Inverse Perturbation for Optimal Intervention in Gene Regulatory Networks

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\section*{ABSTRACT}

Motivation: Analysis and intervention in the dynamics of gene regulatory networks is at the heart of emerging efforts in the development of modern treatment of numerous ailments including cancer. The ultimate goal is to develop methods to intervene in the function of living organisms in order to drive cells away from a malignant state into a benign form. A serious limitation of much of the previous work in cancer network analysis is the use of external control, which requires intervention at each time step, for an indefinite time interval. This is in sharp contrast to the proposed approach, which relies on the solution of an inverse perturbation problem to introduce a one-time intervention in the structure of regulatory networks. This isolated intervention transforms the steady-state distribution of the dynamic system to the desired steady-state distribution.

Results: We formulate the optimal intervention problem in gene regulatory networks as a minimal-perturbation of the network in order to force it to converge to a desired steady-state distribution of gene regulation. We cast optimal intervention in gene regulation as a convex optimization problem, thus providing a globally optimal intervention strategy. We consider a perturbation that minimizes (i) the overall energy of change between the original and controlled networks and (ii) the time needed to reach the desired steady-state of gene regulation. Furthermore, we show that there is an inherent tradeoff between minimizing the energy of the perturbation and the convergence rate to the desired distribution. We apply the proposed control to the Human melanoma gene regulatory network.

Availability: The MATLAB code for optimal intervention in gene regulatory networks can be found online: http://syen.ualr.edu/nxbouaynaya/Bioinformatics2010.html.

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\section*{1 INTRODUCTION}

The cell maintains its function via an elaborate network of interconnecting positive and negative feedback loops of genes and proteins that send different signals to a large number of pathways and molecules. Understanding the dynamic behavior of gene regulatory networks is essential to advance our knowledge of disease, develop modern therapeutic methods, and identify targets in the cell needed to reach a desired goal. In classical biological experiments, cell function is ascertained based on rough phenotypical and genetic behavior. On the other hand, the use of dynamical system models allows one to analytically explore biological hypotheses. Within this context, investigators have sought to discover preferable stationary states, the effect of distinct perturbations on gene dynamics, and the ‘dynamical function’ of genes (Shmulevich\textsuperscript{et al.}, 2002b, Abhishek\textsuperscript{et al.}, 2008, Fathallah-Shaykh, 2005, Ribeiro and Kauffman, 2007, Datta\textsuperscript{et al.}, 2007, Qian\textsuperscript{et al.}, 2009, Qian and Dougherty, 2009, Fathallah-Shaykh\textsuperscript{et al.}, 2009).

The complexity of biological systems and the noisy nature of the sampled data suggest the use of probabilistic methods for system modeling, analysis, and intervention. Markov chain models have been shown to accurately emulate the dynamics of gene regulatory networks (Kim\textsuperscript{et al.}, 2002). In particular, the dynamics of Probabilistic Boolean Networks (PBNs) (Shmulevich\textsuperscript{et al.}, 2002b) and Dynamic Bayesian Networks (Murphy, 2002) can be studied using Markov chains. The long-run behavior of a dynamic network is characterized by the steady-state distributions of the corresponding Markov chain. It has been argued that steady-state distributions determine the phenotype or the state of the cell development, such as cell proliferation and apoptosis (Kauffman, 1993, Ivanov and Dougherty, 2006). The long-run dynamic properties of PBNs and their sensitivity with respect to network perturbations were investigated in several manuscripts (Shmulevich\textsuperscript{et al.}, 2003, Qian and Dougherty, 2009, Qian and Dougherty, 2008).

The ultimate objective of gene regulatory network modeling and analysis is to use the network to design effective intervention strategies for affecting the network dynamics in such a way as to avoid undesirable cellular states. As futuristic gene therapeutic interventions, various control strategies have been proposed to alter gene regulatory network dynamics in a desirable way. Biologically, such alterations may be possible by the introduction of a factor or drug that alters the extant behavior of the cell. Current control strategies can be grouped into three main approaches (Datta\textsuperscript{et al.}, 2007): (i) reboot the network by resetting its initial condition (Shmulevich\textsuperscript{et al.}, 2002c), (ii) introduce external control variables to act upon some control genes, in such a way as to optimize a given cost function (Datta\textsuperscript{et al.}, 2003, Pal\textsuperscript{et al.}, 2006, Datta\textsuperscript{et al.}, 2007, Faryabi\textsuperscript{et al.}, 2008), (iii) alter the underlying rule-based structure of the network in order to shift the steady-state mass of the network from undesirable to desirable states. This last type of intervention is also referred to as structural intervention (Shmulevich\textsuperscript{et al.}, 2002a, Qian and Dougherty, 2008, Qian\textsuperscript{et al.}, 2009).

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The first strategy requires knowledge of the basin of attraction of the desirable steady-state distribution. For large networks, finding the basin of attraction of a given steady-state is a computationally expensive task (Kauffman, 1993, Wuenesch, 1998). The second strategy minimizes a given cost function by controlling the expression level of target genes in the network. In particular, this strategy assumes prior knowledge of the genes to be used as control agents, and the cost associated with each state of the network. More importantly, this strategy produces a recurrent control policy, over a possibly infinite time horizon interval (Datta et al., 2003, Pal et al., 2005b, Pal et al., 2006, Datta et al., 2007, Faryabi et al., 2008). Clinically, such an infinite-horizon intervention can be viewed as connecting the patient to an infinitely recurrent feedback control loop. If the control is applied over a finite time horizon and then stopped, the steady-state distribution of the network (and hence the cell fate) may not change.

The third strategy aims at altering the long-run behavior of the network or its steady-state distributions. A simulation-based study was first conducted in (Shmulevich et al., 2002a), where a procedure to alter the steady-state probability of certain states was implemented using genetic algorithms. Xiao et al. (Xiao and Dougherty, 2007) considered an analytical study, where they explored the impact of function perturbations on the network attractor structure. However, their algorithms are rather cumbersome as they need to closely investigate the state changes before and after perturbations. Moreover, their practical usefulness is limited to singleton attractors, and they do not provide a steady-state characterization for Boolean networks (Qian and Dougherty, 2008). An analytical characterization of the effect on the steady-state distribution caused by perturbation of the regulatory network appears in (Qian and Dougherty, 2008). They relied on the general perturbation theory for finite Markov chains (Kemeny and Snell, 1960) to compute the perturbed steady-state distribution in a sequential manner. They subsequently proposed an intervention strategy for PBNs that affects the long-run dynamics of the network by altering its rule structure. However, they considered rank-one perturbations only. The extension of their method to higher-rank perturbations is iterative and computationally very expensive. Finally, a performance comparison of the above strategies has been conducted in (Qian et al., 2009).

