κ) is an outgoing wall, then $\tau \in \mathcal{F}(\kappa)$. If (τ, κ) is an incoming wall, then $\kappa \in \mathcal{F}(\tau)$. Finally, if κ is an attracting cell, then $\kappa \in \mathcal{F}(\kappa)$.

The wall-domain graph for the system in Figure 1(b) is shown in Figure 1(f). The vertices of this graph are the union of vertices of the wall graph (Figure 1(d)) and the domain graph (Figure 1(e)) and their color matches their color in the original graphs. Except for the self-edge on the vertex corresponding to the attracting cell the graph is bipartite; edges connect wall vertices to domain vertices and domain vertices to wall vertices.

3.3. Dynamical signatures. As indicated in the introduction, to store the dynamics we make use of a more compact representation.

Definition 3.12. A recurrent component (also referred to as a strongly connected path component) of a directed graph \mathcal{F} is a maximal collection \mathcal{C} of vertices such that for any $u, v \in \mathcal{C}$ there exists a nonempty path from u to v in \mathcal{C} . In the context of dynamical systems we refer to a recurrent component of \mathcal{F} as a Morse set of \mathcal{F} and denote it by $\mathcal{M} \subset \mathcal{V}$. The collection of all recurrent components of \mathcal{F} is denoted by

$$\text{MD}(\mathcal{F}) := \{\mathcal{M}(p) \subset \mathcal{V} \mid p \in \mathbf{P}\}$$

and is called a Morse decomposition of \mathcal{F} . Here \mathbf{P} is an index set. Recurrent components inherit a well-defined partial order by the reachability relation in the directed graph \mathcal{F} . Specifically, we may write the partial order on the indexing set \mathbf{P} of $\text{MD}(\mathcal{F})$ by defining

$$q \leq p \quad \text{if there exists a path in } \mathcal{F} \text{ from an element of } \mathcal{M}(p) \text{ to an element of } \mathcal{M}(q).$$

Primarily for clarity we note the following facts regarding recurrent components.

Proposition 3.13. Two elements $\nu, \nu' \in \mathcal{V}$ belong to the same recurrent component of \mathcal{F} if and only if there exist positive integers k, k' such that $\nu' \in \mathcal{F}^k(\nu)$ and $\nu \in \mathcal{F}^{k'}(\nu')$. Distinct recurrent components are disjoint. Not every vertex need belong to some recurrent component. Recurrent components are strongly connected components. The only strongly connected components that are not recurrent components are singleton sets that do not have a self-edge.

Definition 3.14. The Morse graph of \mathcal{F} , denoted $\text{MG}(\mathcal{F})$, is the Hasse diagram of the poset (\mathbf{P}, \leq) . We refer to the elements of \mathbf{P} as the Morse nodes of the graph.

Figure 1(g) shows the strongly connected components of either of the three graphs (wall graph, domain graph, and wall-domain graph) in Figures 1(d), (e), and (f), respectively. The Morse graph representation of the strongly connected components is in Figures 1(g) and (h). The two Morse sets are not comparable and so the partial order P is empty.

We note that the computation of Morse graphs is feasible and can be accomplished via well-known algorithms for strongly connected components [43] and transitive reduction [1]. In particular, for a directed graph with E vertices and V edges, we may compute strongly connected components in $O(E)$ time and the reachability relation among strongly connected components in $O(EV)$ time. In our setting, the number of edges is proportional to the number of vertices (i.e., $E < 2^d V$). For the worst case of $O(V)$ recurrent components this results in quadratic performance. However, if the number of recurrent components is small (as we frequently observe) this becomes a linear time procedure.

Proposition 3.15. *The Morse graphs induced by the wall graph, the wall-domain graph, and the domain graph are isomorphic.*

Proof. In the wall graph, every incoming face maps to every outgoing face. Meanwhile in the wall-domain graph every incoming face maps to the cell, which then maps to each of its exit faces. It follows that the reachability between faces in the wall graph and the wall-domain graph is the same. What is more, every recurrent component in the wall graph either contains a face or else is an attracting domain. From this one may establish the isomorphism between the Morse graph of a wall graph and the Morse graph of the wall-domain graph. Establishing the isomorphism of Morse graphs induced by the wall-domain graph and the domain graph is similar: we observe that the reachability between fundamental domains is the same in either and that each recurrent component must contain at least one domain. Combining these two isomorphisms gives the isomorphism from wall graph to domain graph. ■

In light of the previous result and the simplicity of the domain graph one might wonder why we should bother with the wall graph or wall-domain graph at all. The answer is twofold. First, it is possible to refine our analysis so that in the wall graph not every incoming face (τ, κ) is mapped to every outgoing face (τ', κ) for a cell κ . For example, consider Figure 2. In (a), we see the wall graph obtained in the present work in the situation of a two-dimensional cell with two incoming and two exit faces arranged as indicated. However, for a particular parameter the actual trajectories would be captured to the diagrams sketched in (b) and (c). To achieve combinatorial descriptions capturing the additional information in (b) or (c) the domain graph is inadequate and it will be necessary to use notions such as the wall graph. We leave this to future work.

The idea of using symbolic representation of the dynamics of switching systems motivated the introduction of these systems in the 1970s [27, 28]. Our concept of a domain graph is analogous to the state transition diagram in these papers. The representation of dynamics by maps between walls, in a different context, has been used before [29, 17, 21, 16, 24].

Second, via more sophisticated constructions of the directed graph we can obtain information about unstable dynamics from the Morse graph. These constructions involve using lower-dimensional cells such as faces as vertices and will be detailed in a future work.

3.4. Annotation of Morse nodes. In addition to recording the Morse graph for a parameter $z \in Z$ it is possible to produce extra information in the form of annotations we associate

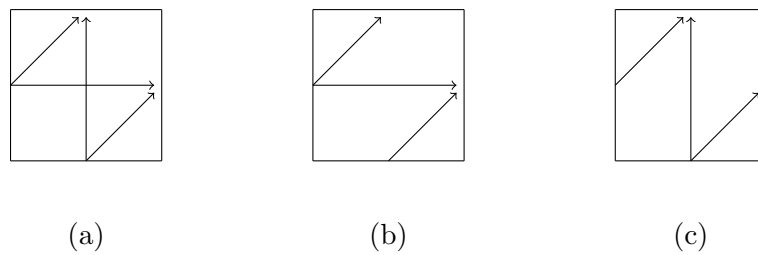


Figure 2. (a) Coarse wall graph where all incoming faces map to all exit faces. (b) Finer wall graph with a disallowed incoming-exit arrow from bottom to top. (c) Finer wall graph with disallowed arrow from left to right.

with the Morse nodes of the Morse graph. We refer to this information, Morse graphs plus annotations, as a *dynamical signature*. Presently, we compute annotations for each Morse node based on the vertices present in the associated Morse set. We briefly describe these annotations. First, we say that a Morse set *makes an x_d transition* if it contains vertices corresponding to cells with differing x_d coordinates. We make annotations according to the set of transitions. In the simplest case, there are no transitions (we have only a single attracting cell) and we annotate the Morse set as a fixed point but respecting three subcases: (a) if the cell is located to the left of each threshold in all dimensions the Morse node is marked FP OFF (*fixed point off*); (b) if the cell is located to the right of at least one threshold in each dimension it is marked FP ON (*fixed point on*); (c) otherwise it is marked just FP. In the other extreme case every transition is made (i.e., x_1, x_2, \dots, x_N). In this case we annotate the Morse node FC (for *full cycle*). Otherwise we annotate the Morse node according to the subset of variables for which there is a transition.

4. Parameter graph. The content of sections 2 and 3 implies that given a regulatory network RN and a parameter $z \in Z$ for the associated switching system (3) it is possible to create an annotated Morse graph.

Since two parameters $z, z' \in Z$ give rise to the same annotated Morse graph provided they give rise to the same wall-labeling (up to the equivalence indicated in Remark 3.3) it is natural to discretize parameter space according to regions that are guaranteed to give the same wall-labeling. How to accomplish this is the topic of this section. As indicated in the introduction, the final results are parameter graphs that are defined below.

The nodes of a parameter graph, which we denote by \mathcal{Z} , are meant to correspond to regions in parameter space and the edges are meant to indicate geometric relations between the regions. The resulting *database of dynamic signatures* can be viewed as a map

$$DB : \mathcal{Z} \rightarrow \text{AnnMG},$$

where AnnMG denotes the collection of annotated Morse graphs. As indicated in section 5, in applications queries to the database are often concerned with finding the set of nodes which correspond to a particular annotated Morse graph.

We remark that the parameter space $Z \subset [0, \infty)^D$, as defined in Definition 2.7, is a semi-algebraic set, i.e., it is expressed in terms of polynomial inequalities. To see this note that by

Definition 2.3, \bar{Z} is a semialgebraic set and Z is the complement in \bar{Z} of the set of parameters that satisfy any one of the equalities

$$(8) \quad \begin{aligned} 0 &= l_{j,i} - u_{j,i}, \\ 0 &= \gamma_i, 0 = \theta_{j,i}, 0 = l_{j,i}, 0 = u_{j,i}, \\ 0 &= \theta_{j,i} - \theta_{j',i} \quad \text{for distinct } j, j' \in \mathbf{T}(i), \text{ or} \\ 0 &= \gamma_i \theta_{j,i} - \Lambda_i(\kappa), \quad \text{where } \theta_{j,i} \text{ defines a face of } \kappa, \end{aligned}$$

where $\kappa \in \mathcal{K}(z)$ and by Remark 2.8 $\Lambda_i(\kappa)$ can be expressed as a sum of monomials involving parameters.

We propose two conceptually different means of identifying Z with \mathcal{Z} . In section 4.1 we describe an approach based on topological considerations; the basic elements are connected components of Z and their adjacency structure is defined in terms of closures of these sets. This gives rise to the GPG.

In section 4.2 we describe an approach that is explicitly computable; we consider subsets of Z described by systems of strict inequalities. We define an adjacency structure by considering reversing the direction of these inequalities one at a time. This gives rise to the CPG.

