An analytical approach to bistable biological circuit discrimination using real algebraic geometry

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1 Summary

Biomolecular circuits with two distinct and stable steady states have been identified as essential components in a wide range of biological networks, with a variety of mechanisms and topologies giving rise to their important bistable property. Understanding the differences between circuit implementations is an important question, particularly for the synthetic biologist faced with determining which bistable circuit design out of many is best for their specific application. In this work we explore the applicability of Sturm’s theorem—a tool from 19th-century real algebraic geometry—to comparing “functionally equivalent” bistable circuits without the need for numerical simulation. We first consider two genetic toggle variants and two different positive feedback circuits, and show how specific topological properties present in each type of circuit can serve to increase the size of the regions of parameter space in which they function as switches. We then demonstrate that a single competitive monomeric activator added to a purely-monomeric (and otherwise monostable) mutual repressor circuit is sufficient for bistability. Finally, we compare our approach with the Routh-Hurwitz method and derive consistent, yet more powerful, parametric conditions. The predictive power and ease of use of Sturm’s theorem demonstrated in this work suggests that algebraic geometric techniques may be underutilized in biomolecular circuit analysis.

2 Key words

biological circuit; bistability; algebraic geometry; Sturm’s theorem; synthetic biology

3 Introduction

The field of synthetic biology has rapidly matured to the point where it is now possible to produce complex synthetic networks with prescribed functions and level of performance [1]. As in other fields of engineering, advances have been enabled by the use of small interchangeable modules that are “functionally equivalent” from an input-output perspective [2]. Bistable circuits—which play a role in essential biological processes including cell fate specification [3], cell cycle progression [4], and apoptosis [5]—make up a particularly large and diverse functionally equivalent set [6]. Effectively characterizing and comparing these biocircuits is crucial for determining which design is in some sense optimal for a particular context.
Ordinary differential equation (ODE) models can be powerful tools for identifying and contrasting biocircuits’ “dynamic phenotypes” (see, e.g., [7]). Numerical simulations are often used in the analysis of these models; however, analytical criteria that focus on topology can provide a more exact assessment of a circuit’s properties [8, 9]. (We use topology to mean the particular set of interactions between regulatory parts.) A novel analytical tool that can provide topology-based insights can be found in Sturm’s theorem [10], developed in 1835 as a solution to the problem of finding the number of real roots of an algebraic equation with real coefficients over a given interval. Despite its predictive power, this “gem of 19th century algebra and one of the greatest discoveries in the theory of polynomials” [11] remains an unexploited tool for analysis of biological circuit models.

In this work we demonstrate an approach to bistable circuit discrimination based on Sturm’s theorem that can give the boundaries of the regions of bistability as exact analytic expressions, eliminating the need for numerical simulation. We compare the regions of bistability for two variants of the classic double-negative toggle switch as well as two positive feedback circuits, one of which is based on the bacteriophage λ promoter P<sub>RM</sub>. We then show that while a purely monomeric version of the genetic toggle cannot be bistable, a single competitive activating species added to the circuit leads to bistability in a noncontiguous region of parameter space. Lastly, we use a model of an RNA aptamer-based bistable switch to compare our Sturm’s theorem approach to another based on the control theoretic Routh-Hurwitz method. Overall our results highlight a new use for Sturm’s theorem for identifying potential differences between functionally equivalent bistable biocircuits, and serve as a (re-)introduction to the method as a general tool for studying the kinds of polynomial expressions that often arise when modeling biological systems.

4 Mathematical preliminaries

4.1 Sturm’s theorem

Sturm’s theorem gives the number of distinct real roots of a univariate polynomial \( f(x) \) in a particular interval. To apply the theorem, we must first construct the Sturm sequence, a set of polynomials \( \mathcal{F} = \{ f_0, f_1, \ldots, f_m \} \) defined as:

\[
\begin{align*}
    f_0 &= f(x), \\
    f_1 &= f'(x), \\
    f_2 &= -\text{rem}(f_0, f_1), \\
    f_3 &= -\text{rem}(f_1, f_2), \\
    &\vdots \\
    f_m &= -\text{rem}(f_{m-2}, f_{m-1}), \\
    0 &= -\text{rem}(f_{m-1}, f_m),
\end{align*}
\]

where \( \text{rem}(f_i, f_{i+1}) \) is the remainder of the polynomial long division of \( f_i \) by \( f_{i+1} \). The sequence ends at \( f_m \) when \( f_{m-1} \) divided by \( f_m \) gives a remainder of zero. For a polynomial of degree \( n \), there are \( m \leq n + 1 \) Sturm polynomials in the sequence.

**Theorem 1 (Sturm’s theorem)** Let \( f(x) \) be a real-valued univariate polynomial and \( a, b \in \mathbb{R} \cup \{-\infty, +\infty\} \), with \( a < b \) and \( f(a), f(b) \neq 0 \). Then the number of zeroes of \( f(x) \) in the interval \( (a, b) \) is the difference

\[
\text{var}(\mathcal{F}, a) - \text{var}(\mathcal{F}, b),
\]

**Theorem 1**
where \( F \) is the Sturm sequence of \( f(x) \), and the variations \( \text{var}(F, a) \) and \( \text{var}(F, b) \) are the number of times that consecutive nonzero elements of the Sturm sequence—evaluated at \( a \) and \( b \), respectively—have opposite signs. (Adapted from [12].)

### 4.2 Number of steady states and bistability

Our approach involves identifying regions of bistability by finding conditions that lead to three steady states, without requiring numerical determination of the exact values or stability of the equilibrium points. While it is in general not possible to draw conclusions on the stability properties of equilibria by simply counting their number, the circuits under consideration enjoy important properties—namely, they are dissipative, their linearizations are positive, and all terms of the characteristic polynomials other than the constant term are positive—that allow us to relate their degree and number of equilibria to the stability properties of each equilibrium (see Supporting Information). For such circuits, when three equilibria are present, two of them must be stable and one must be unstable.

### 5 Overview of the method

To determine the region in parameter space in which a particular circuit exhibits bistability, we first find the polynomial \( f(x) \) that describes the equilibrium state of the system and construct its Sturm sequence \( F_x \). (Here, \( x \) represents the concentration of one of the circuit species.) We then evaluate \( F_x \) at \( x \to 0 \) and \( x \to +\infty \) (since we are only interested in positive concentrations) and find the conditions leading to three steady states by first enumerating all the possible combinations of inequalities that could yield a variation difference \( \text{var}(F_x, 0) - \text{var}(F_x, +\infty) = 3 \) and then testing them for logical consistency (i.e., whether all inequalities can be simultaneously satisfied). It is important to note that, since Sturm’s theorem is only concerned with consecutive nonzero elements and zeroes are ignored, a particular Sturm polynomial may be equal to zero without affecting the total number of sign changes. In cases where zeroes are valid options in the sequence, strict inequalities are made nonstrict (e.g., ‘\( > \)’ \( \rightarrow \) ‘\( \geq \)’).

Symbolic manipulation is done using Mathematica; a sample Mathematica notebook in which the method is applied is included as supplementary material.

### 6 Results

#### 6.1 Genetic toggle circuits

A recent study identified a set of eleven minimal bistable networks (MBNs), simple two-gene circuits with the capacity for bistability that do not also contain a smaller bistable subcircuit [13]. One of these MBNs, a double-negative toggle switch consisting of two dimeric repressors (Fig. 1A, top), was among the very first synthetic biocircuits built and modeled [14]. This dimer-dimer (DD) toggle design has since gone on to be used in a wide range of synthetic biological applications, including the manipulation of fluxes of the \( E. coli \) metabolic network [15] and as part of a circuit involved in programmed autonomous cellular diversification [16]. A second MBN of particular interest, herein referred to as the monomer-dimer (MD) toggle, is a double-negative switch variant in which one of the repressors functions as a monomer (Fig. 1A, bottom). While to our knowledge no MD toggle circuit has been constructed, the components necessary for its implementation exist in monomeric transcription-activator-like effectors (TALEs) and CRISPR/Cas nucleases that have been engineered as transcriptional repressors [17, 18]. Other exotic toggle-like circuit
topologies have also been proposed and/or built (see, e.g., [13, 17, 19–21]). An understanding of the differences in how these various toggles perform can be beneficial, in particular for those circuits that have yet to be built and for which a priori knowledge of their behavior could aid in their development.

![Dimer-dimer toggle](image1)

![Monomer-dimer toggle](image2)

Figure 1: (A) Dimer-dimer (top) and monomer-dimer (bottom) toggle switches. (B) Bistable regions for the monomer-dimer and dimer-dimer toggles. The dimer-dimer toggle exhibits bistability over a much larger range of $\hat{X}_{1\text{tot}}$ and $\hat{X}_{2\text{tot}}$.