In summary, the first two approaches do not guarantee convergence towards the desired steady-state distribution. The third approach, referred to as structural intervention, aims to shift the steady-state mass from undesirable to desirable states. The proposed solutions thus far have been limited to either simulation-based studies (Shmulevich et al., 2002a) or special cases (e.g., rank-one perturbations) (Xiao and Dougherty, 2007, Qian and Dougherty, 2008). In this paper, we provide a general solution to the problem of shifting the steady-state mass of gene regulatory networks modeled as Markov chains. We formulate optimal intervention in gene regulation as a solution to an inverse perturbation problem and demonstrate that the solution is (i) unique, (ii) globally optimum, (iii) non-iterative, and (iv) can be solved efficiently using standard convex optimization methods. The analytical solution to this inverse problem will provide a minimally-perturbed Markov chain characterized by a unique steady-state distribution corresponding to a desired distribution. Such a strategy introduces an isolated, one-time intervention that will require a minimal change in the structure of the regulatory network and converges to a desired steady-state. Moreover, we cast optimal intervention as a convex optimization problem, thus providing a globally-optimal solution that can be efficiently computed using standard toolboxes for convex optimization (Boyd and Vandenberghe, 2003). In particular, we no longer need simulation-based or computationally-expensive algorithms to determine the optimal intervention. The criteria adopted for optimality is designed to minimize potential adverse effects caused by the intervention strategy. Specifically, we will focus on minimization of the change in the structure of the network and maximization of the convergence rate towards the steady-state distribution. We will therefore investigate the following criteria for minimal-perturbation control in the solution of the inverse perturbation problem:

- Reduce the level of change in the expression level of specific genes that are introduced by control agents; that is, we will minimize the overall energy of change between the original and perturbed transition matrices as characterized by the Euclidean-norm of the perturbation matrix.
- Increase the rate of convergence of the network to the desired steady-state distribution; thus, we will minimize the time needed to reach the desired steady-state distribution as evaluated by the second-largest eigenvalue modulus of the perturbed matrix.

This work differs from previous research in optimal structural intervention in at least three ways: First, we do not evaluate the effect of network perturbation on the steady-state distribution (Qian and Dougherty, 2009, Qian and Dougherty, 2008, Qian et al., 2009). Although the subject of perturbation of Markov chains is a well-studied field, unlike the previous works reported in the literature, we do not tackle the subject of perturbation of Markov chains; instead we propose a new framework for the solution of the inverse perturbation problem. That is, the perturbation problem aims to characterize the variation in the stationary distribution in response to a perturbation of the transition matrix (Schweitzer, 1968). The inverse perturbation problem, on the other hand, investigates the perturbation required in order to reach a desired stationary distribution. The proposed approach to the inverse perturbation problem therefore has the potential to have a wide impact in many applications that rely on dynamic systems. Second, unlike the previous work, which is limited to rank-one perturbations, we consider any perturbation that preserves the irreducibility of the original network (Qian and Dougherty, 2008). Third, whereas previous efforts considered unconstrained optimal intervention strategies, we focus on optimal control strategies, which incorporate (energy and rate of convergence) constraints on the protocols employed in gene regulation designed to reduce adverse effects as a result of the intervention strategy.

The mathematical notation used in the paper as well as the proofs of several new results are detailed in the supplementary material of this paper.

2 METHODS

We consider a gene regulatory network with \( m \) genes \( g_1, \ldots, g_m \), where the expression level of each gene is quantized to \( l \) values. The expression levels of all genes in the network defines the state vector...
of the network at each time step. Gene $g_i$ evolves according to a
time-invariant probabilistic law determined by the expression levels of
the genes in the network; i.e., $\Pr(g_i = x_i | g_1 = x_1, \ldots, g_m = x_m)$,
for $x_i \in \{0, 1, \ldots, l - 1\}$ and $j = 1, \ldots, m$. An approach
to obtain the conditional probabilities of the genes from gene
expression data has been presented in Kim et al. [2002], Shmulevich
et al. [2002] based on the coefficient of determination Dougherty
et al. [2000]. The dynamics of this network can be represented as a
finite-state homogeneous Markov chain described by a probability
transition matrix encapsulates the one-step conditional probabiliti es of the
$P$ network at each time step. Gene

$$
\Pr(g_i = x_i | g_1 = x_1, \ldots, g_m = x_m) = \sum_{x_i} \Pr(g_i = x_i, \ldots, g_m = x_m | g_1 = x_1, \ldots, g_m = x_m),
$$

where $\tilde{x}_i$ denotes the set of all $x_i$’s except $x_i$; i.e., $\tilde{x}_i = \{x_1, \ldots, x_{i-1}, x_{i+1}, \ldots, x_m\}$. Consequently, if the probability
transition matrix $P$ is perturbed linearly with a zero-row sum
matrix $E = (\epsilon_{ij})_{1 \leq i, j \leq n}$, then conditional probability of each gene
$\Pr(g_i = x_i | g_1 = x_1, \ldots, g_m = x_m)$ is perturbed linearly by $\sum_{j \in J} \epsilon_{kj}$,
where $h$ is the index of the state vector $[g_1, \ldots, g_m]$ and $J$ is an
isomorphic to $\{1, 2, \ldots, \frac{m}{2}\}$. Thus, we observe that “small”
perturbations $\epsilon_{ij} \ll 1$ of the probability transition matrix that
satisfy the zero-row sum condition $\sum_{j \in J} \epsilon_{kj} = 0$, lead to “small”
perturbations of the genes’ dynamics.

We assume that $P$ is ergodic, i.e., irreducible and aperiodic.
Therefore, the existence and uniqueness of the steady-state
distribution are guaranteed. In practice, there are several fast
algorithms for checking irreducibility and aperiodicity in graphs
(Sharir, 1981). If $P$ is ergodic, then the limiting matrix $P_{\infty} = \lim_{n \to \infty} P_n$ satisfies $P_{\infty} = \pi P_{\infty}$ (Seneta, 2006). In particular, the
rows of the limiting matrix $P_{\infty}$ are identical. This demonstrates
that, in the ergodic case, the initial state of the network has no
influence on the long-run behavior of the chain.