In section 4.3 we present a decomposition of the geometric and combinatorial parameter graphs as a product of smaller graphs in a way which corresponds to the structure of the regulatory network. In section 4.5 we compare the GPG and CPG graphs. There is a canonical graph homomorphism $h : \text{GPG} \rightarrow \text{CPG}$ induced from inclusion, but it is an open question whether this homomorphism is in fact always an isomorphism (i.e., we have not proven in general that these two approaches are equivalent). We are, however, able to give a class of networks for which GPG and CPG are known to be the same. Finally, in section 4.4 we will discuss the computation of CPG.

4.1. Geometric parameter graph. Because we are using the domain graph the state transition diagram $\mathcal{F}(\cdot, z) : \mathcal{V} \rightrightarrows \mathcal{V}$ is defined for each $z \in Z$ and is completely determined by the wall-labeling (7). The ordering O and the wall-labeling are constant on connected components of Z . This motivates a combinatorialization of parameter space via the connected components, which we label by \mathcal{Z} . With this in mind we define the *parameter nodes* of the GPG to be \mathcal{Z} . To complete the definition of the GPG we require a notion of adjacency for parameter nodes.

Definition 4.1. *We say that a parameter value $z \in \bar{Z}$ is k -deficient if exactly k of the equalities of (8) are satisfied. Given a switching network (1) the associated GPG has vertices \mathcal{Z} and edges (ζ, ζ') if there exists $z \in \text{cl}(\zeta) \cap \text{cl}(\zeta')$ such that z is 1-deficient. Here the closures $\text{cl}(\zeta)$ and $\text{cl}(\zeta')$ are taken in \bar{Z} .*

Open Question 4.2. *Does the concept of k -deficient parameter values naturally generate a regular CW-decomposition of \bar{Z} where the elements of \mathcal{Z} represent the cells of dimension D ?*

Adjacency in the GPG is meant to correspond to cells that share a $D - 1$ dimensional face. Extending this construction would provide a means of understanding the topology of regions of parameter space that are associated with specified dynamic phenotypes.

The following example illustrates the concept of the GPG.

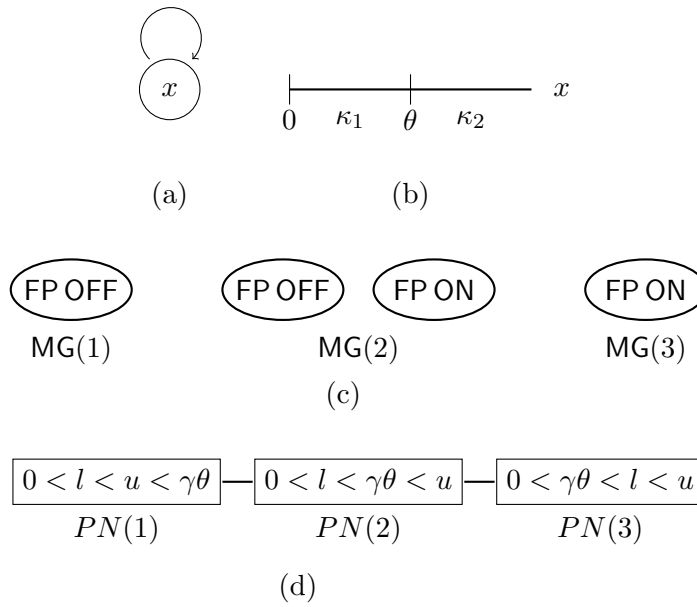


Figure 3. (a) A self-activating one-node network. (b) Phase plane for switching network. (c) Annotated Morse graphs: MG(1) has a single node generated by an attracting cell for which the fixed point is less than the threshold and annotated by FP OFF; MG(3) has a single node generated by an attracting cell for which the fixed point is greater than the threshold and annotated by FP ON; and MG(2) has two minimal nodes generated by attracting cells in one of which the fixed point is less than the threshold and in the other the fixed point is greater than the threshold. (d) Parameter graph.

Example 4.3. Consider the simplest regulatory network $\mathbf{RN} = (V, E)$, where $V = \{1\}$, $1 \rightarrow 1$, and $M_1(x) = x$. The regulatory network is shown in Figure 3(a). Since the $1 \rightarrow 1$ edge is activating, the associated switching network takes the form

$$(9) \quad \dot{x} = -\gamma x + \begin{cases} l & \text{if } x < \theta, \\ u & \text{if } x > \theta. \end{cases}$$

The associated phase space and subdivision are shown in Figure 3(b).

In particular, there are two cells $\kappa_1 := (0, \theta)$ and $\kappa_2 := (\theta, \infty)$. The dimension of the associated parameter space is $D = 1 + 3 = 4$ and

$$Z = \{(l, u, \theta, \gamma) \mid l < u, \gamma\theta \notin \{l, u\}\} \subset [0, \infty)^4.$$

Because of the simplicity of the problem, it is easy to enumerate all the state transition diagrams and compute the associated Morse graphs. Figure 3(c) indicates the annotated Morse graphs. Without the annotations Morse graphs MG(1) and MG(3) are equivalent and consist of a single node. However, in the process of the computation we can identify that the fixed point in κ_1 is less than the threshold; thus the node x is in an “off” state. We denote this information by FP OFF. The annotated Morse graph includes this information at the node. Similarly, the annotated Morse graph MG(3) indicates that the fixed point is in κ_2 . We denote this information by FP ON. The Morse graph MG(2) has two minimal nodes

generated by two attracting cells in one of which the fixed point is less than the threshold and in the other the fixed point is greater than the threshold. Again these fixed points are annotated FP OFF and FP ON, respectively.

The associated regions of parameter space, i.e., their defining inequalities, are indicated in Figure 3(d). Observe that there is a straight line in $PN(i)$ from any point in $PN(i)$ to the origin. Thus one can show that each region $PN(i)$ is connected. Thus, the GPG contains three nodes. Observe that if $z \in \bar{Z}$ satisfies $\gamma\theta - u = 0$, then $z \in \text{cl}(PN(1)) \cap \text{cl}(PN(2))$, and similarly, if $z \in \bar{Z}$ satisfies $\gamma\theta - l = 0$, then $z \in \text{cl}(PN(2)) \cap \text{cl}(PN(3))$. Thus, we have an edge between the nodes corresponding to $PN(1)$ and $PN(2)$ and an edge between the nodes corresponding to $PN(2)$ and $PN(3)$. Therefore, Figure 3(d) is the GPG for this example.

How to extend these arguments in Example 4.3 to more complicated regulatory networks is not obvious. However, consider $Z' := \{(l, u, \theta) \in [0, \infty)^3 \mid (l, u, \theta, 1) \in Z\} \subset [0, \infty)^3$ and observe that characterizing Z' is equivalent to characterizing the complement of the degenerate finite hyperplane arrangement [41]

$$\{\{v \in [0, \infty)^3 \mid a \cdot v = 0\} \mid a \in A\},$$

where

$$A = \{(1, 0, 0), (0, 1, 0), (0, 0, 1), (1, -1, 0), (1, 0, -1), (0, 1, -1)\}.$$

In this setting it is fairly easy to determine that Z' consists of three unbounded connected components for which the origin is contained in their closures. However, we are interested in characterizing all of Z , which implies that we need to consider complements of the nonlinear equations $-\gamma\theta + l = 0$ and $-\gamma\theta + u = 0$. For a general regulatory network the dimension D grows linearly in the number of vertices and edges and the terms of Λ , while multilinear, may consist of higher dimensional products of the parameters. This implies that the problem of understanding Z is at least as challenging as that of determining the cells in a degenerate finite hyperplane arrangement. To deal with these complications, in the next subsections we turn to techniques from computational algebraic geometry. For now we content ourselves with the following result.

Proposition 4.4. *The GPG is a connected graph.*

Proof. Our proof is based on finding a straight line path between points in any two parameter nodes from which we can prove the existence of a corresponding path in the GPG. The burden is to show we can avoid pathologies where the straight line path (a) passes through an accumulation of parameter nodes or (b) passes from one parameter node to another through a point which is not 1-deficient. Since the parameter nodes are all in the strictly positive orthant, we only need to consider deficiencies arising from one of the following equalities being satisfied (the other equalities of (8) do not intersect the strictly positive orthant and need not be considered):

$$(10) \quad \begin{aligned} 0 &= l_{j,i} - u_{j,i}, \\ 0 &= \theta_{j,i} - \theta_{j',i} \quad \text{for distinct } j, j' \in \mathbf{T}(i), \text{ or} \\ 0 &= \gamma_i \theta_{j,i} - \Lambda_i(\kappa) \quad \text{where } \theta_{j,i} \text{ defines a face of } \kappa. \end{aligned}$$

Each of the equalities in (10) has a solution set in $(0, \infty)^D$ (for $D := N + 3 \cdot \#(E)$) which we show is a codimension-1 submanifold. To see this, let $f_1, f_2, \dots, f_k : (0, \infty)^D \rightarrow \mathbb{R}$ be the functions such that the equalities of (10) may be written $f_i = 0$ for $i = 1, 2, \dots, k$, where k is the number of equalities in (10). We leave it to the reader to inspect (10) and see that the gradient of each f_i (for each $i = 1, 2, \dots, k$) is nonvanishing, i.e., 0 is a regular value of f_i . By the regular value theorem, the varieties $f_i^{-1}(0)$ are thus codimension-1 submanifolds S_i of $(0, \infty)^D$ for $i = 1, 2, \dots, k$.

We consider a collection of line segments $\mathcal{L} = \mathcal{L}_a \cup \mathcal{L}_b$, where \mathcal{L}_a denotes the collection of line segments which intersect $\bigcup_{i=1}^k S_i$ infinitely often, and \mathcal{L}_b denotes the collection of line segments which intersect $\bigcup_{i \neq j} S_i \cap S_j$ (i.e., any two of the submanifolds at the same point). Describing line segments by their endpoints we regard \mathcal{L} as a subset of $(0, \infty)^{2D}$. We make the following technical claim: \mathcal{L} is a set of zero measure. We show this by proving that both \mathcal{L}_a and \mathcal{L}_b have zero measure. Note this corresponds to showing the pathologies (a) and (b) are rare.