As a first demonstration of our approach, we apply Sturm’s theorem to the DD and MD toggle circuits. Beginning with a chemical reaction network formulation and assuming mass-action kinetics we derive ODE models of the two toggles (Eqs. (S1) and (S2)). At equilibrium the concentrations of $P_1$ and $P_2$ in the MD system are given by

$$P_{1eq} = \frac{\beta_1 X_{1\text{tot}}}{1 + (P_{2eq}/K_2)^2}, \quad P_{2eq} = \frac{\beta_2 X_{2\text{tot}}}{1 + (P_{1eq}/K_{md})^2},$$

and in the DD system,

$$P_{1eq} = \frac{\beta_1 X_{1\text{tot}}}{1 + (P_{2eq}/K_2)^2}, \quad P_{2eq} = \frac{\beta_2 X_{2\text{tot}}}{1 + (P_{1eq}/K_{dd})^2},$$

where $X_{i\text{tot}}$ is the total amount of gene $i$, $\beta_i = k_{\text{basi}}/k_{\text{degi}}$ is the ratio of basal production rate to degradation rate for protein $i$, $K_{xd} = \{K_{md}, K_{dd}\}$ is the Michaelis constant for $P_1$ (different for the MD and DD toggle cases), and $K_2$ is the Michaelis constant for $P_2$ (see Supporting Information for details). Systems (1) and (2) may be written in terms of $P_{1eq}$ alone, as:

$$\left(\frac{P_{1eq}}{K_{md}}\right)^3 - \left(\frac{\beta_1 X_{1\text{tot}}}{K_{md}} - 2 \left(\frac{P_{1eq}}{K_{md}}\right) - \left(2 \frac{\beta_1 X_{1\text{tot}}}{K_{md}} - \left(\frac{\beta_2 X_{2\text{tot}}}{K_2}\right)^2 - 1 \right) \left(\frac{P_{1eq}}{K_{md}}\right) - \frac{\beta_1 X_{1\text{tot}}}{K_{md}} = 0$$

and

$$\left(\frac{P_{1eq}}{K_{dd}}\right)^5 - \beta_1 X_{1\text{tot}} \left(\frac{P_{1eq}}{K_{dd}}\right)^4 + 2 \left(\frac{P_{1eq}}{K_{dd}}\right)^3 - 2 \frac{\beta_1 X_{1\text{tot}}}{K_{dd}} \left(\frac{P_{1eq}}{K_{dd}}\right)^2 + \left(\frac{\beta_2 X_{2\text{tot}}}{K_2}\right)^2 + 1 \left(\frac{P_{1eq}}{K_{dd}}\right) - \frac{\beta_1 X_{1\text{tot}}}{K_{dd}} = 0,$$
With the following scaling of the DNA and protein concentrations:
\[ \tilde{X}_{1\text{tot}} = \beta_1(X_{1\text{tot}}/K_{xd}) , \quad \tilde{X}_{2\text{tot}} = \beta_2(X_{2\text{tot}}/K_2) \]
and
\[ \tilde{P}_{1eq} = P_{1eq}/K_{xd} , \quad \tilde{P}_{2eq} = P_{2eq}/K_2 , \]
we may write Eqs. (3) and (4) as nondimensional polynomials in \( \tilde{P}_{1eq} \):
\[ \tilde{P}_{1eq}^3 - (\tilde{X}_{1\text{tot}} - 2)\tilde{P}_{1eq}^2 - (2\tilde{X}_{1\text{tot}} - \tilde{X}_{2\text{tot}}^2 - 1)\tilde{P}_{1eq} - \tilde{X}_{1\text{tot}} = 0 \]
and
\[ \tilde{P}_{1eq}^5 - \tilde{X}_{1\text{tot}}\tilde{P}_{1eq}^4 + 2\tilde{P}_{1eq}^3 - 2\tilde{X}_{1\text{tot}}\tilde{P}_{1eq}^2 + (\tilde{X}_{2\text{tot}}^2 + 1)\tilde{P}_{1eq} - \tilde{X}_{1\text{tot}} = 0 , \]
for the MD and DD toggles, respectively. Every positive root of these equilibrium polynomials gives a positive steady state concentration for every other circuit component as well.

We are interested in the two toggles’ regions of bistability in the plane of total (non-dimensionalized) DNA concentrations \( \tilde{X}_{1\text{tot}} \) and \( \tilde{X}_{2\text{tot}} \). We construct the Sturm sequences \( \mathcal{F}_x \) for Eqs. (7) and (8), evaluate \( \mathcal{F}_x \) at \( \tilde{P}_{1eq} \to 0 \) and \( \tilde{P}_{1eq} \to +\infty \), and find the conditions leading to a variation difference \( \text{var}(\mathcal{F}_x, 0) - \text{var}(\mathcal{F}_x, +\infty) = 3 \). For the DD toggle it is necessary to generate two different sequences from Eq. (8)—one with \( \tilde{X}_{1\text{tot}} \neq \sqrt{5} \) and another with \( \tilde{X}_{1\text{tot}} = \sqrt{5} \)—so that all Sturm polynomial denominators are nonzero and the sequence does not terminate prematurely. Sturm sequences are given in the Supporting Information.

The MD toggle Sturm sequence \( \mathcal{F}_{md} \) has a maximum possible variation of 3 and only one combination of inequalities that can give rise to bistability: when \( \text{var}(\mathcal{F}_{md}, 0) = 3 \) and \( \text{var}(\mathcal{F}_{md}, +\infty) = 0 \). In contrast, the DD toggle sequence \( \mathcal{F}_{dd} \) could in principle yield as many as five positive steady states; however, only three are admitted as there are no combinations of inequalities that have a variation difference of 5 or 4 and are logically consistent. All possible inequality sets for the DD toggle are listed (in the compact form \( \{\pm \cdots \pm\} \)) along with their allowabilities in Tables 1 and 2. ‘Allowed’ refers to the logical consistency of the inequality set.

<table>
<thead>
<tr>
<th>( \text{var}(\mathcal{F}_{dd}, 0) )</th>
<th>sign(( \mathcal{F}_{dd}, 0 ))</th>
<th>Allowed?</th>
<th>( \text{var}(\mathcal{F}_{dd}, +\infty) )</th>
<th>sign(( \mathcal{F}_{dd}, +\infty ))</th>
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</tbody>
</table>

Table 1: Sturm sequence inequality sets for the DD toggle when \( \tilde{X}_{1\text{tot}} \neq \sqrt{5} \). The signs of the first two polynomials are fixed at \( \tilde{P}_{1eq} \to 0 \) and \( \tilde{P}_{1eq} \to +\infty \). Neither \( \text{var}(\mathcal{F}_{dd}, 0) = 5 \) nor \( \text{var}(\mathcal{F}_{dd}, +\infty) = 0 \) represent logically consistent sets.

The analytic expressions for the two regions of bistability are Eq. (S24) (MD toggle) and the intersection of Eqs. (S25) and (S26) (DD toggle). We find that the DD toggle operates as a bistable switch over a substantially greater range of (non-dimensionalized) DNA concentrations than does the MD toggle (Fig. 1B), indicating that the DD topology is more functionally robust to variations in DNA concentrations and rate parameters. Furthermore, the DD switch can operate with significantly lower concentrations of DNA: >50% less \( \tilde{X}_{2\text{tot}} \) and >75% less \( \tilde{X}_{1\text{tot}} \).
We may also check the stability of the various steady states using the circuits’ Jacobian $J$ and characteristic polynomial $p_J(\lambda) = \det(\lambda I - J)$. It was recently shown that if all off-diagonal components of the Jacobian are nonnegative (i.e., it is a Metzler matrix), or if the Jacobian may be transformed to have such a form, then any equilibrium is unstable if and only if the constant term of $p_J(\lambda)$ has a sign opposite to that of all other terms in $p_J(\lambda)$ [19]. We use this condition on the constant term of $p_J(\lambda)$ to confirm that each bistable solution set contains one and only one unstable steady state.

The inequalities that satisfy the $p_J(\lambda)$ constant term condition are:

$$P_{1eq}^4 + 4P_{1eq}^3 + 2P_{1eq}(\hat{X}_{2tot}^2 + 3) + P_{1eq}(4 - 2(\hat{X}_{1tot} - 2)\hat{X}_{2tot}^2) - 2(\hat{X}_{1tot} - 1)\hat{X}_{2tot}^2 + \hat{X}_{2tot}^4 + 1 < 0$$

(9)

and

$$(P_{1eq}^4 + 2P_{1eq}^2 + \hat{X}_{2tot}^2 + 1)^2 - 4P_{1eq}(P_{1eq}^2 + 1)X_{1tot}X_{2tot}^2 < 0$$

(10)

for the MD and DD toggles, respectively. For each bistable solution set found we substituted the values $\hat{X}_{1tot}, \hat{X}_{2tot}$, and $P_{1eq}$ into Eqs. (9) and (10) and confirmed that only one of the three solutions satisfies the appropriate instability condition.

It is worth noting that the time required to test $(\hat{X}_{1tot}, \hat{X}_{2tot})$ pairs scales linearly with the number of pairs, so while testing a small number can be done relatively quickly, as the number of pairs becomes appreciable the time can be significant—up to 3 hours to test 600,000 random values of $\hat{X}_{1tot}$ and $\hat{X}_{2tot}$.

### 6.2 Bistable single-gene circuits

The single gene system consisting of bacteriophage $\lambda$ repressor and its promoter $P_{RM}$ (with its three operator sites OR1, OR2, and OR3) also exhibits bistability [22]. Using the dimensionless model given

<table>
<thead>
<tr>
<th>$\var{\mathcal{F}_{dd}}$, 0</th>
<th>sign($\mathcal{F}_{dd}$, 0)</th>
<th>Allowed?</th>
<th>$\var{\mathcal{F}_{dd}}$, $+\infty$</th>
<th>sign($\mathcal{F}_{dd}$, $+\infty$)</th>
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<td>N</td>
<td>0</td>
<td>${++ + + +}$</td>
<td>Y</td>
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Table 2: Sturm sequence inequality sets for the DD toggle when $\hat{X}_{1tot} = \sqrt{5}$. The signs of the first two and three polynomials are fixed at $\hat{P}_{1eq} \to 0$ and $\hat{P}_{1eq} \to +\infty$, respectively. The set with $\var{\mathcal{F}_{dd}}$, 0 = 4 is not logically consistent.

