Stochastic Simulation of Enzyme-Catalyzed Reactions with Disparate Timescales

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ABSTRACT Many physiological characteristics of living cells are regulated by protein interaction networks. Because the total numbers of these protein species can be small, molecular noise can have significant effects on the dynamical properties of a regulatory network. Computing these stochastic effects is made difficult by the large timescale separations typical of protein interactions (e.g., complex formation may occur in fractions of a second, whereas catalytic conversions may take minutes). Exact stochastic simulation may be very inefficient under these circumstances, and methods for speeding up the simulation without sacrificing accuracy have been widely studied. We show that the "total quasi-steady-state approximation" for enzyme-catalyzed reactions provides a useful framework for efficient and accurate stochastic simulations. The method is applied to three examples: a simple enzyme-catalyzed reaction where enzyme and substrate have comparable abundances, a Goldbeter-Koshland switch, where a kinase and phosphatase regulate the phosphorylation state of a common substrate, and coupled Goldbeter-Koshland switches that exhibit bistability. Simulations based on the total quasi-steady-state approximation accurately capture the steady-state probability distributions of all components of these reaction networks. In many respects, the approximation also faithfully reproduces time-dependent aspects of the fluctuations. The method is accurate even under conditions of poor timescale separation.

INTRODUCTION

Stochastic simulation is important for the study of chemical reaction systems where the number of molecules of some species can become small. The exact stochastic simulation algorithm of Gillespie (1, 2) is the standard approach to simulation in this setting, but can become too slow to be useful in cases where fast reactions dominate the calculations but slow reactions determine the interesting evolution of the overall system. This may be the case, for example, in enzyme-catalyzed reactions where binding and unbinding of the enzyme and substrate often occur much more frequently than the product-producing reaction. Gillespie's algorithm will spend most of its time tracking binding and unbinding reactions that essentially cancel each other out and produce no net conversion of substrate into product.

Many approximations to Gillespie's algorithm have been proposed to reduce the bottleneck that occurs in systems with disparate timescales. Tau-leaping (3–6) uses Poisson random variables to allow multiple firings of reaction channels per time step. This approach can speed up simulations, but the explicit version has difficulty with the stiff systems we consider. The implicit version (5) was proposed to solve stiff systems; however, practical τ-selection formulas were only proposed (6) for the explicit version. A τ-selection formula based on reversible reactions was proposed, but a formula for a general stiff system is not yet available. Other approaches rely on separating the system into slow and fast reactions or species. The slow reactions can be simulated with Gillespie's algorithm and the fast reactions can be handled in a more efficient manner. One approach is to simulate fast reactions using chemical Langevin equations (7, 8). Other techniques avoid simulating the fast reactions altogether, instead using their average values to derive the firing propensities of the slow reactions. The average values are obtained using the quasi-steady-state approximation (QSSA) (9) or the quasi-equilibrium approximation (also known as the partial equilibrium assumption) (10–12).

In this article, we apply an idea, called the total QSSA (TQSSA), to provide an algorithmic, fast, and accurate way to stochastically simulate enzyme-catalyzed reactions. The TQSSA was developed in a deterministic setting to remove the restrictions associated with the standard QSSA applied to enzyme-catalyzed reactions (13). The TQSSA accurately tracks intermediate complexes and can be extended straightforwardly to complex systems of interacting enzymes (14). In the stochastic setting considered here, TQSSA allows Gillespie's algorithm to be used to simulate the slow reactions while analytically accounting for the abundances of the fast species, as in the methods mentioned above. Thus, it avoids the bottleneck associated with simulating the fast reactions directly. The main contribution of this article is to show that stochastic TQSSA generalizes the QSSA results of Rao and Arkin (9) and can improve the accuracy of the partial equilibrium assumption employed by others (10–12) by allowing the slow timescales to affect the quasi-steady-state of the fast complexes. We provide a general framework for applying stochastic TQSSA to chemical reaction networks and illus-
trate its usefulness on a complex network exhibiting alternative stable steady states. Our results show that for cases having reasonable timescale separation between fast and slow reactions, the stochastic TQSSA simulations are nearly identical to direct simulation of all the reactions using Gillespie’s stochastic simulation algorithm.