**Definition 1.** A row probability vector $\mu^t = (\mu_1, \ldots, \mu_n)$ is
called a stationary distribution or a steady-state distribution for $P$
if $\mu^t P_{\infty} = \mu^t$.

Because $P$ is stochastic (i.e., its rows sum up to 1), the existence
of stationary distributions is guaranteed (Kemeny and Snell, 1960).

Let $\pi_0$ denote the undesirable steady-state distribution of $P$. We
wish to alter this distribution by linearly perturbing the probability
transition matrix $P$. Specifically, we consider the perturbed
stochastic matrix

$$
P = P_0 + C,
$$

where $C$ is a zero row-sum perturbation matrix. The zero row-
sum condition is necessary to ensure that the perturbed matrix $P$
is stochastic. Let us denote by $\pi_d$ the desired stationary distribution.
We seek to design an optimal zero row-sum perturbation matrix $C$
such that the perturbed matrix $P$ is ergodic and converges to the
desired steady-state distribution $\pi_d$.

## 2.1 The Feasibility Problem

Schweitzer (Schweitzer, 1968) showed that the ergodic perturbed
matrix $P = P_0 + C$ possesses a unique stationary distribution $\pi_d$,
which satisfies

$$
\pi_d = \pi_0 (I - C Z_0)^{-1},
$$

where $Z_0$ is the fundamental matrix of $P_0$ given by $Z_0 = (I - P_0 + P_0^{-\infty})^{-1}$. Equation (3) requires the computation of $\pi_0$, the initial
undesired steady-state distribution, and the fundamental matrix $Z_0$,
which involves the computation of the inverse of an $n \times n$ matrix.
The following proposition shows that Eq. (3) is equivalent to a
simpler and computationally more efficient condition.

**Proposition 1.** Consider a stochastic $n \times n$ ergodic matrix $P_0$
with steady-state distribution $\pi_0$ and fundamental matrix $Z_0$. If $C$
is any $n \times n$ matrix, and $\pi_d$ any probability distribution vector, then
we have

$$
\pi_d = \pi_0 (I - C Z_0)^{-1} \iff \pi_d (P_0 + C) = \pi_d.
$$

In the gene regulatory control problem, we are interested in the
inverse perturbation problem. Namely, given the desired stationary
distribution, $\pi_d$, we wish to determine a perturbation matrix $C$
that satisfies Eq. (4). Notice that there may be multiple solutions
to Eq. (4); i.e., different perturbation matrices $C$ could lead to the
same desired stationary distribution. The problem of finding the set of perturbation matrices satisfying Eq. (4) can be formulated as the
following feasibility problem.

**The feasible set of the control problem:** Given an ergodic network
classified by its probability transition matrix $P_0$ with stationary
distribution $\pi_0$, and given a desired stationary distribution $\pi_d$, then
the set of perturbation matrices $C$, which force the network to
transition from $\pi_0$ to $\pi_d$ satisfy the following constraints:

(i) $\pi_d^t = \pi_d^t (P_0 + C)$, (ii) $C1 = 0$, (iii) $P_0 + C \geq 0$.

Constraints (ii) and (iii) ensure that the perturbed matrix $P$ is a
proper probability transition matrix: constraint (ii) imposes that the
perturbation matrix $C$ is zero-row sum, and hence the perturbed
matrix $P$ is stochastic, and constraint (iii) requires the matrix $P$
to be element-wise non-negative. Let $D$ denote the feasible set of
perturbation matrices, i.e.,

$$
D = \{ C \in \mathbb{R}^{n \times n} : \pi_d^t = \pi_d^t (P_0 + C), C1 = 0, P_0 + C \geq 0 \}.
$$

$D$ is a polyhedra as the solution of a finite number of linear
equalities and inequalities (Boyd and Vandenberghe, 2003). It
is easily shown that polyhedra are convex sets (Boyd and
Vandenberghe, 2003). Observe that $D$ is non-empty because it
contains the perturbation matrix $C = 1 \pi_d - P_0$.

Observe that there are numerous (possibly infinite) perturbation
matrices $C$ which can force the network to transition from an
undesirable steady-state to a desirable one. All such perturbations,
in principle, constitute plausible control strategies and can therefore
be used to drive the network from one steady-state to another.
We impose the minimum-energy and fastest convergence rate
constraints in order to limit the structural changes in the network
and reduce the transient dynamics after perturbation.
2.2 The Minimal-Intervention Problem

Because the feasible set, defined in Eq. (5) is non-empty, there exists at least one perturbation matrix $C$, which forces the network to converge to the desired distribution. A natural question arises then: “Which perturbation matrix should we choose?” In practice, we are interested in perturbation matrices, which incorporate specific biological constraints; e.g., the potential side effects on the patient, and the length of treatment. We translate these limitations into the following optimality criteria.

2.2.1 Minimal-perturbation energy control The minimal perturbation energy control is defined by minimization of the Euclidean-norm of the perturbation matrix. It corresponds, biologically, to the control which minimizes the overall “energy” of change between the perturbed and unperturbed gene regulatory networks. The minimal perturbation energy control is defined by minimization of the Euclidean-norm of $C$ is defined as

$$
\|C\|_2 = \max \{ \|C x\| : x \in \mathbb{R}^n, \|x\| = 1 \}
\tag{7}
$$

$$
= \sqrt{\lambda_{\text{max}}(C^t C)} = \max_x \langle C^t C x, x \rangle, \quad \tag{8}
$$

where $\lambda_{\text{max}}(C^t C) \geq 0$ is the highest eigenvalue of the positive-semi-definite matrix $C^t C$, and $\langle \cdot , \cdot \rangle$ denotes the inner product.

The minimum perturbation energy control can be formulated as the following optimization problem:

**Minimal-perturbation energy control :**

$$
\text{Minimize } \|C\|_2 \text{ subject to } C \in \mathcal{D}, \quad \tag{9}
$$

where $\mathcal{D}$ is the feasible set in Eq. (6).

The optimization problem formulated in Eq. (9) is a convex optimization problem. A convex optimization problem is defined as one that satisfies the following three requirements: (a) the objective function is convex; (b) the inequality constraint functions are convex; and (c) the equality constraint functions are affine (Boyd and Vandenberghe, 2003). A fundamental property of convex optimization problems is that any locally optimal point is also globally optimal. Moreover, because the Euclidean-norm is strictly convex, the optimal solution is unique.