### 6.1.1 Computational support

Recognizing that certain mathematical tools used may be unfamiliar, some computational validation of our results may be of value. For both toggle circuits, and for each of the valid combinations of sign($\mathcal{F}$, 0) and sign($\mathcal{F}$, $+\infty$), 1000 random values of $\hat{X}_{1tot}$ and $\hat{X}_{2tot}$ were selected from inside and outside of the predicted bistable regions and plugged in to the appropriate equilibrium polynomial (Eq. (7) or (8)) which were then solved numerically. In all cases the number of equilibria found matched the number determined by Sturm’s theorem: three equilibria were found inside the bistable regions and only one equilibrium was found outside.

We may also check the stability of the various steady states using the circuits’ Jacobian $J$ and characteristic polynomial $p_J(\lambda) = \det(\lambda I - J)$. It was recently shown that if all off-diagonal components of the Jacobian are nonnegative (i.e., it is a Metzler matrix), or if the Jacobian may be transformed to have such a form, then any equilibrium is unstable if and only if the constant term of $p_J(\lambda)$ has a sign opposite to that of all other terms in $p_J(\lambda)$ [19]. We use this condition on the constant term of $p_J(\lambda)$ to confirm that each bistable solution set contains one and only one unstable steady state.

The inequalities that satisfy the $p_J(\lambda)$ constant term condition are:

$$P_{1eq}^4 + 4P_{1eq}^3 + 2P_{1eq}(\hat{X}_{2tot}^2 + 3) + P_{1eq}(4 - 2(\hat{X}_{1tot} - 2)\hat{X}_{2tot}^2) - 2(\hat{X}_{1tot} - 1)\hat{X}_{2tot}^2 + \hat{X}_{2tot}^4 + 1 < 0$$

(9)

and

$$(P_{1eq}^4 + 2P_{1eq}^2 + \hat{X}_{2tot}^2 + 1)^2 - 4P_{1eq}(P_{1eq}^2 + 1)\hat{X}_{1tot}\hat{X}_{2tot}^2 < 0$$

(10)

for the MD and DD toggles, respectively. For each bistable solution set found we substituted the values $\hat{X}_{1tot}, \hat{X}_{2tot}$, and $P_{1eq}$ into Eqs. (9) and (10) and confirmed that only one of the three solutions satisfies the appropriate instability condition.

It is worth noting that the time required to test $(\hat{X}_{1tot}, \hat{X}_{2tot})$ pairs scales linearly with the number of pairs, so while testing a small number can be done relatively quickly, as the number of pairs becomes appreciable the time can be significant—up to 3 hours to test 600,000 random values of $\hat{X}_{1tot}$ and $\hat{X}_{2tot}$.
in [22] we have that the steady state concentration of protein satisfies
\[\gamma \sigma_1 \sigma_2 P_{eq}^7 + \gamma \sigma_1 P_{eq}^6 - \alpha \sigma_1 P_{eq}^4 + \gamma P_{eq}^3 - P_{eq}^2 + \gamma P_{eq} - 1 = 0,\] (11)
where \(\gamma\) is the rescaled degradation rate constant, \(\alpha\) represents the increase in protein production resulting from dimer binding to OR2, and \(\sigma_1\) and \(\sigma_2\) are the relative (to OR1) affinities for OR2 and the negatively-regulating OR3, respectively. (For simplicity we set the gene copy number equal to one.) With \(\sigma_1 = 2\) and \(\sigma_2 = 0.08\) [22], the associated Sturm sequence \(F_{P_{RM}}\) has only two inequality sets with \(\text{var}(F_{P_{RM}}, 0) = 5\) and one set with \(\text{var}(F_{P_{RM}}, +\infty) = 2\) that are logically consistent and together give bistability (Table 3).

<table>
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Table 3: Sturm sequence inequality sets for the \(\lambda\) repressor–P\(_{RM}\) system. The first four Sturm polynomials have fixed signs at \(P_{eq} = 0\) and \(P_{eq} = +\infty\). Neither \(\text{var}(F_{P_{RM}}, 0) = 6\) nor \(\text{var}(F_{P_{RM}}, +\infty) = 1\) represent logically consistent sets.

We can compare the bistability region of the multi-operator P\(_{RM}\) system with that of a simple positive feedback circuit consisting of a dimeric repressor and only one operator site (MBN \(kqw\) in [13]). Rescaled as in Eq. (11), we have:
\[\gamma P_{eq}^3 - \alpha P_{eq}^2 + \gamma P_{eq} - 1 = 0.\] (12)
As with the MD toggle, this polynomial also has a maximum possible variation of 3 and thus only a single combination of inequalities that give rise to bistability, in the region given by Eq. (S27).

The bistable regions (in \(\alpha–\gamma\) space) for these single gene circuits are shown in Fig. 2A. (The one-dimensional regions of bistability shown in bifurcation diagrams in [22] and [23] are plotted for comparison). It can be seen that the \(\lambda\) repressor circuit is bistable over a larger range and with lower values of the degradation rate constant. Interestingly, with \(\alpha = 11\) ( [22] and references therein) the single operator circuit would just barely function as a bistable circuit, and any small fluctuation in circuit parameters would render it nonfunctional.

We can also use our method to determine how the strength of the negative feedback (\(\sigma_2\)) affects bistability. Keeping \(\sigma_1 = 2\), and using \(\alpha = 11\) and \(\gamma = 4.5\) (centered in the bistable region at \(\alpha = 11\); see Fig. 2A), we find that \(\sigma_2\) can increase twelve-fold to \(\sim 0.96\) before bistability is lost. In general, significant increases in \(\sigma_2\) require similar increases in \(\alpha\) for bistability to be maintained, with the range of allowable \(\gamma\) narrowing as a result (Fig. 2B).

### 6.3 Purely-monomeric toggle circuits

In a recent study [17] it was shown that a toggle variant consisting of two monomeric repressors derived from TALE protein DNA-binding domains cannot support bistability, but that the introduction of positive feedback in the form of two TALE-based activators that compete with the repressors for promoter access.
Figure 2: (A) Bistable regions for the single operator positive feedback circuit and the \(\lambda\) repressor–\(P_{RM}\) system with relative affinities \(\sigma_1 = 2\) and \(\sigma_2 = 0.08\). Vertical dashed lines indicate regions of bistability shown in the single-parameter bifurcation diagrams in [22] and [23]. (B) Region of bistability for the \(\lambda–P_{RM}\) system as a function of protein production enhancement \(\alpha\), protein degradation rate \(\gamma\), and relative affinity for OR3 \(\sigma_2\).

makes the system bistable. However, as we show below, it is not necessary to have two symmetric positive feedback loops for bistability; a single activator added to the monomeric toggle is sufficient.

We begin by combining components from the “mutual repressor” and “competitive feedback” models of [17] so that the two types of promoters—one containing a single binding site for both an activator and a repressor, and a second that is only responsive to a repressor—are present in one hybrid circuit:

\[
\frac{k_K}{1 + [T_{BK}]} + k_b - k_d [T_{AK}] = 0
\]

\[
\frac{k_K [T_{AV}]}{1 + [T_{AK}] + [T_{AV}]} + k_b - k_d [T_{BK}] = 0
\]

\[
\frac{k_V [T_{AV}]}{1 + [T_{AK}] + [T_{AV}]} + k_b - k_d [T_{AV}] = 0
\]

In keeping with the nomenclature of [17], \(T_{AV}\) represents the concentration of the activator species, and \([T_{AK}]\) and \([T_{BK}]\) are the concentrations of the two repressor species. (For simplicity we have not included terms related to the fluorescent reporters as they have no effect on the capacity for bistability.) So that our results may be easily comparable to those in [17], we include a basal rate \(k_b\) to account for expression from the pristinamycin- and erythromycin-inducible promoters incorporated in the original system for external control. Other parameters of the models in [17] set to either zero or one in their simulations are, for clarity of presentation, also set to those same values here.
Eqs. (13) may be combined into a single equilibrium polynomial in $[T_{AK}]$:

$$
\left(k_d^3(k_d + k_b)k_K\right)[T_{AK}]^3
+ \left(k_d^2(k_d^2(k_K - k_V) - k_d(k_d^2 + 2k_bk_V + k_Kk_V) - k_b(k_K(k_V - k_K) + k_b(k_K + k_V)))\right)[T_{AK}]^2
+ \left(k_d(k_d^2(2k_K(k_V - k_K) - k_b(k_K - 2k_V)) + k_d(4k_bk_Kk_V + k_d^2k_Kv + k_d^2(4k_V - 3k_K))
- k_d^2(k_K(k_K - 2k_V) + 2k_b(k_K - k_V))))\right)[T_{AK}]
+ k_b^3(k_b + k_K)(k_K - k_V) + k_d^2(k_b + k_K)(k_b(k_K - k_V) - k_Kk_V) + k_dk_b(2k_b + k_K)(k_b(k_K - k_V) - k_Kk_V) = 0.
$$

(14)

The maximum possible variation that the Sturm sequence associated with Eq. (14) may have is 3 and there is thus only a single combination of inequalities that can yield three steady states.