MICHAELIS-MENTEN KINETICS

Consider a simple chemical reaction where a substrate (S) is converted irreversibly to a product (P) by an enzyme (E):

\[ E + S \xrightarrow{k_1} E:S \xrightarrow{k_2} P + E. \] (1)

In these reactions, \( k_1 \) and \( k_2 \) are the rate constants for formation and dissociation of the enzyme-substrate complex (E:S), and \( k_2 \) is the rate constant for the enzymatic conversion of S to P. With square brackets denoting the concentration of a chemical species, the rate equations corresponding to the above elementary reactions can be written as

\[
\frac{d[S]}{dt} = -k_1[S][E] + k_2[E:S] ;
\]

\[
\frac{d[E:S]}{dt} = -(k_1 + k_2)[E:S] + k_2[S][E] - [E:S] .
\] (2)

where \( [E] + [E:S] \) is the total enzyme concentration. We always use subscript \( T \) to indicate the total amount of a chemical moiety. If the concentration of substrate is very much greater than \( ET \), then the change in concentration of the enzyme-substrate complex (Eq. 3) will be fast in comparison to the change in concentration of the free substrate. Under these conditions, the enzyme-substrate complex rapidly reaches its stationary state, i.e.,

\[
\frac{d[E:S]}{dt} = -(k_1 + k_2)[E:S] + k_2[S][E] - [E:S] = 0 ,
\]

\[
[E:S] = \frac{[E]S}{K_m + [S]},
\] (4)

where \( K_m = (k_1 + k_2)/k_1 \) is called the Michaelis constant of the enzyme. Equation 4 is the QSSA for the enzyme-substrate complex. In this case, the conversion of substrate into product is described by the well known Michaelis-Menten rate law:

\[
\frac{d[S]}{dt} = -k_2[E:S] = \frac{k_2[E][S]}{K_m + [S]} .
\] (5)

The condition for validity of the QSSA is \( ET \ll S_0 + K_m \), where \( S_0 \) is the initial substrate concentration.

When this condition is not satisfied, then both \([S]\) and \([E:S]\) change on the same timescale. However, when \( k_2 \ll k_1 \), the “total” substrate concentration, \([S] = [S] + [E:S]\), changes on a slower timescale than the enzyme-substrate complex (13):

\[
\frac{d}{dt}P(\hat{S},es;t) = \left[-(k_1(\hat{S} - es) + (k_1 + k_2)es) + (k_1 + k_2)es\right]P(\hat{S},es;t) \\
+ k_1(\hat{S} - es + 1)(es + 1 + 1)P(\hat{S},es - 1; t) \\
+ k_2(1+es + 1)P(\hat{S},es + 1; t) \\
+ k_1(1+es + 1)P(\hat{S},es + 1; t) + k_2(es + 1)P(\hat{S} + 1, es + 1; t). (11)
\]
Allowing for a slight abuse of notation, \( es \) refers to the numerical value of the random variable \( N_{ES} \), not the product of two numbers, \( e \) and \( s \). In addition, we use an upper-case italic letter to denote the concentration of a particular species and a lower-case italic letter to denote its number of molecules in a specified volume (e.g., the volume of a yeast cell). For example, \( E_T \) = total concentration of enzyme, whereas \( e_T \) = total number of enzyme molecules.