Next, we express the convex optimization problem as a semi-definite programming (SDP) problem, which can be solved efficiently using standard SDP solvers, such as SDPSOL (Wu and Boyd, 1996), SDPpack (Alizadeh et al., 1997) and SeDuMi (Sturm, 1999). A list of 16 SDP solvers can be found at the SDP website maintained by Helmerk (Helmer, 2003). We can thus rely on SDP solvers to efficiently compute the optimal perturbation of Boolean gene networks consisting of 10 to 15 genes (i.e. $2^{10} = 1024$ to $2^{15} = 32768$ states). Note, however, that the computational efficiency of SDP solvers for larger networks will be lower.

**Semi-definite programming formulation :** Using the fact that

$$
\|C\|_2 \leq t \iff C^t C \preceq t^2 I, \quad t \geq 0,
$$

we can express the problem in Eq. (9) in the following form

$$
\text{Minimize } t \text{ subject to } C^t C \preceq t^2 I, \quad P_0 + C \geq 0 \tag{10}
$$

$$
\pi^t_d(P + C) = \pi^t_d, \quad C I = 0
$$

with variables $t \in \mathbb{R}$ and $C \in \mathbb{R}^{n \times n}$. The problem (10) is readily transformed to a SDP standard form, in which a linear function is minimized, subject to a linear matrix inequality and linear equality constraints. We first observe that, from the Schur complement, we have

$$
C^t C \preceq t^2 I \text{ (and } t \geq 0) \iff \begin{pmatrix} tI & C \\ C^t & tI \end{pmatrix} \succeq 0. \quad \tag{11}
$$

The inequalities in (10) can be expressed as a single linear matrix inequality by using the fact that a block diagonal matrix is positive-semi-definite if and only if its blocks are positive semi-definite.

$$
\text{Minimize } t \text{ subject to } \begin{pmatrix} tI & C \\ C^t & tI \end{pmatrix} \succeq 0 \quad \begin{pmatrix} 0 \ & 0 \\ 0 & \text{vec}(P_0 + C) \end{pmatrix} = 0 \quad \pi^t_d(P + C) = \pi^t_d, \quad C I = 0
$$

At this stage, it is important to notice that we can similarly consider the $L_1$ norm to produce a sparse perturbation matrix (Boyd and Vandenberghe, 2003).

2.2.2 Fastest convergence rate control A clinically-viable optimality criterion is to select the perturbation that yields the fastest convergence rate to the desired steady-state distribution. We know that the convergence rate of ergodic Markov chains is geometric with parameter given by the second-largest eigenvalue modulus (SLEM) of the probability transition matrix (Seneta, 2006). The smaller the SLEM, the faster the Markov chain converges to its equilibrium distribution. The fastest convergence rate control can be casted as the following optimization problem:

**Fastest convergence rate control :**

$$
\text{Minimize } \text{SLEM }(P_0 + C) \text{ subject to } C \in \mathcal{D}, \quad \tag{13}
$$

where $\mathcal{D}$ is the feasible set in Eq. (6). Observe that for a general (non-symmetric) matrix, about the only characterization of the eigenvalues is the fact that they are the roots of the characteristic polynomial. Therefore, the objective function in (13) is not necessarily convex, and thus the optimization problem is not convex.

The following obvious proposition determines the optimal fastest convergence rate perturbation matrix.

**Proposition 2.** The optimal solution of the optimization problem in (13) is given by

$$
C^* = 1 \pi^t_d - P_0. \quad \tag{14}
$$

The optimal SLEM $(P_0 + C^*) = 0$.

That is the perturbation $C^*$ reaches the desired state in a single jump.

The fastest convergent perturbation may, however, result in a large energy deviation between the original and perturbed networks. Next, we will investigate the tradeoffs between minimal-energy and fastest convergence criteria.
2.2.3 Tradeoffs between minimal-energy and fastest convergence rate control

Fig. 1. The Human melanoma gene regulatory network: (a) Plot of SLEM\( (P(s)) \) in Eq. (17) (blue continuous line) and SLEM\( (Q(s)) \) in Eq. (22) (red dashed line) versus \( s \); (b) Plot of \( ||C(s)|| \) in Eq. (16) (blue continuous line) and \( ||C_Q(s)|| \) in Eq. (21) (red dashed line) versus \( s \); (c) Plot of \( ||\pi_d(s) - \pi_d|| \) (blue continuous line), and the upper bound \( \frac{1}{2} ||\pi_0 - \pi_d|| \) (red dashed line) versus \( 0 \leq s \leq 1 \), in Proposition 5. The tradeoff between minimal-energy and fastest convergence rate control is clear from (a) and (b). The parameterized family of perturbed matrices \( P(s) \) in Eq. (15) results in a faster convergence towards the desired steady-state \( \pi_d \) at the expense of a higher norm (energy) of the perturbation matrix. On the other hand, the family of perturbed matrices \( Q(s) \) in Eq. (20) leads to a perturbation matrix with norm (energy) as small as desired, but at the expense of not converging towards the desired steady-state for small \( s \). The distance between the steady-state of \( Q(s) \) and the desired steady-state as a function of \( s \) is shown in Figure (c).

Proposition 4, the norm of the perturbation matrix, and hence the energy deviation between the original and perturbed networks, increases as a function of \( s \). Therefore, we have an inherent tradeoff between the energy of the perturbation matrix and the rate of convergence. The faster we converge towards the desired steady-state, the higher the energy between the initial and perturbed networks.

We would, therefore, like to find the optimal tradeoff perturbation matrix. Specifically, we determine the optimal perturbation matrix, which minimizes the SLEM while keeping the energy bounded. Such a constraint can be imposed, for instance, to minimize the side effects due to the rewiring of the original network. The optimal tradeoff problem is readily written as the following optimization problem:

\[
\text{Minimize } \text{SLEM} (P_0 + C)
\]

subject to \( ||C||_2 \leq \epsilon, \quad C \in D \),

where \( \epsilon \geq ||C(s)|| \) is a given threshold. We consider the solution to the optimization problem in (18) along the line defined in Eq. (15).

A local minimum of the optimization problem in (18) might not belong to the family \( \{P(s)\}_{s \in [0,1]} \). However, the line search seems a reasonable choice, and presents several advantages: (i) it provides a closed-form expression of the SLEM of \( P(s) \) for all \( 0 \leq s \leq 1 \); (ii) Contrary to most eigenvalue problems, which are numerically unstable, the line search has an explicit formula, and hence is numerically stable; (iii) it describes a linear behavior of the optimal solution.