To establish the relationship between the repressor and activator expression strengths ($k_K$ and $k_V$, respectively) and the capacity for bistability, we set $k_b$ to the uninduced steady-state value of $\sim 0.04 \text{ M} \cdot \text{h}^{-1}$ and the degradation rate constant $k_d = 0.1 \text{ h}^{-1}$ (values taken from [17]). Interestingly, we find two unconnected subregions in which the system exhibits bistability: one in which the repressor promoter strength must be greater than $10 \times$ that of the activator, and another in which the activator promoter strength must be greater than $\sim 2 \times$ that of the repressor (Fig. 3). Thus, while a purely monomeric toggle without positive feedback cannot be bistable—indeed, the equilibrium polynomial for such a circuit is only second order and could never have more than two real roots—the addition of a single activator with competitive feedback is all that is needed for bistability. Also noteworthy is that the values of $k_K$ and $k_V$ that give bistability in the simulation of the full “competitive feedback” model of [17] (180 h$^{-1}$ and 30 h$^{-1}$, respectively) only give a monostable response in this hybrid circuit.

![Figure 3](image-url)
6.4 Sturm’s theorem and the Routh-Hurwitz stability criterion: a comparison of analytical methods

The Routh-Hurwitz stability criterion is commonly employed by control theorists to test if a polynomial admits positive roots. Typically applied to characteristic polynomials of linear time invariant systems to establish their stability, the method can also be used to find the parameter range in which a polynomial equilibrium condition admits a desired number of positive roots. Indeed, Sturm’s theorem and the Routh-Hurwitz method are related, and the validity of the latter can be demonstrated using Sturm chains. It may be argued that building a Routh table is a simpler procedure than applying Sturm’s theorem since it only requires basic arithmetic operations on the polynomial coefficients; however, its relative simplicity comes at a price: information about the real or complex nature of the positive roots is lost. In contrast, Sturm’s theorem allows for an exact determination of the number of real positive roots, as well as whether or not they are physically meaningful for a biochemical system.

A recent study of a novel biomolecular circuit that uses modulation of enzyme activity by inhibitory RNA aptamers to achieve bistability [19] provides us with an opportunity to compare our approach with one based on the Routh-Hurwitz. Interestingly, like the monomeric circuit from [17] and its asymmetric variant described above, and proposed MBN \textit{bcdh} in [13], this aptamer-based circuit achieves bistability with monomeric regulation.

The equilibrium condition for the concentration of one of the enzyme species in its active form (given here by $x$; see Supporting Information) is a complicated polynomial of fourth order:

$$
(\gamma^2 \kappa^2 - \beta^2 \gamma^2)x^4 + (\beta \gamma \delta \kappa - 2 \beta \gamma^2 \delta + \gamma \delta \kappa^2 + 2 \beta^2 \gamma^2 E_{tot} - \beta \gamma^2 E_{tot} \kappa - \gamma^2 E_{tot} \kappa^2)x^3 \\
+ (\beta \gamma^2 \kappa - \beta^2 \gamma^2 (E_{tot})^2 + \beta \gamma^2 (E_{tot})^2 \kappa + 4 \beta \gamma \delta E_{tot} - 2 \beta \gamma \delta E_{tot} \kappa - \beta^2 \delta^2)x^2 \\
+ (\beta \gamma \delta (E_{tot})^2 \kappa - 2 \beta^2 \gamma \delta (E_{tot})^2 + 2 \beta^2 \delta^2 E_{tot} - \beta \delta^2 E_{tot} \kappa)x - \beta^2 \delta^2 (E_{tot})^2 = 0 ,
$$

(15)

where $E_{tot}$ is the total concentration of each of the two enzymes used in the circuit and Greek letters indicate reaction rates.

The inequality sets for this circuit are listed along with their allowabilities in Table 6.4. Combining these inequalities we find that for bistability it is necessary that

$$
E_{tot} > \frac{\delta}{\gamma} , \\
\kappa > \frac{\beta \delta^2 + \beta \gamma^2 (E_{tot})^2 + 2 \beta \gamma \delta E_{tot}}{\delta^2 + \gamma^2 (E_{tot})^2 - 2 \gamma \delta E_{tot}}.
$$

(16)

These conditions are identical to those identified in [19] by applying the Routh-Hurwitz table together with imposing conditions to guarantee real positive roots and conditions on the constant term of the characteristic polynomial to achieve bistability. Additional details are provided in the Supporting Information.

<table>
<thead>
<tr>
<th>( var(\mathcal{F}, 0) )</th>
<th>( sign(\mathcal{F}, 0) )</th>
<th>Allowed?</th>
<th>( var(\mathcal{F}, +\infty) )</th>
<th>( sign(\mathcal{F}, +\infty) )</th>
<th>Allowed?</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>{− + − +}</td>
<td>N</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>{− − + +}</td>
<td>Y</td>
<td>0</td>
<td>{++ + + +}</td>
<td>Y</td>
</tr>
</tbody>
</table>

Table 4: Sturm sequence inequality sets for the RNA aptamer-based circuit in [19]. The sign of the first polynomial is fixed at \( x \to 0 \). \( var(\mathcal{F}, 0) = 4 \) is not a logically consistent set.
7 Discussion

In this work we have for the first time applied Sturm’s theorem to the analysis of biological circuits. We take advantage of the fact that the bistable circuits considered here are dissipative, positive in their linearization, and have characteristic polynomials with all non-constant terms positive, properties which allows us to ascertain the stability of their equilibria without computation. It is important to emphasize that these are not unique features of these particular circuits and that many real systems share these properties. And for any such system, once the number of steady-states is determined—by Sturm’s theorem or by other methods—the stabilities are known as well.

So following our analysis, can it be said that some topologies are in some sense “better” than others? Between the two different genetic toggle variants, we see that when rate parameters are fixed a toggle consisting of two dimeric repressor species functions as a bistable switch over a wider range of DNA concentrations than one composed of one dimeric and one monomeric repressor. This result provides a strong motivation for choosing a DD toggle over a MD toggle in any application where there is considerable uncertainty or variability in parameter values or DNA concentrations (e.g., when DNA is in the form of plasmids without strict copy-number control). In single-gene positive feedback circuits exhibiting bistability, we see benefits to having additional operator sites: without both OR1 and OR2 in the λ repressor–P\text{RM} system, the enhancement $\alpha = 11$ would barely be sufficient for bistability. Interestingly, although feedback at OR3 is negative, it is not strong enough to significantly affect the circuit’s ability to function as a bistable switch. Taken together, these two results suggest that the promoter architecture of the λ system may have evolved to allow for both robust bistability due to the positive feedback as well as reduced variability or other benefit of the small negative autoregulation.

Though relatively little-known within the biological sciences, Sturm’s theorem has found applicability in a number of other areas where polynomials play an important role, including computational mathematics [24], dynamical systems [25, 26], robotics [27], and finance [28]. Additional biological applications are also possible, including as a tool to predict new bistable topologies or rule out those that do not have the capacity for bistability (like Chemical Reaction Network Theory, previously [13, 23]). Other methods from algebraic geometry have also found use in biological systems analysis, with recent applications to model discrimination [29] and the study of chemical reaction networks [30, 31] and multisite phosphorylation systems [32], among many others. (This latter paper is particularly relevant to our own work, as it also highlights the benefits of treating parameters symbolically rather than numerically.)

We note that not all bistable circuits of interest may be easily studied using our approach. Firstly, it may not be possible to capture the circuit’s equilibrium state with a single polynomial. When an equilibrium polynomial can be found, those of third-order are the simplest to study since there is only one combination of inequalities that can give rise to bistability. As the order increases, the number of inequality sets to test and the complexity of the Sturm sequences can also increase to the point where application of the theorem is impractical. However, this is highly dependent on the complexity of the polynomial coefficients and the number of Sturm sequences of fixed sign. Indeed, as we demonstrated with our analysis of the λ repressor–P\text{RM} system, even a seventh-order polynomial can be tractable. For particularly large circuits for which an equilibrium polynomial cannot be derived, or for which the Sturm sequence is overly complex, a decomposition of the system into smaller modules can be beneficial.

The analysis of an equilibrium polynomial with a zero root—for example, that describing a recently-developed DNA-based circuit that achieves bistability through a combination of mutual inhibition and autocatalysis [33]—presents challenges, since Sturm’s theorem requires that neither of the limits of the region of interest (in our case, 0 and $+\infty$) be roots. Under these circumstances one could choose an $\epsilon < 0$ for the lower limit; however, in this case there is no guarantee that the three roots found by Sturm’s theorem will all represent physically real, positive quantities (since one or more of the roots could in theory
lie between $\epsilon$ and 0).

### 7.1 Extensions to Sturm’s theorem and related methods

The Sturm’s theorem approach described here can be directly applied to systems that at steady state are described by univariate polynomials with integer exponents and simple roots. While such systems are common, examples of equilibrium polynomials not of this particular form can also be found, and for many of these systems there are extensions or modifications to the described approach that may be used. For example, there are simple extensions to Sturm’s theorem that deal with multiple roots (e.g., [34]) and multivariate models (e.g., [12, 35]). And in limited cases a generalised polynomial with fractional exponents can be turned to a proper polynomial with a simple substitution that does affect the number of zeros (e.g., $u = x^{1/N}$, if all exponents of $x$ are multiples of $1/N$). This last class of systems is perhaps most biologically relevant, since such generalised polynomials may result when functional relationships are modeled with Hill functions fit to empirical data [36–38]. (Integer exponents are typically used in simplified explanatory models and to describe multiple ligand binding reactions with a high degree of positive cooperativity [39]—a situation which may be considered equivalent to strong transcription factor multimerization taking place away from the regulated promoter, as in this work).