To obtain an equation for the probability of the composite variable, \( N_{ES} \), we write the joint probability as

\[
\Pr(es; t) = \Pr(es_1, eT, es_2, t) = \prod_{i=1}^{n} \Pr(es_i; t)
\]

because all terms relating to the fast reactions cancel out. For cases of large timescale separation, the distribution of \( es \) should reach steady state quickly compared to the evolution of \( \tilde{s} \) and we can eliminate the dependence on \( t \) in the expected value of \( es \) in Eq. 13. In this case, Eq. 13 exactly describes the noise in the chemical reaction

\[
\tilde{s} \xrightarrow{k_2(es)} P.
\]

Since in most cases we expect the differential equation description of chemical reactions to track the mean values of the reactants, it seems reasonable to replace \( \langle es|\tilde{s}\rangle \) by the steady-state value of \( es \) from the TQSSA, given by

\[
\langle es|\tilde{s}\rangle = (e_T + K_a + \tilde{s})es + e_T\tilde{s} = 0.
\]

To investigate the accuracy of our approach, we compute the exact value of \( \langle es|\tilde{s}\rangle \) and compare it to the TQSSA value. Assuming that \( \tilde{s} \) is constant on the short timescale, the evolution of the probability of \( N_{ES} \) is governed by the equation

\[
\frac{d}{dt} \Pr(es; t) = -k_1(\tilde{s} - es)(e_T - es) + k_{-1}es\Pr(es; t)
\]

\[
+ k_1(\tilde{s} - es + 1)(e_T - es + 1)\Pr(es + 1; t)
\]

\[
+ k_{-1}(es + 1)\Pr(es + 1; t)
\]

\[
= \Delta J(es + 1; t) - \Delta J(es; t),
\]

where \( \Delta J(es; t) = k_{-1}es\Pr(es; t) - k_1(\tilde{s} - es + 1)(e_T - es + 1) \Pr(es + 1; t) \). At steady state (on the fast timescale), \( \Delta J(es; t) \) is a constant, and (considering the case \( es = 0 \)) the constant must be zero. This equation provides a recurrence relation for \( \Pr(es; t) \) whose solution is

\[
\Pr(es; t) = \prod_{i=1}^{n} \left\{ \frac{k_i(\tilde{s} - i + 1)(e_T - i + 1)}{k_{-1}} \right\} \Pr(0)
\]

\( 0 \leq es \leq \min(\tilde{s}, e_T) \).

The condition that the total probability sums to 1 allows the determination of \( \Pr(0) \), and then the moments \( \langle es|\tilde{s}\rangle \) and \( \langle es^2|\tilde{s}\rangle \) can be computed exactly. Fig. 1a compares the function \( \langle es|\tilde{s}\rangle \) with the deterministic TQSSA result for the case \( e_T = 10 \) with \( k_1 = k_{-1} = 1.0 \) and \( k_2 = 0.1 \) (arbitrary units). Clearly the stochastic mean \( \langle es|\tilde{s}\rangle \) is very close to the deterministic value of \( es \) given by Eq. 15, even for this case of very small numbers of molecules. The variance \( \langle (es - \langle es\rangle)^2|\tilde{s}\rangle \) is shown in Fig. 1b and is quite small, on the order of one molecule or less.

This analysis is quite similar to that in Goutsias (10) and Cao et al. (11,12). Our approach differs in using Eq. 15 for the mean value of the complex. Goutsias and Cao assume that the association and dissociation of the enzyme-substrate complex reaches equilibrium independently of the substrate-to-product conversion reaction, and so their value for \( \langle es|\tilde{s}\rangle \) will not depend on the slow rate constant \( k_2 \), whereas our value does. For cases of extreme timescale separation (\( k_{-1} > 100 k_2 \), say), all these approaches tend to the same solution, and \( \langle es|\tilde{s}; t\rangle \) in Eq. 15 will tend quickly to a constant independent of \( k_2 \). For less extreme cases, the term \( (es - \langle es\rangle)|\tilde{s}\rangle \) approaches a steady value dependent on both fast and slow reactions. Only the TQSSA captures this average value accurately, and the reaction (Eq.

