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Circuit Models of Stochastic Genetic Networks

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Abstract—We use analogies between chemistry and subthreshold electronics to develop fast, scalable, digitally programmable hardware models of stochastic genetic networks. We also present experimental results from a prototype 0.18µm CMOS chip that was designed for rapid simulations of gene activation, transcription and translation.

I. INTRODUCTION

Genetic networks can be usefully modeled and simulated using analog, digital, or hybrid electronic circuit analogs [1], [2]. In this paper we describe scalable circuit models of computationally-intensive stochastic genetic networks that build on our prior work on modeling chemical reactions by using log-domain transistor circuits [3]. Our models are suitable for rapid simulation, parameter discovery and optimization, and may be useful for answering a wide range of questions in both systems and synthetic biology.

II. MODELING

A. Induction and Activation

Genes are transcribed into messenger RNA (mRNA) by an enzyme known as RNA polymerase (RNAP). The rate of transcription is regulated by other proteins, known as transcription factors (TFs), that promote (activate) or hinder (repress) the action of RNA polymerase. TFs bind to specific DNA regions, known as binding sites. Binding sites are usually clustered into so-called enhancer and promoter regions located near the beginning of the gene. Activator proteins bind at the enhancer site, while repressor proteins usually bind somewhere on the promoter, as shown in Figure 1. We will denote such TFs as X. The inducer or substrate S is typically a small molecule that binds to a TF (whether repressor or activator). Inducer binding converts inactive TFs into active forms that can bind to the DNA, i.e., \( S + X \rightleftharpoons X^* \), where X and \( X^* \) are the inactive and active forms of the TF, respectively. We will assume that this reaction is always at equilibrium. As a result, we have [\( X^* \)] = [S][X]/\( K_{SX} \), where \( K_{SX} \) is the dissociation constant of the reaction. We will also assume that [\( X^* \)] < [S] and [X], i.e., that S and X are present in excess. As a result [S] and [X] are not depleted by the reaction, and can be treated as external inputs to the system.

The relationship between the concentrations of active TFs \( X^* \) and the rate of transcription of a gene is known as its input function. Such functions are often assumed to be proportional to the fraction of DNA binding sites \( B \) occupied by an activator, or left vacant by a repressor. We assume that \( n \) activated TF molecules co-operatively bind to \( B \), and that the binding reaction is always at equilibrium. As \( n \) increases the binding fractions transition abruptly from 0 and \( \beta_{max} \) as the value of [\( X^* \)] crosses a threshold \( K_i \), where \( \beta_{max} \) is the maximal rate of transcription. As a result they may be approximated as two-state, or logic functions [4]:

\[
\begin{align*}
  f_A(X^*) &= \beta_{max} \theta(X^* > K_i) \\
  f_R(X^*) &= \beta_{max} \theta(X^* < K_i)
\end{align*}
\]

where \( f_A \) and \( f_R \) are the input functions for an activator or repressor, respectively, and the “threshold” function \( \theta( ) \) is equal to 1 if its argument is true, and 0 otherwise.

Most genes are regulated by multiple TFs, resulting in multidimensional input functions [5]. In our model, concentrations are represented by currents and the genes have \( N \)-dimensional input functions. In general such input functions cannot be factorized into the product of single-variable functions such as those shown in (1). However, arbitrary Boolean functions can always be written in the canonical “sum of products” (SoP) form, i.e., as the logical OR of \( \text{minterms} \). Minterms of Boolean functions are formed by ANDing together each variable or its inverse.

We realize arbitrary input functions by generalizing the SoP form. Firstly, we do not restrict the \( i \)-product term to a minterm, which must include all \( N \) variables, but allow it to be the product of the normal or negated forms of \( n_i \) variables, where \( 0 \leq n_i \leq N \). Secondly, we do not restrict our output to be the logical OR of the product terms, but allow it to be any arbitrary \textit{weighted sum} of them. Thus, our output is no longer a Boolean variable, but a “soft” or analog OR of terms that are formed by “hard” or binary AND operations. Intuitively, each product term sets its own maximal transcription rate \( \beta_{max} \), and the total transcription rate is found by adding the rates of all active terms. The output, which we refer to as the activation strength \( I_{ACTV} \), controls the rate of transcription and is given by

\[
I_{ACTV} = I_0 \sum_{i=1}^{s} \beta_i \left\{ \prod_{j=1}^{n_i} \{ f_A(X_i), f_R(X_i) \} \right\}
\]

where \( s \) is the number of product terms, \( k \in \{1,2,...,N\} \), the \( \beta_i \) are dimensionless numbers, and \( I_0 \) is a constant reference current. The term within curly braces in (2) denotes the selection of either \( f_A \) or \( f_R \) for any TF \( X_i \). We also define the Boolean variable \( ACTV \) as the logical OR of all the product terms in (2). The gene is being actively transcribed whenever \( ACTV = 1 \), with a non-zero activation
strength \( I_{ACTV} \). It is inactive whenever \( ACTV = 0 \), which also implies \( I_{ACTV} = 0 \).