Multivariate [40, 41] and generalised polynomial [42] extensions to the related Descartes’ rule of signs are also available. (It is important to note that although the Descartes’ rule of signs method may be simpler to apply than Sturm’s theorem, it is definitively less powerful in that (1) it can only give an upper bound on the number of real roots of a polynomial, and (2) the multivariate version has been shown to be applicable only to certain classes of systems.)

### 8 Acknowledgements

A large number of people contributed to this work with insights and comments. The authors would like to particularly thank Andras Gyorgy, Yutaka Hori, Scott C. Livingston, Anne Shiu, Eduardo Sontag, Elisenda Feliu, Zvi H. Rosen, Jaap Top, and Brian Ingalls. This research is funded in part by the National Science Foundation through Grant CMMI 1266402, and the Gordon and Betty Moore Foundation through Grant GBMF2809 to the Caltech Programmable Molecular Technology Initiative.

### References


S1 Equilibrium polynomial derivation for toggle variants and single-operator positive feedback circuit

The monomer-dimer toggle, dimer-dimer toggle, and single-operator positive feedback circuits were initially predicted to exhibit bistability using a chemical reaction network (CRN)-based topological survey [1]. Each of these CRNs contains reactions representing basal protein production and degradation:

\[
X_1 \xrightarrow{k_{bas1}} X_1 + P_1, \quad X_2 \xrightarrow{k_{bas2}} X_2 + P_2, \quad P_1 \xrightarrow{k_{deg1}} \emptyset, \quad P_2 \xrightarrow{k_{deg2}} \emptyset
\]

for genes \(X_i\) and proteins \(P_i\). The other reactions that uniquely define each circuit are:

\[
\begin{align*}
X_2 + P_1 & \xrightarrow{k_{cF}} X_2P_1 \\
2P_2 & \xrightarrow{k_{kF}} P_2P_2 \\
X_1 + P_2 & \xrightarrow{k_{nF}} X_1P_2P_2 \\
2P_1 & \xrightarrow{k_{oF}} P_1P_1 \\
2P_2 & \xrightarrow{k_{qF}} P_2P_2 \\
X_2 + P_2 & \xrightarrow{k_{iF}} X_2P_2P_2 \\
X_1 + P_2P_2 & \xrightarrow{k_{nF}} X_1P_2P_2 \\
X_2 + P_1 & \xrightarrow{k_{oF}} X_2P_1P_1 \\
X_2P_2P_2 & \xrightarrow{k_{w}} X_2P_2P_2 + P_2
\end{align*}
\]

\(P_iP_j\) represent dimeric species, and \(X_iP_j\) and \(X_iP_jP_j\) represent monomers and dimers bound to the gene promoters. The various ODE sets were derived from these CRNs under the assumption of mass action kinetics and simplified using the fact that the total concentrations of each gene (in bound and unbound form) are conserved.
From these CRN formulations and assuming mass-action kinetics we can derive the following sets of ordinary differential equations (ODEs) that describe the circuit dynamics. For the MD toggle:

\[ P_1'(t) = -k_{de1}P_1(t) - k_{cF}X_2(t)P_1(t) + k_{bas1}X_1(t) + k_{cR}(X_{2tot} - X_2(t)) \]  
\[ P_2'(t) = 2k_{d2}P_2(t)^2 - k_{de2}P_2(t) + 2k_{cR}P_2(t) + k_{bas2}X_2(t) \]  
\[ P_2'(t) = k_{kF}P_2(t)^2 - k_{kR}P_2(t) - k_{nF}P_2(t)X_1(t) + k_{nR}(X_{1tot} - X_1(t)) \]  
\[ X_1'(t) = k_{nR}(X_{1tot} - X_1(t)) - n_{FR}P_2(t)X_1(t) \]  
\[ X_2'(t) = k_{cR}(X_{2tot} - X_2(t)) - k_{cF}P_1(t)X_2(t) \]  

the DD toggle:

\[ P_1'(t) = -2k_{iF}P_1(t)^2 - k_{de1}P_1(t) + 2k_{iR}P_1(t) + k_{bas1}X_1(t) \]  
\[ P_1'(t) = k_{iF}P_1(t)^2 - k_{iR}P_1(t) - k_{oF}P_1(t)X_2(t) + k_{oR}(X_{2tot} - X_2(t)) \]  
\[ P_2'(t) = -2k_{kF}P_2(t)^2 - k_{de2}P_2(t) + 2k_{kR}P_2(t) + k_{bas2}X_2(t) \]  
\[ P_2'(t) = k_{kF}P_2(t)^2 - k_{kR}P_2(t) - k_{nF}P_2(t)X_1(t) + k_{nR}(X_{1tot} - X_1(t)) \]  
\[ X_1'(t) = k_{nR}(X_{1tot} - X_1(t)) - n_{FR}P_2(t)X_1(t) \]  
\[ X_2'(t) = k_{oR}(X_{2tot} - X_2(t)) - k_{oF}P_1(t)X_2(t) \]  

and single-operator positive feedback circuit:

\[ P_2'(t) = k_{bas2}x_2(t) - k_{de2}P_2(t) - 2k_{kF}P_2(t)^2 + 2k_{kR}P_2(t) + k_{w}(X_{2tot} - X_2(t)) \]  
\[ P_2'(t) = k_{kF}P_2(t)^2 - k_{kR}P_2(t) - k_{nF}P_2(t)X_1(t) + k_{nR}(X_{2tot} - X_2(t)) \]  
\[ X_2'(t) = k_{qR}(X_{2tot} - X_2(t)) - k_{qF}P_2(t)X_2(t) \]  
\[ X_2'(t) = k_{oR}(X_{2tot} - X_2(t)) - k_{oF}P_1(t)X_2(t) \]  

where the variable names are as in [1]: \( X_i \) is the concentration of free (i.e., unbound by repressor) gene \( i \), \( X_{i\text{tot}} \) is the total amount of \( X_i \) in the circuit (bound and unbound), and \( P_i \) and \( P_iP_i \) represent the monomeric and dimeric forms of protein \( i \), respectively. The various \( k_i \) are the reaction rates, and in the positive feedback circuit, \( k_w \) is the repressor binding rate, assumed.

We derive the equilibrium polynomials from (S1), (S2), and (S3) by first setting the left-hand sides equal to zero. For the DD toggle, we subtract (S2d) from (S2e):

\[ 0 = k_{nR}(X_{1tot} - X_1) - k_{nF}P_2P_2X_1 \]
\[ -k_{kF}P_2^2 + k_{kR}P_2P_2 + k_{nF}P_2P_2X_1 - k_{nR}(X_{1tot} - X_1) \]
\[ \Rightarrow P_2P_2 = k_{kF}P_2^2 = k_{kR}P_2^2 = \frac{P_2^2}{k_{kD}} \]  

where for simplicity of notation we use \( X_i \), \( P_i \), and \( P_iP_i \) to mean the equilibrium concentrations. We then plug this expression into (S2e) to get

\[ 0 = k_{nR}(X_{1tot} - X_1) - k_{nF}(P_2^2/k_{kD})X_1 \]
\[ = (X_{1tot} - X_1) - \frac{(P_2^2/((k_{kD}k_{nD})X_{1tot}))X_1}{k_{kD}k_{nD}X_{1tot}} \]
\[ \Rightarrow X_1 = \frac{k_{kD}k_{nD}X_{1tot}^2}{k_{kD}k_{nD}X_{1tot} + P_2^2} \]  

Similarly we subtract (S2b) from (S2f)

\[ 0 = k_{oR}(X_{2tot} - X_2) - k_{oF}P_1P_1X_2 \]
\[ -k_{iF}P_1^2 + k_{iR}P_1P_1 + k_{oF}P_1P_1X_2 - k_{oR}(X_{2tot} - X_2) \]
\[ \Rightarrow P_1P_1 = \frac{k_{iF}P_1^2}{k_{iR}} = \frac{P_1}{k_{iD}} \]
and plug the resulting expression back into (S2f) to get
\[
0 = k_{oR}(X_{2\text{tot}} - X_2) - k_{oF}(P_2^2/k_{iD}) \cdot X_2 \\
= (X_{2\text{tot}} - X_2) - (P_2^2/(k_{iD}k_{oD})) \cdot X_2 \\
\implies X_2 = \frac{k_{iD}k_{oD}X_{2\text{tot}}}{k_{iD}k_{oD} + P_2^2}.
\] (S7)

Substituting Eqs. (S4)–(S7) into (S2a) and (S2c) gives the equilibrium concentrations of \(P_1\) and \(P_2\) in the DD toggle shown in the main text:

\[
P_{1eq} = \left(1 + \left(P_{2eq}/K_2\right)^2\right)^{-1} \beta_1 X_{1\text{tot}}, \quad P_{2eq} = \left(1 + \left(P_{1eq}/K_{dd}\right)^2\right)^{-1} \beta_2 X_{2\text{tot}},
\] (S8)

where \(\beta_i = k_{basi}/k_{degi}\) is the ratio of basal production rate to degradation rate for protein \(i\), and the Michaelis constants \(K_{dd} = (k_{iD}k_{oD})^{1/2}\) and \(K_2 = (k_{kD}k_{nD})^{1/2}\) represent the protein concentrations that yield 50% of the maximum production rate of their respective targets. The combination of these two expressions, with rescaling as described in the main text, gives the DD equilibrium polynomial.