![FIGURE 1 Michaelis-Menten kinetics. (a) Comparison of steady-state mean of enzyme-substrate complex as a function of \( \tilde{s} \) for \( e_T = 10 \). The solid line plots \( \langle es|\tilde{s}\rangle \) calculated from the steady-state probability distribution, \( \Pr(es; t) \), and the dashed line plots deterministic TQSSA results from Eq. 12. (b) Steady-state variance of the enzyme-substrate complex as a function of \( \tilde{s} \), \( \langle (es - \langle es\rangle)^2|\tilde{s}\rangle \), calculated from \( \Pr(es; t) \).](https://example.com/figure1.png)
(14), using the corresponding time-invariant propensity, will satisfy a master equation that approximates the actual master equation (Eq. 11). Thus the TQSSA is expected to be more accurate than the other approaches in cases where the timescale separation is not large. An example of this behavior is shown in the Supplementary Material (Data S1).

We examine the accuracy and efficiency of stochastic TQSSA by simulating both the full reaction mechanism (Eq. 1) (we call this “full Gillespie simulation”) and the simplified reaction (Eq. 14), with propensity $k_2(\varepsilon_1)$ given by the steady-state value of $\varepsilon_1$ from the TQSSA expression (Eq. 15). We do the comparison for different initial numbers of enzyme and substrate molecules. Of two roots obtained from the quadratic equation (Eq. 15), we take only the root that satisfies $0 < \varepsilon_1 < \varepsilon_T$. We have chosen $k_1 = k_{-1} = 1.0$ (arbitrary units), and the numerical simulations are averaged over 50,000 realizations. In Fig. 2, we plot the mean number of product $\langle p \rangle$ and its standard deviation as a function of time for two different cases: 1) $\varepsilon_T \gg \varepsilon_0$ and 2) $\varepsilon_T = \varepsilon_0$.

From the figure, it is clear that in both cases the stochastic TQSSA is in excellent agreement with the full Gillespie simulation. The stochastic QSSA proposed by Rao and Arkin does not give correct results in the two cases considered here. (For the case $\varepsilon_T \ll \varepsilon_0$ (not shown), stochastic QSSA is accurate, as expected.) The approximation proposed by Goutsias gives good results in the cases we consider, but our results are more accurate (see the Supplementary Material, Data S1, Fig. S2, a–c). As expected, the stochastic TQSSA simulation is very much faster than the full Gillespie simulation (15 times faster in the cases considered here).

**GOLDBETER-KOSHLAND SWITCH**

We now extend our stochastic TQSSA method to coupled enzymatic reactions: the well-known Goldbeter-Koshland (GK) ultrasensitive switch (16). The GK switch consists of a substrate-product pair ($S$ and $S_p$) that are interconverted by two enzymes ($D$ and $E$). One can imagine that $S_p$ is the phosphorylated form of the protein $S$, enzyme $E$ is a kinase acting on $S$, and enzyme $D$ is a phosphatase acting on $S_p$. The six reactions involved are:

$$\frac{dS_p}{dt} = G(V_e, V_d, J_e, J_d) = \frac{2V_dJ_d}{\sqrt{(V_d - V_e + V_dJ_e + V_eJ_d)^2 - 4(V_d - V_e)V_eJ_d}} - \frac{2V_dJ_d}{\sqrt{(V_d - V_e + V_dJ_e + V_eJ_d)^2 - 4(V_d - V_e)V_eJ_d}}.$$

The steady-state response of $S_p$ as a function of $E_T$ shows a steep sigmoidal shape if $J_e \ll 1$ and $J_d \ll 1$. The steepness originates from the kinetic interplay of the two substrate-saturated enzymes acting in opposite directions.