To see that \mathcal{L}_a has zero measure, observe first that the union of finitely many codimension-1 submanifolds $\bigcup_{i=1}^k S_i$ has measure zero. It follows that the collection of line segments with either endpoint in $\bigcup_{i=1}^k S_i$ has measure zero. For any line segment with endpoints p and q which are both not in $\bigcup_{i=1}^k S_i$, we consider its parameterization $s : [0, 1] \rightarrow (0, \infty)^D$ defined by $s(t) := (1-t)p + tq$. The compositions $f_i(s(t))$ are polynomials in t which are not identically zero. By the Fundamental Theorem of Algebra they each have a finite number of zeroes. Since the zeros correspond to the intersections of the segment with the submanifolds, the segments are not in \mathcal{L}_a . It follows that \mathcal{L}_a has measure zero.

To see that \mathcal{L}_b has zero measure, define $f_{ij} : (0, \infty)^D \rightarrow \mathbb{R}^2$ via $f_{ij}(x) := (f_i(x), f_j(x))$ for $i \neq j$. We leave it to the reader to inspect (10) and verify that since for $i \neq j$ the set of variables appearing in the expressions for $f_i(x)$ and $f_j(x)$ are not the same, the gradients $\nabla f_i|_p$ and $\nabla f_j|_p$ are linearly independent for any $p \in (0, \infty)^D$, i.e., the Jacobian $Df_{ij}|_p$ has full rank. Thus 0 is a regular value of f_{ij} . By the regular value theorem, $S_i \cap S_j = f_{ij}^{-1}(0)$ is a codimension-2 submanifold of $(0, \infty)^D$. It follows that the set of line segments intersecting $S_i \cap S_j$ for some $i \neq j$ is measure zero. Hence \mathcal{L}_b has measure zero.

Consequently, we may always perturb a line segment by making arbitrarily small adjustments to the location of its endpoints so that it intersects the submanifolds defined by (8) finitely often and never intersects two submanifolds at the same point. Now choose $\zeta, \zeta' \in \text{GPG}$ and let $p : [0, 1] \rightarrow \bar{Z}$ be a straight line path such that $p(0) \in \zeta$ and $p(1) \in \zeta'$. Since ζ and ζ' are open, there exist neighborhoods in which we may perturb $p(0) \in \zeta$ and $p(1) \in \zeta'$; we use this freedom so we can assume without loss of generality that the line segment $p([0, 1])$ satisfies this technical property. By the first technical condition $p(t) \in Z$ for all but finitely many $t \in [0, 1]$ there exists a finite sequence $(\zeta_0 = \zeta, \zeta_1, \zeta_2, \dots, \zeta_n = \zeta')$ through GPG corresponding to a finite sequence of intervals $[t_0 = 0, t_1), (t_1, t_2), \dots, (t_n, t_{n+1} = 1]$ in $[0, 1]$ for which $p((t_i, t_{i+1})) = \zeta_n$. By the second technical condition, $p(t)$ intersects only one submanifold at a time, hence for each $i = 1, \dots, n$ the point $p(t_i)$ is a 1-deficient point in the intersection of the closures of ζ_{i-1} and ζ_i . These facts together yield a path $\zeta = \zeta_0 \rightarrow \zeta_1 \rightarrow \zeta_2 \rightarrow \dots \rightarrow \zeta_n = \zeta'$. Hence GPG is connected. ■

4.2. Combinatorial parameter graph. In this section we show how to assign to each parameter $z \in Z$ a combinatorial description ϕ which is sufficient to construct the wall-labeling (and hence state transition diagrams) induced by z . We call this combinatorial description a *combinatorial parameter* and we denote the collection of combinatorial parameters by Φ . Importantly, we show that for every parameter node $\zeta \in Z$, for any $z, z' \in \zeta$, both z and z' have the same associated combinatorial parameter $\phi \in \Phi$.

Our definition of combinatorial parameters is unfortunately somewhat technical. Overall it amounts to a bookkeeping system to keep track of the directions of the inequalities involving parameters which determine the threshold order and wall-labeling function. The threshold orderings give rise to inequalities comparing θ parameters. To determine the wall-labeling function requires comparing the various Λ_i to $\gamma_i \theta_{j,i}$ (which gives the signs in (7)). For a given regulatory network a fixed number of such comparisons are required; our definition of a combinatorial parameter provides an organizational framework to speak of this collection of inequalities in a rigorous way.

Definition 4.5. Define the input combinations of the node x_i to be the Cartesian product

$$\text{In}_j := \prod_{i \in \mathbf{S}(j)} \{\text{off}, \text{on}\}.$$

Define the indicator function $\chi_j : (0, \infty)^N \rightarrow \text{In}_j$ such that

$$\chi_{j,i}(x) = \begin{cases} \text{off} & \text{if } i \rightarrow j \text{ and } x_i < \theta_{j,i} \text{ or if } i \dashv j \text{ and } x_i > \theta_{j,i}, \\ \text{on} & \text{if } i \rightarrow j \text{ and } x_i > \theta_{j,i} \text{ or if } i \dashv j \text{ and } x_i < \theta_{j,i}, \\ \text{undefined} & \text{otherwise.} \end{cases}$$

Define the valuation function $v_j : \text{In}_j \rightarrow \mathbb{R}^{\mathbf{S}^j}$ via

$$v_{j,i}(A) = \begin{cases} l_{j,i} & \text{whenever } A_i = \text{off}, \\ u_{j,i} & \text{whenever } A_i = \text{on}, \\ \text{undefined} & \text{otherwise.} \end{cases}$$

Note that $\sigma_j = v_j \circ \chi_j$, where σ_j is defined in (5).

Define the output combinations of the node i to be the set

$$\text{Out}_i := \mathbf{T}(i),$$

where we recall from Definition 2.1 that $\mathbf{T}(i)$ is the set of target nodes of i in the regulatory network \mathbf{RN} .

Definition 4.6. A logic parameter is a function

$$L : \prod_{i=1}^N (\text{In}_i \times \text{Out}_i) \rightarrow \{-1, 1\}.$$

We denote the restriction of L onto $\text{In}_i \times \text{Out}_i$ as L_i . An order parameter O is a collection of total orderings O_i of $\mathbf{T}(i)$ for each $i \in X$. A combinatorial parameter is a pair $\phi = (L, O)$,

where L is a logic parameter and O is an order parameter. We denote the collection of combinatorial parameters as Φ . The combinatorial assignment function $\omega : Z \rightarrow \Phi$ is given by $\omega(z) := (L, O)$, where $O = O(z)$ and

$$(11) \quad L_i(A, B) = \text{sgn}(M_i \circ v_i(A) - \gamma_i \theta_{B,i}) \text{ for all } 1 \leq i \leq N.$$

For all $z \in Z$, we say that $\omega(z)$ is the combinatorial parameter associated to the parameter z .

The parameter region associated with the combinatorial parameter ϕ is given by $\omega^{-1}(\phi) \subset Z$ and denoted by $|\phi|$. A combinatorial parameter $\phi \in \Phi$ is realizable if there exists $z \in Z$ such that $\phi = \omega(z)$.

Definition 4.7. Let $\phi = (L, O) \in \Phi$ be a combinatorial parameter. Suppose $z \in Z$ such that $O = O(z)$. We may induce a wall-labeling on $\mathcal{W}(z)$ as follows. Let (τ, κ) be a wall with projection index i and switching index j . We say (τ, κ) is an outgoing wall with respect to ϕ if $L_i(\chi_i(\kappa), j) = -\text{sgn}((\tau, \kappa))$ and an incoming wall if $L_i(\chi_i(\kappa), j) = \text{sgn}((\tau, \kappa))$.

The next result shows we have successfully given a combinatorial description $\phi = \omega(z)$ for each $z \in Z$.

Proposition 4.8. Let $z \in Z$. Define $\phi \in \Phi$ such that $\phi = \omega(z)$. Then the wall-labeling induced by z and the wall-labeling induced by ϕ are the same.

Proof. It suffices to show ϕ and z induce the same wall-labeling, which then in turn will induce the same wall graph. To this end it suffices to show for each wall (τ, κ) with projection index i and switching index j that (1) $L_i(\chi_i(\kappa), j) = +1$ is equivalent to $M_i(\sigma_i(\kappa)) > \gamma_i \theta_{j,i}$, and (2) $L_i(\sigma_i(\kappa), j) = -1$ is equivalent to $M_i(\sigma_i(\kappa)) < \gamma_i \theta_{j,i}$. This follows from the definitions and (11). ■

We supplement combinatorial parameters with a notion of adjacency. To understand this notion of adjacency it helps to remember that a combinatorial parameter is nothing more than a bookkeeping system giving the direction of the inequality for a set of inequalities describing a region of parameter space. Our notion of adjacency corresponds to reversing a single one of these inequalities.

Definition 4.9. Let $\phi = (L, O)$, $\phi' = (L', O') \in \Phi$ be distinct combinatorial parameters. Denote the domains of L and L' by $\mathcal{D} := \bigsqcup_{i=1}^N (\text{In}_i \times \text{Out}_i)$. We say ϕ and ϕ' are adjacent if either (1) $O = O'$ and there exists $x \in \mathcal{D}$ such that ϕ and ϕ' are equal on all of \mathcal{D} except x , or (2) $L = L'$, $O_i = O'_i$ for $i \in V \setminus i^*$, and the total orders O_{i^*} and O'_{i^*} differ only by a single swap of the ordering of consecutive thresholds.

The previous definition renders Φ into a large graph of combinatorial parameters. Since not every combinatorial parameter is realizable (i.e., in the image of ω) we are interested only in a subgraph.

Definition 4.10. The CPG is the undirected graph on the realizable combinatorial parameters with an edge between two parameter nodes ϕ and ϕ' if and only if they are adjacent in the sense of Definition 4.9.