### B. Transcription and Translation

We shall assume that a single mRNA transcript is directly translated into a single protein. The simplest reasonable model that is suitable for describing either transcription or translation is given by \( X \xrightarrow{\ k \ } Y \xrightarrow{\ \gamma \ } \phi \), where \( X \) and \( Y \) are the reactant and product, respectively, of either process, and \( \phi \) is a degraded, inactive form of \( Y \). The symbols \( k \) and \( \gamma \) represent the rates of first-order unidirectional reactions, and are known as the synthesis and degradation rates of \( Y \), respectively [4]. The frequency response of this system is that of a first-order low-pass filter, with DC gain and time constant given by \( k/\gamma \) and \( 1/\gamma \), respectively. We can exactly emulate these dynamics on-chip by using a current-mode first-order low-pass filter [3]. However, the circuit will run a factor of \( \alpha \) faster than the chemical system, where \( \alpha \gg 1 \) is known as the speedup factor.

RNA polymerase moves at a finite velocity along the gene during transcription. As a result it takes a finite amount of time to “read” a gene and produce each mRNA transcript. This time-span is known as the transcription delay. We assume that gene activation turns on mRNA formation after some transcription delay time \( T_p \). In other words, the concentration of mRNA begins increasing at \( t = T_D \) after \( ACTV \) goes high at \( t = 0 \). However, the delay \( T_D \) is sign-sensitive, and occurs only when the gene is activated. The value of \( T_D = L/r \), where \( L \) represents the length of the gene and \( r \) represents the rate at which RNA-polymerase travels along it. We allow the user to independently set the transcription delay for every gene by making \( T_D \) dependent on an external control parameter.

Fluctuations in protein concentration can be attributed to two sources: intrinsic and extrinsic. Intrinsic noise is uncorrelated across genes, and arises from the inherent stochasticity within the processes of gene activation, transcription and translation. Extrinsic noise is correlated across multiple genes, and arises from fluctuations external to the gene itself, such as changes in TF and inducer concentrations with time. Models of intrinsic noise have been proposed for both prokaryotic and eukaryotic cells [6], [7]. Both models can be summarized by the system of reactions shown in Figure 2(a) for a single active TF \( X \). In prokaryotes, activated TF binding and unbinding to DNA (\( k_u \) and \( \gamma_u \) in Figure 2(a) respectively) occurs on time scales that are much faster than transcription or translation. These processes lead to a quickly-varying square wave in the \( ACTV \) parameter of the gene (represented by \( A \) in Figure 2(b)). These fast variations are low-pass filtered to create a mean value for \( A \), shown as \( \overline{A} \) in Figure 2(c).

We would like to find the noise in mRNA and protein concentrations when \( \overline{A} = 1 \), i.e., the gene is ‘on’ and actively transcribing. For simplicity, we also assume that all \( \beta \)'s are \( 1 \) in (2).

A signal-flow block diagram view of the entire model is shown in Figure 2(b). Both transcription and translation are modeled as first-order low-pass filters with DC gain, as in Section II-B. The cutoff frequencies of the filters are given by \( \tau_r = 1/\gamma_r \) and \( \tau_p = 1/\gamma_p \), respectively, while their DC gains are given by \( A_r = k_r/\gamma_r \) and \( A_p = k_p/\gamma_p \), respectively.

We assume that every reaction flux consists of uncorrelated arrivals or departures of molecules. The difference between the mRNA synthesis and degradation fluxes is integrated into the mRNA concentration \( [mRNA] \). Each flux has a power spectral density equal (PSD) to twice the mean flux, like any Poisson variable. The two mean fluxes are \( k_r \overline{A} \) and \( \gamma_r [mRNA] \), leading to PSDs of \( 2k_r \overline{A} \) and \( 2\gamma_r [mRNA] \), respectively. These PSDs are independent, and therefore the total PSD that is filtered by the transcription filter is the sum of the two. However, at steady state the two fluxes must have identical mean values, and so the total PSD is \( 4k_r \overline{A} \). Similarly, the total PSD that is filtered by the translation filter is \( 4k_p [mRNA] \) in steady state. The resultant steady-state noise model is shown in Figure 2(c).