The classic GK ultrasensitive switch requires that substrate concentrations be much larger than enzyme concentrations, which can be expected for metabolic control networks, where concentrations of metabolites are much larger than those of...
enzymes. But the applicability of Michaelis-Menten kinetics is limited in the context of protein interaction networks, because a particular enzyme may act on multiple substrates, a particular substrate may be acted upon by multiple enzymes, and enzymes and substrates often alternate their roles. When the enzyme and substrate swap roles, Michaelis-Menten kinetics is no longer applicable, since the condition $E_T \ll S_0 + K_m$ cannot possibly be true for both reactions simultaneously. In this case it is better to write the rate equations in terms of $[\hat{S}_p] = [S_p] + [D:S_p]$

$$\frac{d[\hat{S}_p]}{dt} = k_2'(E:S) - k_2''[D:S_p]; \quad \text{(22)}$$

$$\frac{d[D:S_p]}{dt} = k_1'(D_T - [D:S_p]) ([\hat{S}_p] - [D:S_p])$$

$$- k_1''[D:S_p] - k_2''[D:S_p]; \quad \text{(23)}$$

$$\frac{d[E:S]}{dt} = k_1''(E_T - [E:S]) ([\hat{S}_p] - [E:S])$$

$$- k_1''[E:S] - k_1''[E:S]; \quad \text{(24)}$$

where $S_T = [\hat{S}_p] + [\hat{S}]$ and $[\hat{S}] = [S] + [E:S]$. Invoking the TQSSA on Eqs. 23 and 24, as done by Ciliberto et al. (14), we find that the reactions portrayed in Eqs. 18 and 19 are governed by a single “slow” differential equation (Eq. 22) with $[E:S]$ and $[D:S_p]$ given as functions of $[\hat{S}_p]$ by the solution of the pair of quadratic equations:

$$[D:S_p]^2 - (D_T + K_m + [\hat{S}_p])[D:S_p] + D_T[\hat{S}_p] = 0; \quad \text{(25)}$$

$$[E:S]^2 - (E_T + K_m + S_T - [\hat{S}_p])[E:S] + E_T([S_T - [\hat{S}_p]] = 0, \quad \text{(26)}$$

with $0 \leq [D:S_p] \leq D_T$ and $0 \leq [E:S] \leq E_T$.

As before, we can write a chemical master equation for the entire system and then sum over the fast variables to get a master equation for the slow composite variable $N_{\hat{S}_p} = \hat{S}_p$:

$$\frac{d}{dt}P(\hat{S}_p; t) = -[k_2'(E:S) + k_2''d_{sp}[\hat{S}_p; t])P(\hat{S}_p; t)$$

$$+ k_2'(d_{sp}[\hat{S}_p; t + 1; t]P(\hat{S}_p + 1; t)$$

$$+ k_2''(E:S) - k_2''[D:S_p; t]P(\hat{S}_p - 1; t)], \quad \text{(27)}$$

(for details, consult the Supplementary Material, Data S1). Assuming that the expected values equilibrate on a faster timescale, we can reduce Eq. 27 to a master equation for the reactions

$$\frac{d}{dt}N_{\hat{S}_p} = k_2'(d_{sp}[\hat{S}_p; t])S, \quad \text{(28)}$$

with propensities $k_2'(d_{sp}[\hat{S}_p; t])$ and $k_2''(E:S)[\hat{S}_p; t]$ for the forward and backward reactions, respectively. If we look at the evolution of the fast variables in between transitions of the slow variables, the equations for $es$ and $ds_p$ are analogous to the equation for $es$ in the simple enzyme-catalyzed reaction.