Example 4.11. To illustrate the idea of the combinatorial parameter graph, we return to the one-node regulatory network, indicated in Figure 3(a), of Example 4.3. Recall that the

switching network is (9), the phase space is indicated in Figure 3(b), and Figure 3(c) indicates the annotated Morse graphs. For the purposes of this example it is instructive to construct the collection of combinatorial parameters Φ and see which ones are realizable. There is only a single node, so

$$\text{In} = \{\text{off}, \text{on}\} \quad \text{and} \quad \text{Out} = \{1\}$$

and the order parameter is similarly trivial $O = \{\theta\}$. The set of logic parameters is

$$(12) \quad (\text{off}, 1) \mapsto -1 \quad \text{and} \quad (\text{on}, 1) \mapsto -1,$$

$$(13) \quad (\text{off}, 1) \mapsto 1 \quad \text{and} \quad (\text{on}, 1) \mapsto -1,$$

$$(14) \quad (\text{off}, 1) \mapsto -1 \quad \text{and} \quad (\text{on}, 1) \mapsto 1,$$

$$(15) \quad (\text{off}, 1) \mapsto 1 \quad \text{and} \quad (\text{on}, 1) \mapsto 1,$$

and therefore Φ contains four elements.

To determine the realizable combinatorial parameters we note that the indicator function $\chi: (0, \infty) \rightarrow \text{In}$ is

$$\chi(x) = \begin{cases} \text{off} & \text{if } x < \theta, \\ \text{on} & \text{if } \theta < x, \\ \text{undefined} & \text{otherwise} \end{cases}$$

and the valuation function $v: \text{In} \rightarrow \mathbb{R}$ is

$$v(A) = \begin{cases} l & \text{if } A = \text{off}, \\ u & \text{if } A = \text{on}. \end{cases}$$

Note that M is the identity and thus to understand the image of the combinatorial assignment function $\omega: Z \rightarrow \Phi$ we only need to consider

$$L(A, B) = \text{sgn}(v(A) - \gamma\theta).$$

Observe that if $A = \text{off}$, then

$$L(\text{off}, 1) = \text{sgn}(l - \gamma\theta) = \begin{cases} -1 & \text{if } l < \gamma\theta, \\ 1 & \text{if } \gamma\theta < l, \end{cases}$$

and if $A = \text{on}$, then

$$L(\text{on}, 1) = \text{sgn}(u - \gamma\theta) = \begin{cases} -1 & \text{if } u < \gamma\theta, \\ 1 & \text{if } \gamma\theta < u. \end{cases}$$

Observe that if $l < u < \gamma\theta$, $l < \gamma\theta < u$, and $\gamma\theta < l < u$, then we obtain the combinatorial parameters with logic parameters (12), (14), and (15), respectively. Notice that the combinatorial parameter with logic parameter (13) is not realizable since it would require $\gamma\theta < l$ and $u < \gamma\theta$, which contradicts $l < u$. Thus CPG has three nodes. The edges are as indicated in Figure 3(d), since these correspond to the switching of a single inequality. Thus, as promised (see Theorem 4.18) the CPG agrees with the GPG of Example 4.3.

4.3. Product structure of parameter graph. Both GPG and CPG have a product structure. To describe this structure we need to define what is meant by a product of graphs.

Definition 4.12. Given a collection of graphs $\{G_\alpha\}_{\alpha \in J}$ the graph product $\prod_{\alpha \in J} G_\alpha$ is the graph with nodes which are J -tuples x such that $x_\alpha \in G_\alpha$ for $\alpha \in J$ and two J -tuples x and y are adjacent if and only if $x_\alpha = y_\alpha$ for all but possibly one exceptional value $\alpha^* \in J$, and for that exceptional α^* , x_{α^*} , and y_{α^*} are adjacent in G_{α^*} .

Definition 4.13. For each $i \in V$, define the geometric factor graph GPG_i to be the connected components of the complement of the solutions of the equalities

$$\begin{aligned} 0 &= l_{j,i} - u_{j,i}, \\ 0 &= \gamma_i, 0 = \theta_{j,i}, 0 = l_{j,i}, 0 = u_{j,i}, \\ 0 &= \theta_{j,i} - \theta_{j',i} \quad \text{for distinct } j, j' \in \mathbf{T}(i), \text{ or} \\ 0 &= \gamma_i \theta_{j,i} - \Lambda_i(\kappa), \quad \text{where } \theta_{j,i} \text{ defines a face of } \kappa, \end{aligned}$$

in

$$Z_i := \{(l, u, \theta, \gamma) \in [0, \infty)^{1+3\#\mathbf{T}(i)}\}.$$

Two connected components are considered to be adjacent if they admit a 1-deficient point in the intersection of their closures.

Definition 4.14. Let $i \in V$. Define Φ_i to be the collection of pairs (L_i, O_i) , where O_i is an ordering of $\mathbf{T}(i)$ and L_i is a function

$$L_i : \text{In}_i \times \text{Out}_i \rightarrow \{-1, 1\}.$$

Define a function $\omega_i : Z_i \rightarrow \Phi_i$ via $\omega(z) := \phi_i = (L_i, O_i)$ whenever

$$(16) \quad L_i(A, B) = \text{sgn}(M_i \circ v_i(A) - \gamma_i \theta_{B,i}).$$

We say two elements $\phi_i, \phi'_i \in \Phi_i$ are adjacent if they differ in only one value. We say ϕ_i is realizable if $\omega_i^{-1}(\phi_i)$ is nonempty. We denote the subgraph of realizable elements in Φ_i as CPG_i and call it the combinatorial factor graph.

Theorem 4.15. For either $\text{PG} = \text{GPG}$ or $\text{PG} = \text{CPG}$, we have the following factor decomposition:

$$\text{PG} = \prod_{i=1}^N \text{PG}_i.$$

Proof. We show it first for the GPG. First we demonstrate a one-to-one correspondence between the vertices of $\prod \text{GPG}_i$ and GPG. Observe that $Z = \prod Z_i$. By straightforward topological arguments a connected component in Z is a product of connected components of Z_i and vice versa. This establishes a one-to-one correspondence of vertices between $\prod \text{GPG}_i$ and GPG. Now we show that vertices ζ, ζ' are adjacent in GPG if and only if they are adjacent in $\prod \text{GPG}_i$. Assume first that ζ and ζ' are adjacent in GPG. Then there exists a 1-deficient point in the intersection of their closure. At all such 1-deficient points there is a single equality of (8) which is satisfied; it involves parameters corresponding to a definite factor GPG_i for some

i . For all but a single exceptional $i = i^*$ we have $\pi_i(\zeta) = \pi_i(\zeta')$ and in the exceptional case $\pi_{i^*}(\zeta)$ and $\pi_{i^*}(\zeta')$ are adjacent in GPG_{i^*} . By Definition 4.12 this means ζ and ζ' are adjacent in $\prod \text{GPG}_i$. The converse more or less runs this argument in reverse. Hence $\text{GPG} = \prod \text{GPG}_i$.

Next we show $\text{CPG} = \prod \text{CPG}_i$. In this case it follows very immediately from the definitions; $\Phi = \prod \Phi_i$ and the definition of adjacency in Definition 4.9 is compatible with Definition 4.12. What remains is to see that a combinatorial parameter ϕ is realizable only if ϕ_i is realizable for all $i \in V$. To this end we leave it to the reader to verify from the definitions that $\omega^{-1}(\phi) = \prod \omega_i^{-1}(\phi_i)$, from which the result follows. ■

The utility of this decomposition theorem is that it allows us a way of understanding parameter space piecewise; given a network we can consider a single node i that has $n = \#\mathbf{S}(i)$ inputs, $m = \#\mathbf{T}(i)$ outputs, and an associated logic function M_i . Given these three things we may construct the factor graph. Thus, we can store a library of such factor graphs once and for all, and given a network we can immediately understand the combinatorial decomposition of parameter space by assembling this product structure.

4.4. Computation of parameter graph. Theorem 4.15 allows us to construct parameter graphs as products of factor graphs, so the problem of computing a parameter graph reduces to the problem of constructing the factor graphs. In light of Proposition 4.4 the factor graph is connected so we may explore it, and hence construct it, via any graph traversal technique, e.g., breadth-first-search, depth-first-search. To use this approach we require two ingredients: we must have a starting point, and we must be able construct the list of adjacent parameters.

For a starting point we may choose the combinatorial parameter $\phi = (L, O)$, where $L \equiv -1$ and O may be chosen freely. This is guaranteed to be realizable: we obtain a realization if we choose u, l , and γ values freely and then choose θ values in the desired ordering and sufficiently large so that $\Lambda_i < \gamma_i \theta_{j,i}$ for all $(i, j) \in E$.

To compute adjacency lists we first obtain from Definition 4.9 a list of candidate adjacent combinatorial parameters in Φ . Not every candidate adjacency is in CPG since not every combinatorial parameter is realizable. To obtain the adjacencies we filter the list of candidates in search of the realizable combinatorial parameters. In particular, we employ the computational algebra technique CAD [15], which provides a finite, recursive description of the geometric region ζ' associated with each candidate adjacent combinatorial parameter ϕ' (i.e., $\zeta' = \omega^{-1}(\phi')$). Since ζ' is given in terms of strict inequalities, we use the algorithm of [42] as implemented in Mathematica.

These algorithms can be quite expensive (worst-case bounds are doubly exponential in the number of variables) but are tractable for networks built out of components given in Table 1. From a CAD description of ζ' we can determine if it is empty and hence if ϕ' is realizable. If it is, then ζ' is added into the list of adjacent vertices of CPG .

In this manner we can traverse the graph and construct the factor graphs and hence the graph. Note that given a network component there is an $\#\mathbf{T}(i)!$ -fold symmetry due to the number of possible rearrangements of the θ variable orderings. This can be used to reduce the number of CAD calculations; in particular we can compute the parameter graph only for a single ordering of thresholds and from this we can extrapolate the rest of the factor graph via symmetry in a straightforward manner. In particular, we can make a disjoint union of $\#\mathbf{T}(i)!$ independent copies of the graph and then connect them together via adjacencies

corresponding to swapping threshold orderings whenever none of the Λ values are between them.