It is straightforward to see that the optimal tradeoff perturbation matrix on the line, defined by Eq. (15), is given by \( C^* \equiv C(s^*) \), where \( s^* \) is the unique solution to \( ||C(s)||_2 = \epsilon \). However, the optimal tradeoff perturbation matrix requires a numerical computation of the minimal energy perturbed matrix \( ||P_0||_2 \). More importantly, if the bound on the energy \( \epsilon < ||C(s)||_2 \), then we have no solution for the problem (18). Indeed, in some cases, we might want to constrain the energy of the perturbation matrix \( C \) to be no larger than a “small” specified threshold (i.e., \( \epsilon < ||C||_2 \)). We

\[
P(s) = (1 - s)P_E^* + s1\pi_d^*.
\]
will show that, in this case, we might not be able to reach the desired steady-state distribution. Intuitively, if the energy of the perturbation matrix is constrained to be too small, then we might not be able to force the network to transition from one steady-state to another. In this case, we will quantify how far we are from the desired steady-state.

Mathematically, the general energy constrained optimization problem can be formulated as follows

Energy-constrained fastest convergence rate control:

\[
\text{Minimize } \text{SLEM} \left( P_0 + C \right) \quad (19)
\]

subject to \( ||C||_2 \leq \epsilon, \ C \mathbf{1} = \mathbf{0}, \ (P_0 + C) \geq 0, \)

where \( \epsilon \geq 0. \) Observe that the optimization problem in (19) is different from the problem in (18) in two points: First, the bound \( \epsilon \) can be any non-negative number. Second, the perturbation matrix \( C \) does not necessarily belong to \( \mathbb{D}. \) Observe that the optimization problem in (19) is not a convex optimization problem as the SLEM of a general (non-symmetric) matrix is not necessarily convex. We will look for a solution on the line between \( P_0 \) and \( \pi_d, \) i.e., we consider the family

\[
Q(s) = (1 - s)P_0 + s\pi_d, \quad 0 \leq s \leq 1. \quad (20)
\]

The perturbation matrix, \( C_Q, \) is therefore given by

\[
C_Q(s) = Q(s) - P_0 = s(\pi_d - P_0). \quad (21)
\]

In particular, the norm \( ||C_Q(s)|| \) can be made arbitrarily small by choosing a small \( s. \) On the other hand, it is easy to see that

\[
\text{SLEM} \left( Q(s) \right) = (1 - s) \text{SLEM} \left( P_0 \right). \quad (22)
\]

The proof of Eq. (22) follows the same steps of the proof of Proposition 3. Therefore, it seems that the family \( \{Q(s)\}_{0 \leq s \leq 1} \) provides a perturbation matrix with an arbitrarily small energy, and an explicit formula for the SLEM of the perturbed network as a function of the SLEM of the original network. The drawback, however, is that \( Q(s) \) does not necessarily converge to the desired steady-state distribution. The following proposition quantifies the difference between the steady-state of \( Q(s) \) and the desired steady-state \( \pi_d. \)

\textbf{Proposition 5.} The family of matrices \( Q(s), \) given in Eq. (20), is ergodic for all \( 0 \leq s \leq 1, \) and therefore converges towards a unique steady-state distribution \( \pi_d(s) \) given by

\[
\pi_d(s) = s(1 - s)(I - (1 - s)P_0)\text{P}_0^{-1}P_0\pi_0 + (1 - s)\pi_0 + s\pi_d. \quad (23)
\]

That is

\[
\pi_d(s) - \pi_d = (1 - s)(I - s(I - s)P_0)\text{P}_0^{-1}P_0\pi_0 - \pi_0 + s\pi_d. \quad (24)
\]

Furthermore, we have

\[
||\pi_d(s) - \pi_d|| \leq A(P_0)(1 - s)||\pi_0 - \pi_d||, \quad 0 \leq s \leq 1, \quad (25)
\]

where \( A(P_0) = 1 + \sup_{k \geq 1} ||P_0^k||_2, \) which is finite because \( P_0 \) has a limit as \( k \to \infty. \) If \( P_0 \) is symmetric, then we have a simpler upper bound given by

\[
||\pi_d(s) - \pi_d|| \leq \frac{2(1 - s)}{2 - s}||\pi_0 - \pi_d||, \quad 0 \leq s \leq 1. \quad (26)
\]

From Proposition 5, it is clear that when \( s \to 1, \pi_d(s) \to \pi_d. \)

### 3 Optimal Intervention in the Human Melanoma Gene Regulatory Network

The inverse perturbation control is applicable in every gene regulatory network that can be modeled as a Markov chain. In particular, we note that two of the most popular gene regulatory network models, Probabilistic Boolean Networks (PBNs) and Dynamic Bayesian Networks (DBNs) can be modeled as Markov chains (Llodesmikia et al., 2006). In this paper, we consider the Human melanoma gene regulatory network, which is one of the most studied gene regulatory networks in the literature (Pal et al., 2006, Datta et al., 2007, Qian and Dougherty, 2008). The abundance of mRNA for the gene WNT5A was found to be highly discriminating between cells with properties typically associated with high versus low metastatic competence. Furthermore, it was found that an intervention that blocked the Wnt5a protein from activating its receptor, the use of an antibody that binds the Wnt5a protein, could substantially reduce Wnt5A's ability to induce a metastatic phenotype (Pal et al., 2006). This suggests a control strategy that reduces WNT5A’s action in affecting biological regulation.

A seven-gene probabilistic Boolean network model of the melanoma network containing the genes WNT5A, pirin, S100P, RET1, MART1, HADHB, and STC2 was derived in (Pal et al., 2005a). Figure 2(a) illustrates the relationship between genes in the Human melanoma regulatory network. This diagram is a conceptual abstraction and is not intended as an explicit mechanistic diagram of regulatory actions. The influences depicted may be the result of many intervening steps that are not shown. Some generalizations that emerge from this conceptual diagram, such as the wide influence of the state of WNT5A on the states of other genes, have been confirmed experimentally (Kim et al., 2002). Note that the Human melanoma Boolean network consists of \( 2^7 = 128 \) states ranging from \( 00 \cdots 0 \) to \( 11 \cdots 1, \) where the states are ordered as WNT5A, pirin, S100P, RET1, MART1, HADHB, and STC2, with WNT5A and STC2 denoted by the most significant bit (MSB) and least significant bit (LSB), respectively. The probability transition matrix of the Human melanoma network, derived in (Zhou et al., 2004) and used in this paper, is courtesy of Dr. Ranadip Pal.