For the stochastic TQSSA, numerical simulation of the reactions in Eq. 28 is carried out using Gillespie’s stochastic simulation algorithm. At each time step, we calculate the propensities by solving the quadratic equations (Eqs. 25 and 26) in terms of numbers of molecules rather than concentrations. Of the multiple roots obtained from the solution of the quadratic equations we take only that root for which $0 \leq es \leq e_T$ and $0 \leq ds_p \leq d_T$. To compare the full Gillespie simulation involving six reactions (Eqs. 18 and 19) with the stochastic TQSSA simulation, we consider the parameter set $s_T = 900$ molecules, $k_2' = 0.05555$, $k_2'' = 0.83$, $k_2''' = 0.17$, $k_2'''' = 0.05$, $k_2''' = 0.8$, and $k_2'''' = 0.1$ (all the rate constants with subscript “1” have the unit molecule$^{-1}$ min$^{-1}$, and the other rate constants have the unit min$^{-1}$). These parameter values are consistent with Ciliberto et al. (14). In Fig. 3, we plot the time evolution of mean $\langle \hat{S}_p \rangle$ for three different total numbers of kinase and phosphatase: (a), $e_T = 45$, $d_T = 9$; (b), $e_T = 450$, $d_T = 90$; and (c), $e_T = 4500$, $d_T = 900$. In each case, the middle line corresponds to the mean value of $\hat{S}_p$ and the lines above and below are $\pm \sigma$ (1 SD) from the mean. For the initial condition, we used $N_{\hat{S}_p}(0) = 900$, $N_{\hat{S}_p}(0) = 0$, $N_{DS_p}(0) = 0$, and $N_{ES}(0) = 0$. To get the mean, we have averaged each case over 50,000 trajectories. In Fig. 3, we have plotted the results from both the full Gillespie and stochastic TQSSA simulations; the agreement is so good that it is difficult to distinguish between the two results.

Fig. 4 compares the steady-state probability distributions of $N_{\hat{S}_p}$ calculated from full Gillespie and stochastic TQSSA simulations. We have chosen $d_T = 90$ and $e_T = 450$, with the same initial conditions as used previously. The agreement between the two distributions is so good that it is difficult to distinguish between the two lines. We have found that the steady-state probability calculated from stochastic TQSSA matches very well with full Gillespie simulations for both $\hat{S}_p$ and $\hat{S}$ for any numbers of total enzymes, $d_T$ and $e_T$.
FIGURE 4 Goldbeter-Koshland switch. Steady-state probability distributions of $\hat{S}_p$ for $s_T = 900$, $d_T = 90$, and $e_T = 450$. The solid line represents the full Gillespie, and the dashed line the stochastic TQSSA.

In Fig. 5, we compare full Gillespie simulations with stochastic TQSSA for steady-state means and standard deviations of $\hat{S}_p$ and $D:Sp$. We choose the same parameters as before, with $s_T = 900$, $d_T = 90$ and with different values of $e_T$. It is clear that for any values of initial enzyme and substrate concentrations, stochastic TQSSA always predicts the mean accurately for both $\hat{S}_p$ and the complex $D:Sp$. It always predicts the standard deviation for $\hat{S}_p$ accurately. For the complex, the standard deviation of $D:Sp$ predicted by stochastic TQSSA is generally good. It fails for the cases where the number of free enzyme molecules is very small (see Fig. 5 b, $E_T = 180$ and 270; see also Table S3 in the Supplementary Material, Data S1). Fig. 5 a also illustrates the switching behavior of $\hat{S}_p$ with increasing $e_T$. (In the Supplementary Material (Data S1), we present more comparisons of full Gillespie with stochastic TQSSA simulations.)

In Fig. 6, we report the time autocorrelation function, $C(t)$, of steady-state fluctuations around the mean for different variables. We define the autocorrelation function for any variable $x$ as

$$C_x(\tau) = \frac{\langle (x(t) - x_m)(x(t + \tau) - x_m) \rangle}{\langle (x(t) - x_m)^2 \rangle},$$