Remark 4.16. As noted above CAD computations can be quite expensive. Currently we know of no general way of constructing a CPG graph without using CAD unless we are content with performing computations for combinatorial parameters which are not realizable, i.e., do not correspond to some actual parameter $z \in Z$. This may seem like a defect of this approach but we should emphasize that these CAD computations do not need to be repeated for each network we analyze. Rather, we construct a library of CAD results corresponding to each single node in a network with a given number of inputs, number of outputs, and logic. From this library of CAD computations we may analyze any network which may be built up from these components without doing any further computational algebraic geometry. Because this library has been constructed for all nodes of the form in Table 1, we may analyze any network built out of components with these node types without performing any additional CAD computations. The Mathematica scripts we executed to produce our library of CAD results may be found in the DSGRN software package [32].

4.5. Relationship between geometric and combinatorial parameter graphs. For all $z, z' \in \zeta \in \text{GPG}$, $\omega(z) = \omega(z')$, there is a well-defined map

$$h : \text{GPG} \rightarrow \text{CPG}$$

such that $\zeta \mapsto \omega(\zeta)$. Moreover, the existence of a 1-deficient point in the intersection of the closures of the connected components $\zeta, \zeta' \in \mathcal{Z}$ immediately implies $h(\zeta)$ and $h(\zeta')$ are described by sets of inequalities which differ by only a single inequality reversal and hence are adjacent in CPG. This renders h into a *graph homomorphism* as it maps vertices to vertices and edges to edges. Note that h behaves nicely with the product structure indicated in the previous section.

Open Question 4.17. *Is h in general an isomorphism?*

An answer to the question ultimately comes down to two issues: (1) Are the geometric regions associated with the realizable combinatorial parameters connected? (2) Do the geometric regions associated with adjacent vertices of CPG always admit a 1-deficient point in the intersection of their closures? While we cannot at this time answer these questions in complete generality, we do have the following result and a method for checking specific cases.

Theorem 4.18. *The homomorphism h is an isomorphism for regulatory networks for which each node in the regulatory network is described by an entry of Table 1.*

Proof. By Theorem 4.15 it suffices to consider only the factor graphs; all that we must show is that

$$\text{GPG}_i \simeq \text{CPG}_i.$$

This in turn becomes a finite computation for each entry of Table 1. In particular we use CAD to show the following:

1. For each $\phi_i \in \text{CPG}_i$, the associated geometric region is connected.
2. For each pair of adjacent $\phi_i, \phi'_i \in \text{CPG}_i$, the associated geometric regions admit a 1-deficient point in the intersection of their closures.

For (1), observe that each node ϕ_i in CPG_i is associated with a parameter region $|\phi_i|$ in Z defined by a collection of polynomial inequalities. These inequalities are obtained by writing threshold inequalities $\theta_{j_1,i} < \theta_{j_2,i}$ according to the threshold ordering O_i and by writing either $M_i \circ v_i(A) < \gamma_i \theta_{B,i}$ or $M_i \circ v_i(A) > \gamma_i \theta_{B,i}$ for each $(A, B) \in \text{In}_i \times \text{Out}_i$ according to the sign of $L_i(A, B)$.

For (2), we consider the inequalities associated with adjacent parameters ϕ_i and with ϕ'_i in CPG_i . These two sets of inequalities are identical apart from a single inequality with its sign flipped. Dropping this inconsistent inequality yields a set of inequalities \mathcal{I} describing a region in \bar{Z}_i which contains the union of $|\phi_i|$ and $|\phi'_i|$. Since all the inequalities are strict, the solutions of \mathcal{I} make up an open set. If this open set is connected, then as an open subset of Euclidean space it is also path-connected; in particular take any path from a point in $|\phi_i|$ to a point in $|\phi'_i|$ in $|\mathcal{I}|$. By the intermediate value theorem it must pass through a point where equality is obtained on the inequality ϕ_i and ϕ'_i disagree on (but on none of the other inequalities). Thus we have identified a 1-deficient point on the intersection of the closures of $|\phi_i|$ and $|\phi'_i|$.

Using Mathematica, we have performed these connectedness computations and verified (1) for each vertex and (2) for each adjacency of the factor graphs corresponding to network components indicated in the rows of Table 1. The scripts which accomplish these computations may be found in the DSGRN software package [32]. ■

5. Applications. We demonstrate the use of the DSGRN database with two elementary three-node networks, the repressilator and bistable repressilator models, and an example associated with the p53 network. These databases, along with a growing number of other examples, can be found at the DSGRN database [32].

5.1. The repressilator. We begin with the repressilator as shown in Figure 4(a) for two reasons: first, it is of biological interest in that it has been constructed in *E. coli* by [22], and second, it is extremely simple as it consists of three genes that repress each other in a cycle and thus we can draw the entire parameter graph (see Figure 5).

Applying Definition 2.4 the switching equations for the repressilator take the form

$$(17) \quad \begin{aligned} \dot{x}_1 &= -\gamma_1 x_1 + \sigma_{1,3}^-(x_3), \\ \dot{x}_2 &= -\gamma_2 x_2 + \sigma_{2,1}^-(x_1), \\ \dot{x}_3 &= -\gamma_3 x_3 + \sigma_{3,2}^-(x_2), \end{aligned}$$

where the notation for the nonlinearities σ follows the notation of (5). We write σ^- as a reminder that the interaction represents repression, the second choice in (5).



Figure 4. Left: Repressilator. Right: Bistable repressilator.

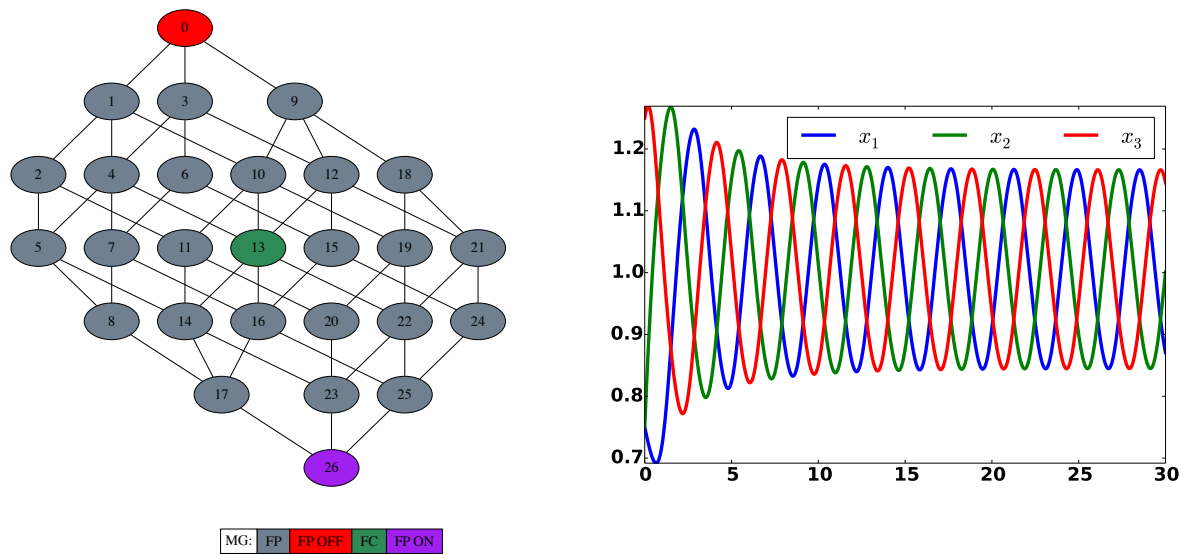


Figure 5. Left: Repressilator parameter graph with same-colored parameter nodes corresponding to the same Morse graph. Right: Hill function simulation for the repressilator satisfying the inequalities of parameter node 13 with $l_{i,j} = 0.5$, $\theta_{i,j} = 1.0$, $u_{i,j} = 1.5$ (see (21)). The Hill exponent is $n = 9$.

The repressilator model has a single regulatory threshold for each variable, dividing the phase space (which is the positive orthant of \mathbb{R}^3) into 8 cells and 12 walls. The domain and wall graphs from section 3.2 vary with parameter choice and therefore we start with a discussion of the space of parameters and the parameter graph.

For each i the function $\sigma_{i+1,i}^-$ representing the edge $i \rightarrow i + 1$ is parameterized by three parameters $u_{i+1,i}$, $l_{i+1,i}$ and $\theta_{i+1,i}$ (i is taken mod 3). In addition there are three decay constants $\gamma_1, \gamma_2, \gamma_3$. Therefore the parameter space $Z \subset \bar{Z} \subset \mathbb{R}^{12}$. To determine the parameter graph we first construct the combinatorial parameter graph CPG using the factor decomposition theorem, Theorem 4.15. For each factor of the CPG we use Table 1. Since the repressilator model has a single threshold for each variable, the threshold orders O_i as in Definition 4.14 are trivial. The choice of logic M_i (see (4)) is also trivial. Therefore for each $i = 1, 2, 3$ we consider the network node component in the first row in Table 1. The last column of the table shows that $\#PG_i = 3$ for each variable x_i . To make this explicit, for variable x_2 these choices correspond to

$$l_{2,1} < u_{2,1} < \gamma_1 \theta_{2,1}, \quad l_{2,1} < \gamma_1 \theta_{2,1} < u_{2,1}, \quad \text{and} \quad \gamma_1 \theta_{2,1} < l_{2,1} < u_{2,1}.$$

By the factor decomposition theorem, Theorem 4.15, the combinatorial parameter graph CPG has $3^3 = 27$ nodes and edges as shown in Figure 5 (left). Finally, by Theorem 4.18 the combinatorial and geometrical parameter graphs are the same and therefore Figure 5 (left) also indicates GPG.