A naive control strategy, which would exclusively target the gene WNT5A by reducing its expression level while keeping the expression levels of the other genes in the network unchanged will inevitably fail as it basically resets the initial state of the underlying process and does not alter the network structure. Biologically, the complex gene interactions in the network will almost certainly bypass this gene perturbation and return to their initial cancerous state. On the other hand, determining the optimal gene intervention by a brute-force approach is computationally intractable and experimentally infeasible: Even within the context of Boolean regulation (two-level quantization), the number of experiments to perform increases exponentially in the number of genes in the network. For instance, in the 7-gene melanoma network, an extensive control search amounts to performing 2186 laboratory experiments; i.e. downregulate and upregulate the expression level of every gene, every pair of genes, every triple of genes, etc., thus requiring \( \sum_{k=1}^7 \binom{7}{k} 2^k = 2186 \) laboratory experiments. The proposed inverse perturbation control provides the optimal one-time intervention that rewires the network in order to force it to converge to the desired steady-state.
Using the breadth first search algorithm (Russell and Norvig, 2003), we found that the melanoma probabilistic Boolean network is irreducible. Therefore, it has a unique stationary distribution, and we can apply the inverse perturbation control developed in this paper. Because the control objective is to downregulate the WNT5A gene, we consider the desired steady-state distribution where the probability of the states having WNT5A upregulated is $10^{-4}$ and the probability of the other states, which correspond to WNT5A downregulated is set equal to 0.015525 (see Fig. 2(b)). Observe that the states from 0 to 63 have WNT5A downregulated (0) and hence are desirable states, as compared to states 64 to 127 that have WNT5A upregulated (1) and hence are undesirable. The probability transition matrices of the Human melanoma networks corresponding to the original and perturbed networks are portrayed in Fig. 3. Observe that the family of perturbed matrices $\{P(s)\}_{s \in [0,1]}$, defined in Eq. (15), converges towards the desired steady-state distribution $\pi_d$, in the sense that $P(s)^n \rightarrow 1\pi_d^n$ as $n \rightarrow \infty$. On the other hand, the family of matrices $Q(s)$, defined in Eq. (20), does not converge to the desired distribution $\pi_d$, for $0 \leq s < 1$. Figure 1(c) shows the norm difference between the steady-state distribution of $Q(s)$, $\pi_d(s)$, and $\pi_d$ as a function of $s$. As the parameter $s$ increases, $\pi_d(s) \rightarrow \pi_d$. The advantage of considering the family $\{Q(s)\}$ resides in the ability to design perturbation matrices $C(s)$ with arbitrary small norms (energy) (see Fig. 1(b)). This is in contrast to the family $\{P(s)\}$ where the norm of the perturbation matrices is lower bounded by the minimal-energy perturbation matrix norm $\|C_E\|$. The tradeoff between the minimal-energy and fastest convergence rate is depicted in Figs 1(a) and 1(b). The steady-state distribution of the Human melanoma network of the original and perturbed networks are shown in Fig. 2. Observe that the after-control steady-state is identical to the desired steady-state. Therefore, the control has enabled us to shift the steady-state probability mass from the undesirable states to states with lower metastatic competence.

The minimal-energy perturbed matrix, which optimally solves the SDP problem in (12), is $\|C_E\|_2 = 1.20667$ and its SLEM $= 0.4696$. We have shown that the optimal SLEM of the fastest convergence rate control is equal to 0, and its energy is given by $\|C\|_2 = \|1\pi_d^0 - P_0\|_2 = 1.81854$. The SDP problem has been implemented in MATLAB and uses the CVX software for convex optimization (Grant and Boyd, 2010).

The mathematical findings derived were confirmed by computer simulation experiments by demonstrating that the optimal perturbation of a known melanoma gene regulatory network leads to a desired steady-state. In order to reach the full impact of the proposed research on gene regulation in biological systems, we plan to investigate changes in the cell that induce the optimal perturbed transition matrix. In particular, our current and future work will focus on determining the optimal biological design rules needed to induce the optimal intervention strategy while limiting ourselves to biologically-viable design rules that rely on modern methods for molecular control in cellular systems.

4 CONCLUSION

In this paper, we introduced a novel method for optimal intervention in gene regulatory networks posed as an inverse perturbation problem. The optimal perturbation has been derived such that the regulatory network will transition to a desired stationary, or steady-state, distribution. Biological evidence suggests that steady-state distributions of molecular networks reflect the phenotype of the cell. In other words, both malignant (e.g. cancer) and benign phenotypes correspond to steady-state distributions in dynamic system models of gene regulatory networks.

We developed a mathematical framework for the solution of the inverse perturbation problem for irreducible and ergodic Markov chains. Our aim was to derive a minimal-perturbation intervention policy designed to introduce an isolated, one-time intervention
and induce few changes in the original network structure, thus minimizing potential adverse effects on the patient as a consequence of the intervention strategy. The mathematical analysis presented provides a general framework for the solution of the inverse perturbation problem for arbitrary discrete-event ergodic systems.

**ACKNOWLEDGEMENT**

The authors would like to extend their gratitude to Dr. Ranadip Pal from Texas Tech University for providing the Human melanoma gene regulatory network dataset. The authors are also grateful to Dr. S. Friedland from the University of Illinois at Chicago for illuminating discussions about this work.

This project is supported by Award Number R01GM096191 from the National Institute Of General Medical Sciences (NIH/NIGMS). The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institute Of General Medical Sciences or the National Institutes of Health.