![Fig. 2. A model of intrinsic noise in gene expression. The model is shown as (a) a set of chemical reactions, (b) a feedback block diagram and (c) simplified block diagram with noise sources. In (c) wavy lines represent the power spectral density of noise in steady state.](image)

The circuits that simulate the induction and binding of transcriptional activators and repressors to DNA can be divided into three blocks: connection matrix, inducer, and activator. Each gene can be

\[
\begin{align*}
\sigma^2_{mRNA} &= k_r \overline{A} \gamma_r = \overline{mRNA} \\
\sigma^2_{prn} &= prn \left( 1 + \frac{k_p}{\gamma_p} \right) \equiv prn(1 + b) \tag{3}
\end{align*}
\]

where \( \overline{mRNA} \) and \( prn = (k_p/\gamma_p) \overline{mRNA} \) are the mean concentrations of mRNA and protein, respectively, and the second equation is valid when \( \gamma_p \ll \gamma_r \), which is typical. The concentration of mRNA has variance equal to its mean, i.e., behaves like a Poisson random variable. However, the upstream noise from transcription increases the variance of the protein concentration by a factor \( (1 + b) \) over a Poisson variable, where \( b = k_p/\gamma_p \) is known as the burst parameter. It is the average number of protein molecules produced from a single mRNA transcript, and is analogous to the current gain \( \beta \) of a bipolar transistor (BJT). As in a BJT, the total output noise power is the sum of \( \beta \)-amplified shot noise from the input (base or mRNA) and shot noise at the output (collector or protein).

### III. Chip Design

Our prototype genetic network chip is 1.5mm \( \times \) 1.5mm in size, contains 6 gene circuits, and accepts up to 16 external (off-chip) input signals. It was fabricated in a standard 0.18\( \mu \)m CMOS process. A connection matrix, consisting of an array of programmable switches, allows any external input to act as an inducer or TF for any gene. Each gene circuit contains programmable circuits for modeling activation, transcription and translation. Programmable parameter values are set on-chip by individually-addressable transistor or capacitor DACs. The outputs generated by the chip consist of each gene’s activation signal \( ACTV \), and associated mRNA and protein concentrations.

#### A. Input Selection and Activation

The circuits that simulate the induction and binding of transcriptional activators and repressors to DNA can be divided into three blocks: connection matrix, inducer, and activator. Each gene can be
activated or repressed by a maximum of $N$ active TFs, which are induced from their inactive forms by inducers. We allow each TF to have its own inducer, resulting in a total of $2N$ input signals for each gene. The connection matrix allows these signals to be connected to any of the $2N$ external inputs. We use previously-described log-domain circuits [3] to mimic the chemical reactions of inducer-TF binding. Single-variable activation functions are generated by simple current comparators that compare the concentration of active TF with a programmable threshold.

We implemented the complete activation function using a generalization of the well-known Programmable Logic Array (PLA) approach for synthesizing arbitrary Boolean functions. In our modification, AND gates are replaced by more complex combinational logic blocks. Each block accepts the previous block’s output, and ANDs it with a member of the set $\{f_A, f_R, 0, 1\}$, thus building up a product term. Two control bits are fed into each block and determine which member of the set is selected for product formation. Once the product terms have been formed they can be combined by an S-input OR gate to create $ACTV$, or added together with programmable weighting factors to create $I_{ACTV}$.

B. Transcription and Translation

The circuits that simulate the processes of transcription and translation of a single gene can be divided into three blocks. The first block contains our model of transcription delay, the second simulates the dynamics of transcription, and the third simulates translation. We used translinear low-pass filter circuits to model the dynamics of both transcription and translation. Translation delays were produced by the circuit shown in Figure 3. The output of the D-type flip-flop is $ACTD$, the delayed version of $ACTV$. It is immediately reset when $ACTV$ goes low. However, it can only go high on rising clock edges, which are delayed by a time $T_D$. The delay was implemented with an $M$-bit long shift register driven by an oscillator. The oscillator frequency can be set by the user over a 10-bit range, thus allowing $T_D$ to be programmed.

The Poisson electronic shot noise statistics of our circuits controllably mimic the molecular shot noise in cells at high SNR. Obtaining very low values of SNR, however, requires impractically small capacitors and bias currents. We designed a feedback loop to solve this problem by artificially generating controllable electronic fluctuations that are consistent with Poisson statistics. We have implemented the loop on-chip for the special case of transcription, since the concentration of mRNA within a cell is usually low enough for it to be the noise-limiting molecular species.