As indicated in section 3.2 the domain and wall graphs and hence the Morse graph (Definition 3.14) are potentially different at each of the 27 parameter nodes. However, for the

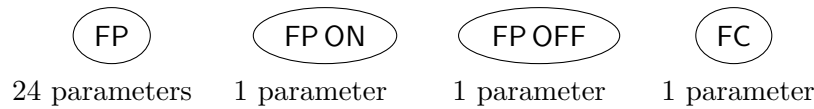


Figure 6. DSGRN Morse graphs for the repressilator and the number of parameter regions at which each Morse graph is realized.

repressilator there is a single Morse graph each consisting of a single node. This emphasizes the value of annotating the nodes of the Morse graph. As described in section 3.4 this annotation indicates whether the variables transition in the associated Morse set and, if not, whether their values are high or low. As shown in Figure 6 in the repressilator the Morse graph can assume four different annotations: a fixed point, FP; a fixed point where all variables are above the threshold, FP ON; a fixed point where all variables are below the threshold, FP OFF; and a full cycle FC where all variables pass through their threshold. In Figure 6, the number of parameters associated to each Morse graph is listed, with 24 of the 27 exhibiting the Morse graph FP.

Returning to Figure 5 (left), which shows the entire parameter graph for the repressilator, each node is color-coded according to the associated Morse graph. We note that a single parameter node gives rise to this Morse graph FC indicating a periodic orbit. To further investigate the parameter set $R_{FC} \subset Z$ which is represented by this parameter node, the DSGRN database provides us with the set of inequalities that define R_{FC} :

$$(18) \quad \begin{aligned} l_{1,3} &< \gamma_1 \theta_{2,1} < u_{1,3}, \\ l_{2,1} &< \gamma_2 \theta_{3,2} < u_{2,1}, \\ l_{3,2} &< \gamma_3 \theta_{1,3} < u_{3,2}. \end{aligned}$$

While this unique combination of parameters is represented by a single parameter graph node, R_{FC} is clearly a substantial and unbounded component in the parameter space $Z \subset \mathbb{R}^{12}$. To see if R_{FC} predicts well oscillations for a smooth repressilator model, we replace the switching model by a model that uses Hill function nonlinearities, which are closely related to the switching nonlinearities. Observe that for each $z \in |\zeta|$, where ζ is a node in the parameter graph, there is a natural one-parameter family of Hill functions. For an activating step function $\sigma^+(x)$ this takes the form

$$(19) \quad h_n^+(x) = l + (u - l) \frac{x^n}{\theta^n + x^n},$$

and for a repressing step function $\sigma^-(x)$ it takes the form

$$(20) \quad h_n^-(x) = l + (u - l) \frac{\theta^n}{\theta^n + x^n},$$

where the undetermined parameter is the Hill exponent n . Note that

$$\lim_{n \rightarrow \infty} h_n^\pm(x) = \sigma^\pm(x)$$

pointwise for all $x \neq \theta$.

We sample a point from R_{FC} that satisfies the inequalities (18), select a Hill exponent, and simulate a Hill function model with these parameters.

$$\begin{aligned}
 \dot{x}_1 &= -x_1 + 0.5 + \frac{1}{1 + x_3^n} = -x_1 + g_3(x_3), \\
 \dot{x}_2 &= -x_2 + 0.5 + \frac{1}{1 + x_1^n} = -x_2 + g_1(x_1), \\
 \dot{x}_3 &= -x_3 + 0.5 + \frac{1}{1 + x_2^n} = -x_3 + g_2(x_2).
 \end{aligned}
 \tag{21}$$

The choice of n will affect the dynamics, and so it is important to choose an n large enough so that the Hill function model (21) is reasonably representative of the switching model (3). For the repressilator, there is analysis available that allows us to suggest the minimum allowable n , which we calculate below. Tyson and Othmer [46] proved a necessary secant condition for stability of a global fixed point E in an I -dimensional cyclic feedback system (see also Thron [44]), given by

$$\frac{|g'_1(E) \cdots g'_I(E)|}{\gamma_1 \cdots \gamma_I} < \sec\left(\frac{\pi}{I}\right)^I,$$

where the Jacobian of the system is

$$\begin{pmatrix}
 -\gamma_1 & 0 & 0 & \cdots & 0 & g'_I(E) \\
 g'_1(E) & -\gamma_2 & 0 & \cdots & 0 & 0 \\
 0 & g'_2(E) & -\gamma_3 & \cdots & 0 & 0 \\
 \cdots & & & & & \\
 0 & 0 & 0 & \cdots & g'_{I-1}(E) & -\gamma_I
 \end{pmatrix}.$$

This condition is sharp when all of the decay rates are equal, which is the case here.

Applying this condition to (21), we see that for the equilibrium is $E = (1, 1, 1)$ the secant formula becomes

$$|g'_1(1)g'_2(1)g'_3(1)| < \sec\left(\frac{\pi}{3}\right)^3.
 \tag{22}$$

It is readily verified that $g'_i(1) = -n/4$ so that (22) takes the form

$$n^3 < 4^3 2^3$$

and therefore $E = (1, 1, 1)$ is stable when $n < 8$. Since the condition is sharp, it is easy to show that at $n = 8$ there is a Hopf bifurcation at which the equilibrium destabilizes and a stable periodic orbit is born. So for $n > 8$ in (21), there is a stable periodic orbit. In Figure 5 (right) we show a periodic orbit for $n = 9$.

5.2. The bistable repressilator. The bistable repressilator is slightly more complicated than the repressilator in that it has an additional negative feedback. Figure 4 represents these regulatory networks in graphical form. Because of the double feedback loop we expect that for appropriate parameter values this system may exhibit bistability, hence the name. The associated switching system is given by

$$\begin{aligned}
 \dot{x}_1 &= -\gamma_1 x_1 + \sigma_{1,2}^-(x_2) \sigma_{1,3}^-(x_3), \\
 \dot{x}_2 &= -\gamma_2 x_2 + \sigma_{2,1}^-(x_1), \\
 \dot{x}_3 &= -\gamma_3 x_3 + \sigma_{3,2}^-(x_2).
 \end{aligned}
 \tag{23}$$

The logic M_i for each node is trivial except for the first equation, where we have chosen AND logic, i.e., the negative influences of x_2 and x_3 are multiplicative.

In the bistable repressilator the variable x_2 represses both x_3 and x_1 , and so there are two choices for O_2 , $\theta_{3,2} < \theta_{1,2}$ or $\theta_{1,2} < \theta_{3,2}$. Because of the extra threshold in the bistable repressilator, there are 12 cells and 20 walls dividing the phase space.

The parameter space $Z \subset \mathbb{R}^{15}$ and we seek to understand the GPG that represents arrangement of components of Z . To construct the CPG we again consult Table 1. The variable x_1 has two inputs and one output and the logic is multiplication; this corresponds to row 6 in the table and hence $\#PG_1 = 6$. The variable x_2 has one input and two outputs and so it corresponds to row 2. However, since the last column in Table 1 lists $\#PG_i$ divided by all possible permutations of output variable thresholds, $\#PG_2 = 12$. Finally, the variable x_3 has one input and one output, which corresponds to the first row and thus $\#PG_3 = 3$. By the factor decomposition theorem, Theorem 4.15, the CPG has $6 * 12 * 3 = 216$ parameter nodes, and by Theorem 4.18 the CPG and GPG are the same. Since the parameter graph is sizable we show in Figure 7 only half of the parameter graph corresponding to one of the two orders of the thresholds in O_2 ($\theta_{3,2} < \theta_{1,2}$).

Using very similar annotation to the repressilator, there are seven distinct classes indicated in Figure 8. For simplicity in Figure 7 we group the annotations FP, FP ON, and FP OFF together and call this group FP. We further group the parameter nodes based on the following four annotations: type A nodes have a Morse graph with a single fixed point FP; nodes in class B have a Morse graph with two fixed points FP and hence signal the presence of bistability; nodes of type C have Morse graph FC; and nodes of type D have the Morse graph with the lower Morse set FP and the upper Morse set FC.

The collection of Morse graphs immediately signals the presence of richer dynamics across parameter space than for the repressilator system. In addition to the four Morse graphs seen in the repressilator example, there are two new dynamical signatures present: bistability (the existence of two stable fixed points, type B) and an unstable full cycle with an attracting fixed point (type D). The comparison of Figures 6 and 8 shows that the addition of a single edge to a regulatory network **RN** can radically change the dynamical signature of **RN** across parameter space.

From Figure 8, we see that there are six parameter vertices with the Morse graph FC, which suggest a presence of a stable periodic oscillation. The DSGRN database provides us with the inequalities that define these regions in the parameter Z . For the purpose of illustration, we select one of them (parameter 151), which represents a region in $Z \subset \mathbb{R}^{15}$ given by

$$\begin{aligned}
 l_{1,2} l_{1,3} &< \left\{ \begin{array}{l} u_{1,2} l_{1,3} \\ l_{1,2} u_{1,3} \end{array} \right\} < \gamma_1 \theta_{2,1} < u_{1,2} u_{1,3}, \\
 l_{2,1} &< \gamma_2 \theta_{3,2} < u_{2,1} < \gamma_2 \theta_{1,2}, \\
 l_{3,2} &< \gamma_3 \theta_{1,3} < u_{3,2}.
 \end{aligned}
 \tag{24}$$