For the MD toggle, we have again that
\[
P_2P_2 = \frac{P_2^2}{k_{kD}}
\] (S9)

and
\[
X_1 = \frac{k_{kD}k_{nD}X_{1\text{tot}}}{k_{kD}k_{nD} + P_2^2},
\] (S10)

but also (from Eq. (S1e)) that
\[
X_2 = \frac{k_{cD}X_{2\text{tot}}}{k_{cD} + P_1}.
\] (S11)

Substituting Eqs. (S9)–(S11) into (S1a) and (S1b) gives the equilibrium concentrations of \(P_1\) and \(P_2\) in the MD toggle shown in the main text:

\[
P_{1eq} = \left(1 + \left(P_{2eq}/K_2\right)^2\right)^{-1} \beta_1 X_{1\text{tot}}, \quad P_{2eq} = \left(1 + \left(P_{1eq}/K_{md}\right)^2\right)^{-1} \beta_2 X_{2\text{tot}}.
\] (S12)

As previously, \(\beta_i\) is the ratio of basal production rate to degradation rate for protein \(i\), and the Michaelis constants \(K_{md} = k_{cD}\) and \(K_2 = (k_{kD}k_{nD})^{1/2}\) represent the protein concentrations that yield 50% of the maximum production rate of their respective targets. The combination of these two expressions, again with rescaling as described in the main text, gives the MD equilibrium polynomial.

The equilibrium polynomial for the single-operator positive feedback circuit may be similarly derived.

### S2 Aptamer-based bistable system with monomeric inhibition

The system described in [2] is composed by two enzymes \(E_1\) and \(E_2\) which transcribe at a constant rate two RNA aptamers \(R_1\) and \(R_2\). The aptamers mutually inhibit the enzymes; enzyme activity is recovered and aptamers are degraded with first order reaction rates. The reactions are:

\[
E_i + g_i \xrightarrow{k_i} E_i + R_i + g_i
\] (S13)

\[
R_i \xrightarrow{\delta} 0
\] (S14)

\[
E_i + R_j \xrightarrow{\gamma} E_i^*
\] (S15)

\[
E_i^* \xrightarrow{\beta} E_i
\] (S16)
for \( i = 1,2 \) and \( j = 2,1 \), where \( E_i \) are active enzymes, \( E_i^* \) are inactive enzymes, \( R_i \) are RNA species, \( g_i \) are genes (constant). We assume the fraction of enzyme bound to its gene substrate is negligible. Using mass action laws, we can derive the ODEs describing the dynamics of the system:

\[
[R_i]' = k[E_i][g_i] - \delta[R_i] - \gamma[E_i][R_i] \\
[E_i]' = \beta([E_i^{\text{tot}}] - [E_i]) - \gamma[E_i][R_j]
\]

We switch to a more compact notation, and define \( x_1 = [R_1], x_2 = [E_1], x_3 = [R_2], \) and \( x_4 = [E_2] \). Because the concentration of genes is constant, we define \( \kappa_1 = k g_1 \) and \( \kappa_2 = k g_2 \), and we assume \( \kappa_1 = \kappa_2 = \kappa \). The system of ODEs becomes:

\[
\begin{cases}
  x'_1 = \kappa x_2 - \delta x_1 - \gamma x_4 x_1 \\
  x'_2 = \beta x_2^{\text{tot}} - x_2 - \gamma x_2 x_3 \\
  x'_3 = \kappa x_4 - \delta x_3 - \gamma x_2 x_3 \\
  x'_4 = \beta x_4^{\text{tot}} - x_4 - \gamma x_4 x_1
\end{cases}
\]

Equilibrium conditions are derived in detail in [2] from these expressions:

\[
\begin{align*}
\beta \gamma x_1^2 + (\gamma \kappa x_2 - \beta \gamma x_4^{\text{tot}} + \beta \delta) x_4 - \beta \delta x_4^{\text{tot}} &= 0, \\
\beta \gamma x_2^2 + (\gamma \kappa x_4 - \beta \gamma x_2^{\text{tot}} + \beta \delta) x_2 - \beta \delta x_2^{\text{tot}} &= 0.
\end{align*}
\]

From here we can derive a fourth order polynomial in either \( x_2 \) or \( x_4 \):

\[
(\gamma^2 \kappa^2 - \beta^2 \gamma^2)x^4 + (\beta \gamma \delta \kappa - 2 \beta^2 \gamma \delta + \gamma \delta \kappa^2 + 2 \beta^2 \gamma^2 E^{\text{tot}} - \beta \gamma \delta E^{\text{tot}} \kappa - \beta \gamma \delta E^{\text{tot}} \kappa - \gamma^2 E^{\text{tot}} \kappa^2)x^3 \\
+ (\beta \delta^2 \kappa - \beta^2 \gamma^2 (E^{\text{tot}})^2 + \beta \gamma^2 (E^{\text{tot}})^2 \kappa + 4 \beta^2 \gamma \delta E^{\text{tot}} - 2 \beta \gamma \delta E^{\text{tot}} \kappa - \beta^2 \delta^2)x^2 \\
+ (\beta \gamma \delta (E^{\text{tot}})^2 \kappa - 2 \beta^2 \gamma \delta (E^{\text{tot}})^2 + 2 \beta^2 \delta^2 E^{\text{tot}} - \beta \delta^2 E^{\text{tot}} \kappa - \beta^2 \delta^2 (E^{\text{tot}})^2) x + \beta^2 \delta^2 (E^{\text{tot}})^2 = 0,
\]

where we have used \( x \) to represent the steady-state concentration of an enzyme species in its active form and have put \( x_2^{\text{tot}} = x_4^{\text{tot}} = E^{\text{tot}} \) (as in [2]).

The inequality sets for this circuit along are listed along with their allowabilities in the main text (Table ??), as are the conditions on the parameters that yield bistability.

## S3 Sturm polynomials

Sturm polynomials can be rather long and complicated functions; however, they can be easily generated using software capable of symbolic manipulation (e.g., Mathematica). We present here the Sturm sequences (in \( x \), for simplicity of notation) for the MD toggle, DD toggle, and single-operator positive feedback circuits.

The Sturm polynomials associated with the MD toggle equilibrium polynomial are:

\[
\begin{align*}
 f_0(x) &= (x - \tilde{X}_{1\text{tot}})(x + 1)^2 + x \tilde{X}_{2\text{tot}}^2 \\
 f_1(x) &= \tilde{X}_{2\text{tot}}^2 + (x + 1)(3x - 2\tilde{X}_{1\text{tot}} + 1) \\
 f_2(x) &= \frac{1}{9} \left( 2(x + 1)(\tilde{X}_{1\text{tot}} + 1)^2 - (6x + \tilde{X}_{1\text{tot}} - 2)\tilde{X}_{2\text{tot}}^2 \right) \\
 f_3(x) &= \frac{9\tilde{X}_{2\text{tot}}^2 - 4\tilde{X}_{2\text{tot}}^4 + (\tilde{X}_{1\text{tot}}(\tilde{X}_{1\text{tot}} + 20) - 8)\tilde{X}_{2\text{tot}} - 4(\tilde{X}_{1\text{tot}} + 1)^3}{4((\tilde{X}_{1\text{tot}} + 1)^2 - 3\tilde{X}_{2\text{tot}}^2)^2}.
\end{align*}
\]
The DD Sturm polynomials are:

\[ f_0(x) = (x^2 + 1)^2(x - \hat{X}_{1tot}) + x\hat{X}_{2tot}^2 \]
\[ f_1(x) = (x^2 + 1)(5x^2 - 4x\hat{X}_{1tot} + 1) + \hat{X}_{2tot}^2 \]
\[ f_2(x) = \frac{1}{25}
    \left( 4(x^2 + 1)(x(\hat{X}_{1tot}^2 - 5) + 6\hat{X}_{1tot}) - \hat{X}_{2tot}^2(20x + \hat{X}_{1tot}) \right) \]
\[ f_3(x) = \frac{1}{q_3} \left( \hat{X}_{2tot}^2 \left( 2\hat{X}_{1tot}^2 - 4 + 4x(\hat{X}_{1tot}^2 - 5) - 3x\hat{X}_{1tot}(3 + \hat{X}_{1tot}^2) \right) - 4(x^2 + 1)(\hat{X}_{1tot}^2 + 1)^2 \right) \]
\[ f_4(x) = \frac{1}{q_4} \left( 4(\hat{X}_{1tot}^2 - 5)^2\hat{X}_{2tot}^2(20x + \hat{X}_{1tot}) - (\hat{X}_{1tot}^2 - 5)^2\hat{X}_{2tot}^4(x(9\hat{X}_{1tot}^4 + 35\hat{X}_{1tot}^2 - 64) - 2\hat{X}_{1tot}(\hat{X}_{1tot}^2 - 62)) \right. \\
\left. + 16(\hat{X}_{1tot}^4 - 4\hat{X}_{1tot}^2 - 5)^2\hat{X}_{2tot}^2(\hat{X}_{1tot}^2 - x) \right) \]
\[ f_5(x) = \frac{1}{q_5} \left( 256\hat{X}_{1tot}^6 - 3\hat{X}_{1tot}^4(9\hat{X}_{2tot}^4 + 32\hat{X}_{2tot}^2 - 256) - 96\hat{X}_{1tot}^2(\hat{X}_{2tot}^4 + 29\hat{X}_{2tot}^2 - 8) + 256(\hat{X}_{2tot}^2 + 1)^3 \right) \\
\times 25 \left( (\hat{X}_{1tot}^2 + 1)^2 + (\hat{X}_{1tot}^2 - 5)\hat{X}_{2tot}^2 \right)^2 \]

where

\[ q_3 = \frac{4}{25} (\hat{X}_{1tot}^2 - 5)^2 \]
\[ q_4 = 100 \left( (\hat{X}_{1tot}^2 - 5)\hat{X}_{2tot}^2 + (\hat{X}_{1tot}^2 + 1)^2 \right)^2 \]
\[ q_5 = (\hat{X}_{1tot}^2 - 5)^2 \left( 16(\hat{X}_{1tot}^2 + 1)^2 + (9\hat{X}_{1tot}^4 + 35\hat{X}_{1tot}^2 - 64)\hat{X}_{2tot}^2 - 80\hat{X}_{2tot}^4 \right)^2 \]