**REFERENCES**


SUPPLEMENTAL FILE

5 MATHEMATICAL NOTATION

In this paper, we consider real variables. We use \( \mathbb{R} \) to denote the set of real numbers. Scalars are denoted by lower case letters, e.g., \( s, t \). Vectors in \( \mathbb{R}^n \) are denoted by either bold letters and numbers, or lower-case Greek letters, e.g., \( \mathbf{x}, \pi, \). \( \mathbf{1} \) denotes a vector all of whose components are equal to one. \( \mathbf{x}^\top \) denotes the transpose of the vector \( \mathbf{x} \). The notation \( x_i \) refers to the \( i \)th component of the vector \( \mathbf{x} \). Matrices in \( \mathbb{R}^{m \times n} \) are denoted by either capital letters or upper-case Greek letters, e.g., \( C, P, A, I \) stands for the identity matrix. If \( P \in \mathbb{R}^{m \times n} \), then \( \text{vec}(P) \) transforms \( P \) into an \( mn \)-dimensional vector by stacking the columns. The equality and inequality symbols, \( =, \leq \) and \( \geq \) denote component-wise equality and inequality, respectively, for arrays of the same size. For example, if \( C \) is an \( m \times n \) matrix, then \( C \geq 0 \) denotes the \( mn \) inequalities: each element of the matrix \( C \) is nonnegative. The curved inequality symbols, \( \leq, \prec, \succ \), denote generalized matrix inequalities associated with the positive semi-definite cone. That is, if \( A, B \in \mathbb{R}^{n \times n} \), then \( A \geq B \) (resp., \( A \succ B \)) means that \( A - B \) is positive-semi-definite (resp., positive definite); and \( A \preceq B \) (resp., \( A \prec B \)) means that \( A - B \) is negative-semi-definite (resp., negative definite). We recall that a matrix \( A \in \mathbb{R}^{n \times n} \) is called positive-semi-definite (resp., positive definite) if \( x^\top Ax \geq 0 \) (resp., \( x^\top Ax > 0 \)) for all \( x \in \mathbb{R}^n \) (resp., \( x \neq 0 \)). If \( -A \) is positive-definite (resp., positive definite), then \( A \) is called negative-semi-definite (resp., negative definite).

PROOF OF PROPOSITION 1.

\[
\pi_d^i = \pi_0^i (I - C Z_0)^{-1} \\
\iff \pi_d^i (I - C Z_0) - \pi_0^i = 0 \\
\iff \pi_d^i (I - C Z_0) = \pi_0^i [I_Z^{-1}] = 0 \\
\iff \pi_d^i (I - P_0 + P_0^c) = \pi_d^i C - \pi_0^i (I - P_0 + P_0^c) = 0 \\
\iff \pi_d^i = \pi_d^i (P_0 + C),
\]

where Eq. (27) follows from the fact that \( Z_0 \) is invertible, and Eq. (28) follows from the properties: \( \pi_d^i P_0^c = \pi_d^i \mathbf{1} \pi_0^i = \pi_0^i; \pi_0^i P_0 = \pi_0^i; \) and \( \pi_0^i P_0^c = \pi_0^i \).

PROOF OF PROPOSITION 2. It is straightforward to check that the perturbation matrix \( C = (1 \pi_d^i - P_0) \in D \). That is we have (i) \( \pi_d^i = \pi_d^i (P_0 + C); \) (ii) \( C \mathbf{1} = 0; \) (iii) \( P_0 + C \geq 0 \). Moreover, the perturbed matrix \( P_0 + C = 1 \pi_d^i \) is a stochastic matrix with rank one. Therefore, it has a simple eigenvalue 1 corresponding to eigenvector \( \mathbf{1} \), and eigenvalue 0 with multiplicity \( n - 1 \). Hence, its SLEM = 0.

PROOF OF PROPOSITION 3. For any vector \( f \), we introduce its unique direct sum decomposition \( f = \alpha_f \mathbf{1} + f^\perp \), where \( \alpha_f = \pi_d^i f \) and \( f^\perp = \pi_0^i f \). It is easy to check that \( f^\perp \) is proportional to \( \mathbf{1} \) if and only if \( f^\perp = 0 \).

Let \( \psi = \alpha_\phi \mathbf{1} + \psi^\perp \) be a non-trivial eigenvector (i.e., \( \psi^\perp \neq 0 \)) of \( P_E^* \) with eigenvalue \( \mu \). We will look for a vector \( \phi \), in the form \( \phi = \psi + c \mathbf{1} \), that satisfies \( P(s) \phi = (1 - s) \mu \phi \). We have

\[
P(s) \phi = P(s) \psi + c \mathbf{1} \\
= (1 - s) \mu \psi + s \alpha_\phi \mathbf{1} + c \mathbf{1} \\
= (1 - s) \mu \psi + (s \alpha_\phi + c - (1 - s) \mu) \mathbf{1},
\]

where Eq. (29) follows from the fact that \( P(s) \) is stochastic, i.e., \( P(s) \mathbf{1} = 1 \), and Eq. (30) is obtained by replacing \( P(s) \) by its expression in Eq. (15). Therefore, if we chose \( c = \frac{s \alpha_\phi}{1 - s} \), we obtain \( P(s) \phi = (1 - s) \mu \phi \).

Let now \( \phi = \alpha_\phi \mathbf{1} + \phi^\perp \) be a non trivial eigenvector of \( P(s) \) with eigenvalue \( \lambda \). In particular, \( \phi^\perp \neq 0 \). We first show that \( \lambda \neq (1 - s) \). From Eq. (15), we have

\[
P(s) \phi = (1 - s) P_E^* \phi + s \alpha_\phi \mathbf{1} \\
= (1 - s) P_E^* \phi^\perp + (1 - s) \alpha_\phi \mathbf{1} + s \alpha_\phi \mathbf{1}.
\]

On the other hand, if \( \lambda = 1 - s \), then we would have

\[
P(s) \phi = (1 - s) \phi \\
= (1 - s) \alpha_\phi \mathbf{1} + (1 - s) \phi^\perp.
\]

By equating Eqs. (33) and (35), we obtain

\[
P_E^* \phi^\perp = \phi^\perp - \frac{s \alpha_\phi}{1 - s} \mathbf{1}.
\]

If \( \alpha_\phi = 0 \), which implies, from Eq. (36), that \( P_E^* \phi^\perp = \phi^\perp \). Hence, \( \phi^\perp \) is an eigenvector of \( P_E^* \) corresponding to eigenvalue 1. Therefore, \( \phi^\perp \) must be proportional to \( \mathbf{1} \). We recall that \( \phi^\perp \) is proportional to \( \mathbf{1} \) if and only if \( \phi^\perp = 0 \). This results in a contradiction because of the fact that \( \phi^\perp \neq 0 \). Therefore, we conclude that \( \lambda \neq 1 - s \).