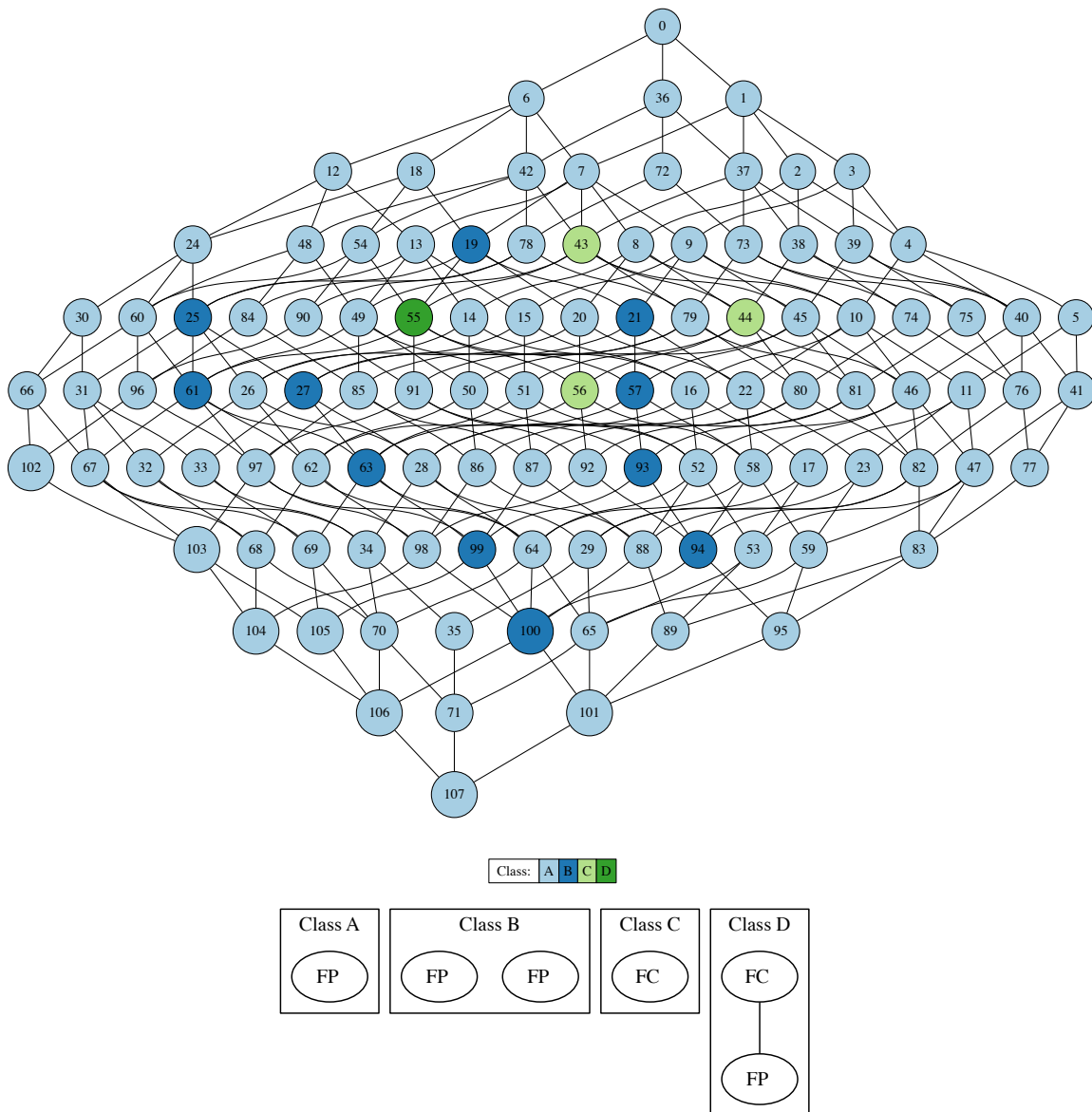


Figure 7. Bistable repressilator parameter graph with colors corresponding to partitioned Morse graph continuation classes. Class A: single stable fixed points; Class B: bistability; Class C: stable cycle; Class D: unstable cycle with a stable fixed point.

The curly braces denote an undetermined (arbitrary) order: the relative order of $u_{1,2}l_{1,3}$ and $l_{1,2}u_{1,3}$ does not change the wall graph and hence the Morse graph as long as both remain below $\gamma_1\theta_{2,1}$.

We sample this parameter region at the values $\gamma_1 = \gamma_2 = \gamma_3 = 1$, $l_{i,j} = 1$, $\theta_{1,3} = 2$, $\theta_{3,2} = 3$, $\theta_{2,1} = 4$, $\theta_{1,2} = 6$, $u_{1,2} = 2$, $u_{1,3} = 3$, $u_{3,2} = 4$, and $u_{2,1} = 5$. Substituting these

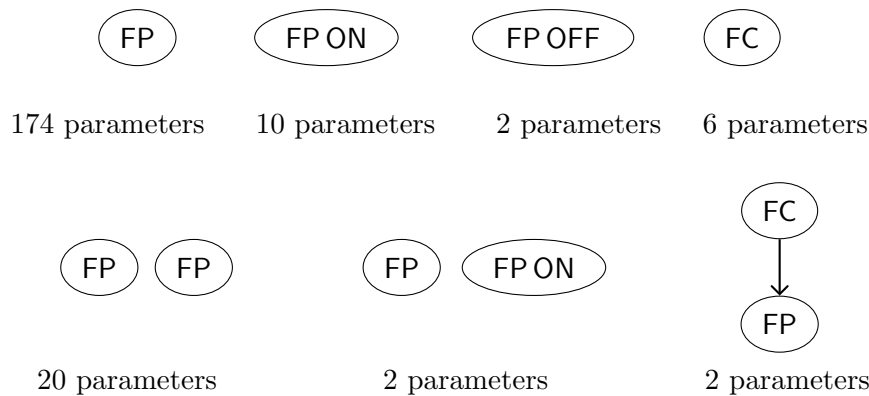


Figure 8. DSGRN Morse graphs for the bistable repressilator.

values into (20), we have the following system of smooth equations:

$$\begin{aligned}
 \dot{x}_1 &= -x_1 + \left(1 + \frac{6^n}{6^n + x_2^n}\right) \left(1 + \frac{2^{n+1}}{2^n + x_3^n}\right), \\
 \dot{x}_2 &= -x_2 + 1 + \frac{4^{n+1}}{4^n + x_1^n}, \\
 \dot{x}_3 &= -x_3 + 1 + \frac{3^{n+1}}{3^n + x_2^n}.
 \end{aligned}
 \tag{25}$$

Results of the simulations are shown in Figure 9 for $n = 10$. There is no condition analogous to the secant condition that would provide an estimate for n that would produce a stable periodic orbit. Numerically, we found that at these parameter values $n = 7$ is sufficient for periodicity and $n = 6$ is not.

We want to finish this section with an important observation about a relationship between the parameter graph for the repressilator and the parameter graph for the bistable repressilator. Since the repressilator network is a subnetwork of the bistable repressilator, a natural question is whether there is a similar correspondence between their parameter graphs. To begin to answer this question we investigate the parameter node given by (24) in the bistable repressilator. We first note that because $l_{2,1} < u_{2,1} < \gamma_2 \theta_{1,2}$ the second component of any target point in the system $\frac{\sigma_{2,1}^-(\cdot)}{\gamma_2} < \theta_{1,2}$. Therefore, perhaps after a transient, $x_2(t) < \theta_{1,2}$ for all $t \geq T$ and some $T > 0$. Consequently, the value of the function $\sigma_{1,2}^-(x_2)$ will be $u_{1,2}$ for all $t \geq T$ and x_2 will effectively cease regulation of x_1 . Therefore the network that is *effectively* represented by this parameter node is not a bistable repressilator but a repressilator where the edge from x_2 to x_1 is erased. This brings up a set of interesting questions about how to recognize subnetworks that are effectively represented by each node in the parameter graph and whether it is possible to build parameter graphs of larger networks from parameter graphs of their subnetworks. The answers to these questions are beyond the scope of this paper but will be addressed in the near future.

5.3. p53 network. While the first two examples of this section were aimed at illustrating the main concepts of the paper on small networks, DSGRN is being used on larger and more

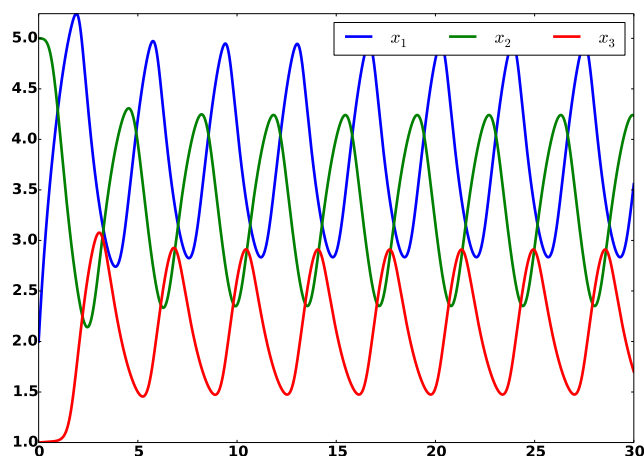


Figure 9. Hill function simulation for the bistable repressilator at parameter node 151. See the text for parameter choices. The Hill exponent is $n = 10$.

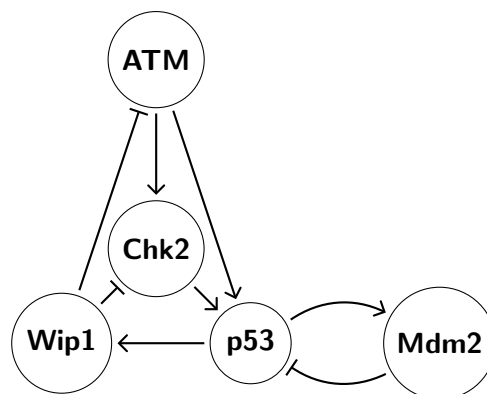


Figure 10. Subnetwork of key species of the p53 signaling network.

complicated networks with greater biological urgency. We briefly comment on simulations of a subnetwork of the p53 signaling network from [37]. The point of including this model in this paper is to indicate the ease with which DSGRN handles a network of this size and then allows the dynamics to be interrogated.

As indicated in Figure 10 this network has 5 nodes and 8 edges and thus the parameter space $Z \subset \bar{Z} \subset \mathbb{R}^{29}$. As in the previous cases we construct the geometric parameter graph GPG by first computing the CPG using the factors listed in Table 1. This results in a GPG with 803,520 nodes. Next we construct a DSGRN database over this parameter graph. This problem is quite tractable and the database construction took only 37 seconds on a mid-2014 Macbook Pro laptop (Intel Core i7-4870HQ CPU, 2.50 GHz). We note that for larger parameter graphs, our software can scale to HPC cluster environments, but in this case this is clearly not necessary.