For all values of \( \hat{X}_{1tot} \neq \sqrt{5} \), the sequence consisting of the \( f_i(x) \) above may be used to determine the number of steady states. However, when \( \hat{X}_{1tot} \to \sqrt{5} \), the sequence terminates prematurely (since \( f_4(x) \to 0 \)) and there are problematic zeroes in the denominators of \( f_3(x) \) and \( f_5(x) \). We thus set \( \hat{X}_{1tot} = \sqrt{5} \) in the equilibrium polynomial to get

\[ \hat{P}_{1eq}^5 - \sqrt{5}\hat{P}_{1eq}^3 + 2\hat{P}_{1eq} - 2\sqrt{5}\hat{P}_{1eq}^2 + \hat{P}_{1eq}\hat{X}_{2tot}^2 + \hat{P}_{1eq} - \sqrt{5} = 0 \tag{S23} \]

and generate a second Sturm sequence to use at \( \hat{X}_{1tot} = \sqrt{5} \) only:

\[ f_0(x) = x^5 - \sqrt{5}x^4 + 2x^3 - 2\sqrt{5}x^2 + x\hat{X}_{2tot}^2 + x - \sqrt{5} \]
\[ f_1(x) = 5x^4 - 4\sqrt{5}x^3 + 6x^2 - 4\sqrt{5}x + \hat{X}_{2tot}^2 + 1 \]
\[ f_2(x) = \frac{1}{25} \left( 24\sqrt{5}x^2 - 20x\hat{X}_{2tot}^2 - \sqrt{5}\hat{X}_{2tot}^2 + 24\sqrt{5} \right) \]
\[ f_3(x) = -\frac{5}{1728} \left( 40\sqrt{5}x\hat{X}_{2tot}^6 - 168\sqrt{5}x\hat{X}_{2tot}^4 - 288\sqrt{5}x\hat{X}_{2tot}^2 + 10\hat{X}_{2tot}^6 - 285\hat{X}_{2tot}^4 + 1440\hat{X}_{2tot}^2 \right) \]
\[ f_4(x) = \frac{-27(256\hat{X}_{2tot}^6 - 387\hat{X}_{2tot}^4 - 15552\hat{X}_{2tot}^2 + 55296)}{40\sqrt{5}(5\hat{X}_{2tot}^4 - 21\hat{X}_{2tot}^2 - 36)^2} \]

The single-operator positive feedback circuit Sturm sequence contains the following polynomials:

\[ f_0(x) = \alpha x^2 - \gamma \left( x^3 + x \right) + 1 \]
\[ f_1(x) = 2\alpha x - \gamma \left( 3x^2 + 1 \right) \]
\[ f_2(x) = \frac{1}{9} \left( \frac{-2\alpha^2 x}{\gamma} + \alpha + 6\gamma x - 9 \right) \]
\[ f_3(x) = \frac{9\gamma \left( 4\alpha^3 - \alpha^2 \gamma^2 - 18\alpha \gamma^2 + 4\gamma^4 + 27\gamma^2 \right)}{4 (\alpha^2 - 3\gamma^2)^2} \]
Recall that $\gamma$ is the rescaled degradation rate constant for the $\lambda$ repressor and $\alpha$ represents the increase in protein production resulting from repressor dimer binding to OR2.

## S4 Analytic expressions for regions of bistability

The regions of bistability were determined by combining the valid inequality sets and reducing to a single pair of inequalities in the plane of the relevant variables. Reduction was done using Mathematica.

There is only one combination of inequalities that can give rise to bistability for the MD toggle: when $\var(F_{md}, 0) = 3$ and $\var(F_{md}, +\infty) = 0$. This gives the region of bistability in $\hat{X}_{1\text{tot}}-\hat{X}_{2\text{tot}}$ space as

$$\hat{X}_{1\text{tot}} > 8 \frac{1}{8} \left( 20\hat{X}_{1\text{tot}} + \hat{X}_{1\text{tot}}^2 - 8 - f(\hat{X}_{1\text{tot}}) \right) < \hat{X}_{2\text{tot}}^2 < \frac{1}{8} \left( 20\hat{X}_{1\text{tot}} + \hat{X}_{1\text{tot}}^2 - 8 + f(\hat{X}_{1\text{tot}}) \right),$$

(S24)

with $f(\hat{X}_{1\text{tot}}) = (\hat{X}_{1\text{tot}} - 8)^{3/2}(\hat{X}_{1\text{tot}})^{1/2}$.

In the case of the DD toggle, there are two different Sturm sequences we need to consider depending on the value of $\hat{X}_{1\text{tot}}$ (see Section ‘Sturm polynomials’ above). When $\hat{X}_{1\text{tot}} \neq \sqrt{5}$, the Sturm sequence $F_{dd}$ contains six polynomials, and there are three sets of inequalities with $\var(F_{dd}, 0) = 4$ and two with $\var(F_{dd}, +\infty) = 1$ that are logically consistent (Table ??). When $\hat{X}_{1\text{tot}} = \sqrt{5}$, the sequence $F_{dd}$ contains five polynomials, and only one set of inequalities with $\var(F_{dd}, 0) = 3$ and one with $\var(F_{dd}, +\infty) = 0$ are allowed (Table ??). These inequalities may be combined to give a continuous region of bistability as the intersection of

$$0 < \hat{X}_{2\text{tot}}^2 \leq \frac{1}{160} \left( 9\hat{X}_{1\text{tot}}^4 + 35\hat{X}_{1\text{tot}}^2 - 64 + 3 \left( 9\hat{X}_{1\text{tot}}^8 + 70\hat{X}_{1\text{tot}}^6 + 577\hat{X}_{1\text{tot}}^4 + 640\hat{X}_{1\text{tot}}^2 + 1024 \right)^{1/2} \right)$$

(S25)

and

$$256 \left( \hat{X}_{1\text{tot}}^6 + (\hat{X}_{2\text{tot}}^2 + 1)^3 \right) < 3\hat{X}_{1\text{tot}}^4 \left( 9\hat{X}_{2\text{tot}}^4 + 32\hat{X}_{2\text{tot}}^2 - 256 \right) + 96\hat{X}_{1\text{tot}}^2 \left( \hat{X}_{2\text{tot}}^4 + 29\hat{X}_{2\text{tot}}^2 - 8 \right).$$

(S26)

As with the MD toggle, the equilibrium polynomial for the single-operator positive feedback circuit also has a maximum possible variation of 3, which means that the circuit exhibits bistability only in the region

$$\alpha > 9 \frac{1}{8} \left( \alpha^2 + 18\alpha - 27 - (\alpha - 9)^{3/2}(\alpha - 1)^{1/2} \right) < \gamma^2 < \frac{1}{8} \left( \alpha^2 + 18\alpha - 27 + (\alpha - 9)^{3/2}(\alpha - 1)^{1/2} \right).$$

(S27)

With the exception of the DD toggle at $\hat{X}_{1\text{tot}} = \sqrt{5}$ (which was treated by analyzing a second Sturm sequence; see ‘Sturm polynomials’ section above), none of the potential zeroes in the Sturm polynomial denominators required special treatment nor did they present any problems in determining the regions of bistability.
S5 Number of steady states and stability analysis

S5.1 Preliminaries

**Definition 1 (Positive systems)** A linear system \( \dot{x} = Ax \) is positive if for every nonnegative initial state the solution \( x(t) \) is nonnegative.

The following is a well known condition for positivity [3]:

**Theorem 1** A linear system \( \dot{x} = Ax \) is positive if and only if matrix \( A \) is a Metzler matrix, i.e., its elements satisfy: \( a_{ij} \geq 0, \forall (i,j) \) such that \( i \neq j \).

The Jacobian matrices of all systems considered here are, for any choice of parameters, similar to Metzler matrices via linear transformations. For example, the Jacobians \( J_i \) of each of the MD toggle, DD toggle, single-operator positive feedback, and aptamer-based inhibition circuits may be transformed to Metzler matrices \( (J_M)_i = P_i^{-1} J_i P_i \), with \( P_{md} = \text{diag}(-1,1,1,-1,1) \), \( P_{dd} = \text{diag}(-1,-1,1,1,-1,1) \), \( P_{pf} = \text{diag}(1,1,-1) \), and \( P_{apt} = \text{diag}(1,1,-1,-1) \) respectively. Thus, the linearizations of the circuit ODEs are positive.