Now, we consider \( \lambda \neq 1 - s \), we will find \( \psi \) in the form \( \psi = \phi + c \mathbf{1} \), that satisfies \( P(s) \psi = \frac{\lambda}{1 - s} \psi \). We have

\[
P(s) \psi = P(s) \phi + c \mathbf{1} \\
= \frac{\lambda}{1 - s} \psi - \frac{s \alpha_\phi}{1 - s} \mathbf{1} + c \mathbf{1} \\
= \frac{\lambda}{1 - s} \psi + (- \frac{\lambda}{1 - s} c - \frac{s \alpha_\phi}{1 - s} + c) \mathbf{1},
\]

where Eq. (39) follows from the stochasticity of \( P_E^* \) and Eq. (40) is obtained by replacing \( P_E^* \) by its expression in Eq. (15) and using the fact that \( \phi \) is an eigenvector of \( P(s) \) with eigenvalue \( \lambda \). Finally, Eq. (41) follows by replacing \( \phi = \psi - c \mathbf{1} \). Therefore, if we chose \( c = \frac{s \alpha_\phi}{1 - s} \), we obtain \( P(s) \psi = \frac{\lambda}{1 - s} \psi \).
We also provide an alternative proof as follows: Let \( A = P_E - P_0 \) and \( B = \mathbf{1}_d \sigma_d^2 - P_E \). Then, from Eq. (16),

\[
C(s) = A + Bs.
\]  
(42)

By construction, we have for all \( s \geq 0 \),

\[
\|C(s)\|_2 \geq \|P_E^* - P_0\|_2 = \|C(0)\|_2 = \|A\|_2 \iff \max_{x, \|x\|_1 = 1} < (A + sB)\hat{\phi}(A + sB)x, x > \geq \max_{x, \|x\|_1 = 1} < A^tAx, x > ,
\]  
(43)

where the right hand side equivalence follows from the definition of the spectral norm given in Eq. (8). Let \( x_\ast \) be such that \( \|x_\ast\| = 1 \) and

\[
\max_{x, \|x\|_1 = 1} < (A + sB)\hat{\phi}(A + sB)x, x > \geq (A + sB)^t(A + sB)x_\ast, x_\ast > .
\]  
(44)

Then,

\[
< (A + sB)^t(A + sB)x_\ast, x_\ast > \geq \langle A^tAx_\ast, x_\ast > ,
\]  
(45)

which means

\[
< (A^tB + A^tsB^tB)x_\ast, x_\ast > \geq 0.
\]  
(46)

Let \( \tilde{s} \geq s \). We need to show that

\[
\max_{x, \|x\|_1 = 1} < (A + \tilde{s}B)^t(A + \tilde{s}B)x, x > \geq < (A + sB)^t(A + sB)x_\ast, x_\ast > .
\]  
(47)

It is sufficient to show that

\[
< (A + \tilde{s}B)^t(A + \tilde{s}B)x_\ast, x_\ast > \geq < (A + sB)^t(A + sB)x_\ast, x_\ast > .
\]  
(48)

But,

\[
< (A + \tilde{s}B)^t(A + \tilde{s}B)x_\ast, x_\ast > - < (A + sB)^t(A + sB)x_\ast, x_\ast > = (\tilde{s} - s) < (A^tB + A^tsB^tB)x_\ast, x_\ast > ,
\]

which is positive because of Eq. (46).

**Proof of Proposition 5.** For \( 0 \leq s \leq 1 \), we have the following three properties

\[
\pi_d(s)^tQ(s) = \pi_d(s)^t
\]  
(49)

\[
\pi_d(s)1 = 1
\]  
(50)

\[
\pi_d(s) \geq 0.
\]  
(51)

Because \( Q(s) \) is ergodic, we know that such \( \pi_d(s) \) exists and is unique. Let

\[
\phi = \pi_d(s) - ((1-s)\pi_0 + s\pi_d).
\]  
(52)

Then, we have

\[
\phi^tQ(s) = \pi_d(s)^t - (1-s)^2\pi_0^t - s(1-s)\pi_d^tP_0 - s(1-s)\pi_0^t - s^2\pi_d^t
\]  
(53)

\[
= \phi^t + s(1-s)(\pi_0^t - \pi_d^tP_0)
\]  
(54)

\[
= \phi^t + s(1-s)(\pi_0^t - \pi_d^t)P_0,
\]  
(55)

where Eq. (55) follows from the fact that \( \pi_0^tP_0 = \pi_0^t \). Next, we notice that \( \phi^t1 = 0 \). Thus, from Eq. (20), we obtain

\[
\phi^tQ(s) = (1-s)\phi^tP_0.
\]  
(56)

By equating Eqs. (55) and (56), we obtain

\[
\phi^t[I - (1-s)P_0] = s(1-s)(\pi_0^t - \pi_d^t)P_0.
\]  
(57)

Observe that for \( s > 0 \), 1 is not an eigenvalue of \( (1-s)P_0 \). Hence, \( I - (1-s)P_0 \) is invertible, and we have

\[
\phi = s(1-s)[I - (1-s)P_0]^{-1}\pi_0^t(\pi_0 - \pi_d).
\]  
(58)

From Eq. (52), we have

\[
\pi_d(s) - \pi_d = \phi + (1-s)(\pi_0 - \pi_d).
\]  
(59)

Replacing \( \phi \) by its expression in Eq. (58), Eq. (59) can be written as

\[
\pi_d(s) - \pi_d = (1-s)\left( I - s(I - (1-s)P_0)\right)^{-1}\pi_0^t(\pi_0 - \pi_d).
\]  
(60)

That is, by factoring by \( (I - (1-s)P_0)^{-1} \),

\[
\pi_d(s) - \pi_d = (1-s)(I - (1-s)P_0)^{-1}(I - P_0^t)(\pi_0 - \pi_d).
\]  
(61)

If \( P_0 \) is symmetric, then by the spectral theorem we have \( \|P_0\|_2 = \lambda_{\max}(P_0) = 1 \), and by the triangle inequality,

\[
\|(I - (1-s)P_0)^{-1}(I - P_0^t)\|_2 \leq \frac{2}{2 - s},
\]

and thus

\[
\|\pi_d(s) - \pi_d\| \leq \frac{2(1-s)}{2 - s}\|\pi_0 - \pi_d\|.
\]

In the case of a non-symmetric matrix \( P_0 \), we use geometric progression:

\[
[I - (1-s)P_0]^{-1} = \sum_{k=0}^{\infty} (1-s)^kP_0^k.
\]  
(62)

We note that the last series is convergent for any \( 0 < s \leq 1 \) because \( P_0^k \) has a limit as \( k \to \infty \). Equation (60) becomes then

\[
\pi_d(s) - \pi_d = (1-s)\left( I - s\sum_{k=0}^{\infty} (1-s)^k(P_0^k)^{k+1}\right)(\pi_0 - \pi_d).
\]  
(63)

By noting that \( \sup_{k \geq 1}\|P_0^k\|_2 = \sup_{k \geq 1}\|P_0^k\|_2 \) is finite, we have the desired upper bound on \( \|\pi_d(s) - \pi_d\| \).