Our circuit is shown in Figure 4. A current $2I_A$ that is proportional to the mRNA degradation rate $\gamma$ is shut off by a pseudo-random waveform generated by the linear-feedback shift register (LFSR). LFSRs produce waveforms with properties close to random telegraph signals, i.e., random, uncorrelated transitions between two states, which in this case result in currents of value 0 or $2I_A$. The waveform has 50% duty cycle, resulting in an average degradation current of $I_A$, as assumed in previous sections. However, in this case the current is not constant, but generated by a square wave that switches pseudorandomly between $2I_A$ and 0. The clock frequency of the LFSR is controlled by a current-controlled oscillator (CCO) whose input is a scaled version of the mRNA concentration, i.e., $\beta_{snr}\gamma_mRNA$, where $\beta_{snr}$ is a dimensionless number that can be programmed by the user. The output frequency of the CCO is given by $f_{cco} = \beta_{snr}\gamma_mRNA / q_{osc}$, where $q_{osc}$ is characteristic to the CCO, and may be viewed as the total charge supplied to the capacitors within the oscillator during one oscillation cycle. The total noise power in $I_{mRNA}$ and $I_{prn}$, the protein concentration, are given by

$$\overline{i^2_{mRNA}} = \left( \frac{q_{osc}/\beta_{snr}}{2\tau_r} \right) I_{mRNA}$$

$$\overline{i^2_{prn}} = b \left( \frac{q_{osc}/\beta_{snr}}{2\tau_r} \right) I_{prn}$$

where $b = k_p/\gamma_r$, as in (3), and the second equation is valid when $\gamma_p \ll \gamma_r$. We see that total noise variances are proportional to means, just like Poisson random variables. In fact, both variances are $\propto 2I_{cco}\gamma_f \Delta f$, where $q_{cco} = \gamma_mRNA/\beta_{snr}$ is the effective charge and $\Delta f = 1/\tau_r$ is the bandwidth. Thus, our circuit replaces the electronic charge $q$ by the much larger charge $q_{cco}$, resulting in lower SNR for the same mean level of mRNA or protein. On this chip we obtained a typical value of $q_{osc} = 9.2 \times 10^5 q$. The SNR at a given mean level can be set by the user through the parameter $\beta_{snr}$ since $SNR \propto \beta_{snr}$ when all other parameters are fixed. The current circuit only models noise from transcription, not translation. We can model translation noise in future chips if necessary with a similar circuit.

IV. CHIPS MEASUREMENTS

The gene network simulator chip was mounted on a printed circuit board and interfaced with user interface software (written in MATLAB) running on a desktop computer. We successfully programmed the input function, activation threshold, transcription delay, transcription and translation rates and delay times of each gene circuit. We also tested the SNR-adjustment circuitry. We know that the SNR of a standard Poisson variable $N$ is equal to its mean value $\mathbb{N}$, i.e., the mean number of random events that occur during a counting (integration) period. The mRNA concentration $I_{mRNA}$ can be normalized into a standard Poisson variable $\mathbb{N}_{mRNA} = I_{mRNA} / (I_{mRNA} / \mathbb{N}_{mRNA}) = SNR (I_{mRNA} / \mathbb{N}_{mRNA})$. Thus $\mathbb{N}_{mRNA}$ should obey the well-known Poisson probability distribution, and only SNR should affect it. We estimated probability distributions at various values of SNR by varying $\beta_{snr}$ and measuring $I_{mRNA}$ during a time interval of $T = 200\mu$s when transcription was active. Figure 5 shows both measured and theoretically-predicted probability distributions. The excellent match between the two distributions shows that the stochastic behavior of our chip is as expected.

Equation (4) shows that the SNR of mRNA, i.e., $I_{mRNA}^2 / \mathbb{N}_{mRNA}$, is proportional to the product $\beta_{snr}I_{mRNA}$. Therefore, it should increase linearly with $I_{mRNA}$ when $\beta_{snr}$ is fixed. The protein

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Fig. 3. A simple circuit that generates sign-sensitive transcription delay.

Fig. 4. Block diagram of the SNR adjustment circuit.
concentration should behave similarly, but with an excess noise factor. We verified this prediction experimentally by changing the input to the transcription filter, i.e., the activation strength $I_{ACTV}$. The mean value and SNR of both mRNA and protein should be proportional to $I_{ACTV}$. We varied $I_{ACTV}$ over a 32:1 range while maintaining a fixed value of $\beta_{snr} = 10$. We also kept the following parameter values fixed: $\gamma_p = \gamma_X$, $A_p = 2$ and $b = 2$. The results are shown in Figure 6. We see that the SNR increases linearly with mean concentration in both cases, as expected.

We used three of the gene circuits on our chip to create a coherent FFL. The gene producing $Z$ was programmed to have an AND-like input function. Figure 7 shows measured concentrations of the proteins transcribed from genes $X$, $Y$ and $Z$ in response to square inducer pulses $S_X$ of two different durations. The circuit behaves as expected. It exhibits sign-sensitive delay: $Z$ follows rising transients of $Y$, but only after $\approx 60 \mu s$. However, it immediately responds to falling transients. In addition, the circuit distinguishes between short inducer pulses (which do not activate $Z$) and long pulses.

We have described integrated circuits for high-speed simulations of stochastic genetic networks. They may be useful for analyzing complex networks, particularly those involved in disease.

**REFERENCES**