The general definition of dissipativity (see, e.g., [4]) is based on the existence of compact, forward invariant subsets of \( \mathbb{R}^n_+ \) that absorb the system trajectories. The following definition (from [5]) is equivalent and easier to verify:

**Definition 2 (Dissipative systems)** A system \( \dot{x} = f(x) \) is dissipative if its solutions are eventually uniformly bounded, i.e., there exists a constant \( k > 0 \) such that:

\[
\lim_{t \to +\infty} \sup x_i(t) \leq k.
\]

The systems analyzed in this work are all dissipative. As an example, we verify the definition for the MD toggle model (S1). Because the total mass of each of the DNA species \( X_1 \) and \( X_2 \) is constant, we know that \( X_1(t) \leq X_{\text{max}} \) and \( X_2(t) \leq X_{\text{max}} \) \( \forall t \), where \( X_{\text{max}} = \max\{X_{1\text{tot}}, X_{2\text{tot}}\} \). The concentration of \( P_1 \) can be upper bounded as follows:

\[
P_1'(t) = -k_{\text{deg} \, 1} P_1(t) - k_{cF} X_2(t) \cdot P_1(t) + k_{\text{bas} \, 1} X_1(t) + k_{cR} (X_{2\text{tot}} - X_2(t)) \\
\leq -k_{\text{deg} \, 1} P_1(t) - k_{cF} X_2(t) \cdot P_1(t) + k_{\text{bas} \, 1} X_{\text{max}} + k_{cR} X_{\text{max}} \\
\leq -a P_1(t) + b,
\]

where \( a = k_{\text{deg} \, 1} \) and \( b = (k_{\text{bas} \, 1} + k_{cR}) X_{\text{max}} \). The right hand side of the last inequality above is a linear, asymptotically stable system whose solution is eventually uniformly bounded (\( b \) is a finite constant). Using the comparison principle [6], we conclude that \( P_1(t) \) is bounded and can find a constant \( k \) that satisfies the definition.

\( P_2 \) may be similarly upper bounded. We first consider the dynamics of \( P_2'(t) = P_2(t) + 2P_2P_2(t) \), the total amount of unbound \( P_2 \) in the system:

\[
P_2'(t) = -k_{\text{deg} \, 2} P_2(t) - 2k_{nF} P_2 P_2(t) \cdot X_1(t) + 2k_{nR} (X_{1\text{tot}} - X_1(t)) + k_{\text{bas} \, 2} X_2(t) \\
\leq -k_{\text{deg} \, 2} P_2(t) - 2k_{nF} P_2 P_2(t) \cdot X_1(t) + (2k_{nR} + k_{\text{bas} \, 2}) X_{\text{max}} \\
\leq -k_{\text{deg} \, 2} P_2(t) + (2k_{nR} + k_{\text{bas} \, 2}) X_{\text{max}}.
\]
The dynamics of monomeric $P_2$ satisfy:

$$P_2'(t) = -2k_{kF}P_2(t)^2 - k_{deg} P_2(t) + 2k_{kR} P_2(t) + k_{bas} X_2(t)$$

$$\leq -k_{deg} P_2(t) + 2k_{kR} P_2(t) + k_{bas} X_{max}$$

$$\leq -k_{deg} P_2(t) + k_{kR}(P_2(t) + 2P_2(t)) + k_{bas} X_{max}.$$

Together, we have:

$$\left( \begin{array}{c} P_2'(t) \\ P_2^f'(t) \end{array} \right) \leq \left( \begin{array}{cc} -k_{deg} & k_{kR} \\ -k_{deg} & 0 \end{array} \right) \left( \begin{array}{c} P_2(t) \\ P_2^f(t) \end{array} \right) + \left( \begin{array}{c} k_{bas} X_{max} \\ (2k_{kR} + k_{bas}) X_{max} \end{array} \right).$$

The variables $P_2^f(t)$ and $P_2(t)$ are upper bounded by a linear system with eigenvalues

$$\lambda_{1,2} = \frac{1}{2} \left( -k_{deg} \pm \sqrt{k_{deg}^2 - 4k_{kR}k_{deg}} \right),$$

whose real part is always negative for any value of the (positive) parameters. Being upper bounded by an asymptotically stable linear system, the concentrations $P_2$ and $P_2^f$ are eventually uniformly bounded. It follows that $P_2P_2^f$ is also eventually uniformly bounded, since $P_2P_2^f \geq P_2^f$.

Therefore, the ODE model of the monomer-dimer toggle system is dissipative. Note that the same conclusion cannot be reached in the absence of degradation ($k_{deg} = 0$) since the total amount of protein will grow unbounded.

We can also easily show that system (S19) is dissipative: the total concentration of enzymes is constant, thus bounded at all times. The RNA species are also asymptotically bounded. The dynamics of RNA $x_1$, for instance, can be upper bounded by an asymptotically stable linear time invariant system:

$$x_1' = \kappa x_2 - \delta x_1 - \gamma x_4 x_1$$

$$\leq \kappa x_2^{tot} - \delta x_1.$$ 

We can derive a similar upper bound for $x_2'$.

The other systems considered in this work can be similarly shown to be dissipative.

### S5.2 Stability of equilibria

Sturm’s theorem applied to our circuits’ polynomial equilibrium conditions reveals that each system admits three positive equilibria. The stability properties of these equilibria can be determined by degree theory [5].

**Definition 3 (Regular equilibrium)** A point $\bar{x}$ of system $\dot{x} = f(x)$ is regular if $\text{det}(J(\bar{x})) \neq 0$ (in other words, $J(\bar{x})$ must be invertible; alternatively, $J(\bar{x})$ must not have eigenvalues at the origin).

**Definition 4 (Index of an equilibrium point)** The index of a regular equilibrium point $\bar{x}$ is the sign of the determinant of $-J(\bar{x})$:

$$\text{ind}(\bar{x}) = \text{sign}(\text{det}(-J(\bar{x})))$$
Definition 5 (Degree of a system) The degree of a dynamical system \( \dot{x} = f(x) \), over a set \( U \subseteq \mathbb{R}^n \), having equilibria \( \bar{x}_i, i = 1, \ldots, m \), is defined as:

\[
\operatorname{deg}(f) = \sum_{i=1}^{m} \{ \operatorname{ind}(\bar{x}_i), \bar{x}_i \in U, f(\bar{x}_i) = 0 \},
\]

where \( \bar{x}_i \) are regular equilibria.

Theorem 2 A dissipative dynamical system \( \dot{x} = f(x) \) defined on \( \mathbb{R}^n \) has degree +1 with respect to any bounded open set containing all its equilibria.

Since the systems are all dissipative, by Theorem 2 all have degree +1. We further note that the Jacobian matrices are row equivalent to the identity matrix (as confirmed with Mathematica) and thus always invertible—\( \det(J) \neq 0 \) for all (positive) parameters and equilibria. Therefore, all equilibria are regular. To determine the index of each equilibrium point, we need not know the value of the equilibrium itself, since in general

\[
\operatorname{ind}(\bar{x}) = \operatorname{sign} \left( \det \left( -J(\bar{x}) \right) \right) = \operatorname{sign} \left( \det \left( \lambda I - J(\bar{x}) \right) \right),
\]

with \( \lambda = 0 \).

Therefore, the index of an equilibrium corresponds to the sign of the constant term in the system’s characteristic polynomial \( p_J(\lambda, \bar{x}) = \det(\lambda I - J(\bar{x})) \).

For the Jacobians under consideration, the sign of the constant term in the characteristic polynomial also determines the stability properties of the corresponding equilibrium; we can state the following lemma [2]:

Lemma 1 Any single equilibrium \( \bar{x} \) of a system under consideration in this work is unstable if and only if the constant term of the characteristic polynomial \( p_J(\lambda, \bar{x}) \) is negative. Instability can only be driven by a simple, real (positive) eigenvalue.

Proof The linearization of our circuit ODEs define positive linear systems, where the Jacobians are similar to Metzler matrices. Therefore, these Jacobians always have a real dominant eigenvalue, i.e. \( \lambda_{\text{max}} \geq \Re(\lambda_i), \forall \lambda_i \in J \) [7].

The coefficients of the characteristic polynomials are all real and all positive with the exception of the constant terms, which can be positive or negative. If the constant term of each \( p_J(\lambda, \bar{x}) \) is negative, then we know that \( p_J(0, \bar{x}) < 0 \) and it is real. In the limit \( \lambda \to \infty \), \( p_J(\lambda, \bar{x}) > 0 \) because all other coefficients are positive. Thus, there must be at least one point in the right half plane that is a root of the \( p_J(\lambda, \bar{x}) \), all our systems are unstable, and because the Jacobians are similar to Metzler matrices, their largest roots must be real.

If a system is unstable, then its characteristic polynomial must have at least one root with positive real part. \textit{Ab absurdo}, suppose the constant term is positive. Then instability can only occur with a pair of complex conjugate eigenvalues with positive real part. This is impossible because the Jacobian is a Metzler matrix and the dominant eigenvalue must be real. Thus, the constant term of the characteristic polynomial must be negative.

We can now finish our stability analysis. Our systems all have degree +1 (Theorem 2), thus when three equilibria are present their indices must be equal to +1, +1, and -1 so that their sum is +1 (we recall that all equilibria of our systems are regular). Since the index is equal to the sign of the constant term in the characteristic polynomial, a positive index is associated with a stable equilibrium and a negative index is associated with an unstable equilibrium, and we can conclude that, with three equilibria, our systems are bistable. Note that the unstable point does not admit local oscillatory behaviors, because local instability is driven by a real eigenvalue (Lemma 1). As an alternative argument, we can also simply note that our systems are monotone—for any choice of parameters the Jacobians are similar to Metzler matrices,
a property that defines a monotone system with respect to the positive orthant $[8, 9]$—and a monotone system does not admit oscillatory behaviors.

References