We remark that there is interest in the question of stable oscillations in this system [26]. Given the size of the GPG, visualization is impractical. However, as indicated in the introduction DSGRN produces an SQL database. The query for a Morse graph with a minimal node annotated by FC identifies parameter nodes associated with stable recurrent dynamics where all the species pass thresholds. There are 6904 nodes in the GPG which satisfy this query, of which 3204 are associated with a Morse graph consisting of a single node.

Similar to the previous example we sample a parameter from the parameter region corresponding to one of these combinatorial parameters and perform a numerical simulation. But unlike the previous example, it is less obvious how to obtain such a sample. The parameter region is a semialgebraic set defined by a collection of strict inequalities which do not have an obvious solution technique. However, as discussed in section 4.4 we have computed CADs of the semialgebraic sets describing combinatorial parameters. These descriptions give a more convenient representation of the semialgebraic set which allow us to readily produce solutions. For a rigorous account of CADs we refer the reader to [14] and [34], but we give a brief explanation as follows. First, the notion of CAD requires we impose an ordering on variables. With respect to an imposed variable ordering, a *cylindrical component* is a semialgebraic set described by a sequence of inequalities providing lower and upper bounds (either strict or non-strict) for each variable in sequence according to the imposed ordering. The lower and upper bounds for each variable are algebraic functions of the preceding variables, and the crucial property is that given valid choices for the preceding variables the lower bound cannot exceed the upper bound (and hence there is always a choice for the next variable given choices of previous variables). This gives rise to a simple procedure for finding solutions points whereby we find a value for each variable in turn and substitute the variables we have chosen so far into the lower and upper bounds of the next inequality in order to choose the next variable. A CAD of an algebraic set S represents the set S as a disjoint union of cylindrical components. Applying this to the present situation, we present a CAD description of the semialgebraic set corresponding to a combinatorial parameter (which is indexed by our algorithms as node 40535) in the p53 parameter graph. This description is given in Table 2. Notice we simplified the subscript notation. Each edge in the graph $k \rightarrow j$ is now associated to a single index, i , and each edge has three numerical values associated to it: $U_i := u_{j,k}$, $L_i := l_{j,k}$, and $T_i := \gamma_j \theta_{j,k}$. From this table we see there are three cylindrical components comprising the semialgebraic set in \mathbb{R}^{29} . We remark that it is possible from CAD to decide connectedness [34] of the semialgebraic set, though we omit details on how this is done. Choosing any column, numerical values U, L, and T can be iteratively chosen. For example, in the top section, T_7 , L_1 , and L_2 are chosen to be completely independent values. Then L_3 is chosen to be less than an algebraic combination of those three values, followed by U_1 dependent on the previous four values, and so forth. An acceptable geometric parameter will be found by iteratively choosing values from any one of the three columns (or connected components). The property that these are cylindrical components means this procedure cannot fail (i.e., no matter what choices are made, one cannot reach an inequality where the lower bound exceeds the upper bound and it is impossible to continue). A solution which can be found using this technique is given in Table 3. We also remark that the factorization of the GPG discussed in section 4.3 is visible in the CAD description of Table 2. We have indicated this by partitioning the inequalities into sections with horizontal lines. For example, the top section describes the parameter for the factor graph of node p53, the second for node Chk2, etc.

Table 2

CAD description of a parameter node for the p53 network. We use the following numeric scheme to identify the edges: 1 = (ATM → p53), 2 = (Chk2 → p53), 3 = (Mdm2 → p53), 4 = (Wip1 → ATM), 5 = (ATM → Chk2), 6 = (Wip1 → Chk2), 7 = (p53 → Wip1), 8 = (p53 → Mdm2). The upper values, lower values, and thresholds (or product of threshold and decay rate) for each edge correspond to U_i , L_i , and T_i respectively. Example: The upper value associated with the edge (ATM → Chk2) corresponds to U_5 .

| Component 1 | Component 2 | Component 3 |
|--|---|--|
| $0 < T_7$ $0 < L_1$ $0 < L_2$ $0 < L_3 < \frac{T_7}{L_1+L_2}$ $L_1 < U_1 \leq \frac{L_3(L_1-L_2)+T_7}{2L_3}$ $\frac{T_7-L_3U_1}{L_3} < U_2 < \frac{T_7-L_1L_3}{L_3}$ $L_3 < U_3 < \frac{T_7}{L_1+U_2}$ $T_7 < T_8 < L_3(U_1 + U_2)$ | $0 < T_7$ $0 < L_1$ $0 < L_2$ $0 < L_3 < \frac{T_7}{L_1+L_2}$ $\frac{L_3(L_1-L_2)+T_7}{2L_3} < U_1 < \frac{T_7-L_2L_3}{L_3}$ $\frac{T_7-L_3U_1}{L_3} < U_2 < L_2 - L_1 + U_1$ $L_3 < U_3 < \frac{T_7}{L_2+U_1}$ $T_7 < T_8 < L_3(U_1 + U_2)$ | $0 < T_7$ $0 < L_1$ $0 < L_2$ $0 < L_3 < \frac{T_7}{L_1+L_2}$ $\frac{L_3(L_1-L_2)+T_7}{2L_3} < U_1 < \frac{T_7-L_2L_3}{L_3}$ $L_2 + U_1 - L_1 \leq U_2 < \frac{T_7-L_1L_3}{L_3}$ $L_3 < U_3 < \frac{T_7}{L_1+U_2}$ $T_7 < T_8 < L_3(U_1 + U_2)$ |
| $0 < T_2$ $0 < L_5$ $0 < L_6 < \frac{T_2}{L_5}$ $L_5 < U_5 < \frac{T_2}{L_6}$ $\frac{T_2}{U_5} < U_6 < \frac{T_2}{L_5}$ | $0 < T_2$ $0 < L_5$ $0 < L_6 < \frac{T_2}{L_5}$ $L_5 < U_5 < \frac{T_2}{L_6}$ $\frac{T_2}{U_5} < U_6 < \frac{T_2}{L_5}$ | $0 < T_2$ $0 < L_5$ $0 < L_6 < \frac{T_2}{L_5}$ $L_5 < U_5 < \frac{T_2}{L_6}$ $\frac{T_2}{U_5} < U_6 < \frac{T_2}{L_5}$ |
| $0 < T_6$ $0 < U_7 < T_6$ $0 < L_7 < U_7$ $L_7 < T_4 < U_7$ | $0 < T_6$ $0 < U_7 < T_6$ $0 < L_7 < U_7$ $L_7 < T_4 < U_7$ | $0 < T_6$ $0 < U_7 < T_6$ $0 < L_7 < U_7$ $L_7 < T_4 < U_7$ |
| $0 < L_4$ $L_4 < U_4$ $L_4 < T_1 < U_4$ $T_1 < T_5 < U_4$ | $0 < L_4$ $L_4 < U_4$ $L_4 < T_1 < U_4$ $T_1 < T_5 < U_4$ | $0 < L_4$ $L_4 < U_4$ $L_4 < T_1 < U_4$ $T_1 < T_5 < U_4$ |
| $0 < T_3$ $0 < L_8 < T_3$ $T_3 < U_8$ | $0 < T_3$ $0 < L_8 < T_3$ $T_3 < U_8$ | $0 < T_3$ $0 < L_8 < T_3$ $T_3 < U_8$ |

Table 3

p53 network parameters.

| Edge | u-value | l-value | θ-value |
|-------------|---------|---------|-----------|
| ATM → Chk2 | 1 | 1/2 | 1/2 |
| ATM → p53 | 7/8 | 7/32 | 1/4 |
| Chk2 → p53 | 7/8 | 7/32 | 3/4 |
| Mdm2 → p53 | 7/8 | 21/32 | 1 |
| Wip1 → ATM | 1 | 1/2 | 1/2 |
| Wip1 → Chk2 | 2 | 1/2 | 2 |
| p53 → Mdm2 | 2 | 1/2 | 1127/1024 |
| p53 → Wip1 | 1 | 1/4 | 539/512 |

As in the previous examples we perform a numerical simulation of this system using Hill functions. Using the CAD description of Table 2 we choose decay rates $\gamma_i = 1$ and the remaining parameters of the switching system as presented in Table 3. It remains to choose the Hill exponent n for each nonlinearity, i.e., one exponent for each edge in the network.

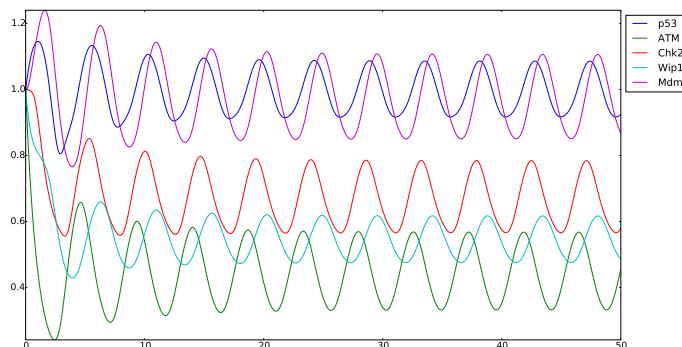


Figure 11. Hill function simulation for the p53 model at parameter node 40535. See the text for parameter choices. The Hill exponent for every nonlinearity is $n = 8$.

Setting all Hill exponents to be 8 the solution exhibits the oscillations depicted in Figure 11. The uniform choice of $n = 6$ does not produce oscillations, but many other choices, e.g., setting $n = 2$ for the connections $\text{Mdm2} \rightarrow \text{p53}$ and $\text{p53} \rightarrow \text{Mdm2}$ and $n = 10$ for all other nonlinearities, also produces oscillations. It is worth noting that the peaks of p53 in Figure 11 come slightly ahead of the peaks of Mdm2, which agrees with one of the key experimental observations in [37].

This last example is meant to indicate the usefulness of the CAD description of the nodes in parameter space. It provides a scheme for efficiently sampling parameter points with specific combinatorial properties. This opens the possibility for efficiently studying numerical simulations to understand the finer structure of the dynamics that satisfies the local and global properties of the annotated Morse graph.

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