where $x_m$ is the steady-state value of $x$. To get a smooth profile for an autocorrelation function, we averaged results over 10,000 trajectories. In general, the autocorrelation functions for various species in this network decay exponentially. In Fig. 6 a we show that the autocorrelation function of $\hat{S}_p$ calculated from stochastic TQSSA simulations matches very well with full Gillespie simulations. In Fig. 6 b, we compare autocorrelation functions for free species $Sp$. TQSSA always predicts a single exponential decay of the autocorrelation function, but in some cases, full Gillespie calculations show biexponential decay with a short and a long component of the decay. From the figures, it is clear that stochastic TQSSA predicts the long time component accurately but it can’t predict the short time component. We find a similar trend for the complex $D:Sp$ (Fig. 6 c) as well. The failure of stochastic TQSSA simulations to capture the fast timescale of decaying autocorrelations is to be expected, because all fast-timescale events are averaged out in the quasi-steady-state approximations (Eqs. 25 and 26).

GENERAL FORMULATION OF THE TQSSA

To provide a general setting for the TQSS approach to stochastic simulation taken in this article, we first develop a general framework for the deterministic TQSSA and then extend it to a stochastic formulation. Systems of enzyme-catalyzed reactions, the focus of this article, fit naturally into this general framework, but the framework encompasses a much broader class of systems.

Consider a chemical reaction network with $N$ species participating in $M$ independent reactions. The deterministic chemical rate equations can be written as $dx/dt = S \cdot r(x(t))$, where $x(t)$ is an $N$-vector of concentrations of reacting chemical species, $S$ is an $N \times M$ matrix of stoichiometric coefficients, and $r$ is a vector of time-dependent fluxes through the $M$ reactions of the network. (These fluxes are, in
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general, nonlinear functions of the concentrations of the reacting species.) We assume that the reactions have been partitioned into \( M_s \) slow reactions and \( M_f \) fast reactions, \( r = \left( \frac{r_s}{r_f} \right) \), where \( M_s + M_f = M \). If the reactions are ordered in this fashion, then the stoichiometric matrix can be separated into two submatrices, \( S = [S_{\text{slow}} S_{\text{fast}}] \), where \( S_{\text{slow}} \) is an \( N \times M_s \) matrix of stoichiometric coefficients for the slow reactions, and \( S_{\text{fast}} \) is an \( N \times M_f \) matrix of stoichiometric coefficients for the fast reactions.

Although it may be possible to separate reactions into slow and fast categories, it is usually impossible to separate chemical species into these categories because most chemical species participate in both fast and slow reactions. Using the TQSS idea, we seek linear combinations of the original chemical species (e.g., \( \hat{x}_s = \mathbf{1}^T \cdot x \), where \( \mathbf{1} \) is an \( N \times N \) matrix of constants) such that \( d\hat{x}_s/dt \) involves only slow reactions, \( r_s \). In such a case,

\[
d\hat{x}_s/dt = \mathbf{1}^T d\mathbf{x}/dt = \mathbf{1}^T [S_{\text{slow}} S_{\text{fast}}] \begin{bmatrix} \mathbf{r}_s \\ \mathbf{r}_f \end{bmatrix} = [S_s \mathbf{0}] \begin{bmatrix} \mathbf{r}_s \\ \mathbf{r}_f \end{bmatrix} . \tag{30}
\]

Hence, the condition for a slow combination of species, \( \hat{x}_s = \mathbf{1}^T \cdot x \), is that the columns of \( \mathbf{1} \) span the left null space of the \( N \times M_f \) matrix \( S_{\text{fast}} \). The number of slow variables, \( N_s \), is the dimension of this left null space. A linear transformation matrix \( \mathbf{L} \) can be constructed by choosing the first \( N_s \) rows of \( \mathbf{L} \) as appropriate linearly independent components of the null space, and the remaining rows as standard basis vectors that are linearly independent of the first \( N_s \) rows. (It is clear that \( \mathbf{L} \) is not unique, and this freedom can be used for other purposes.) The transformation \( \hat{x}_s = \mathbf{L} \cdot x \) produces chemical rate equations of the form (the hat is dropped for notational simplicity):

\[
\frac{d}{dt} x_s(t) = S_s \begin{bmatrix} \mathbf{r}_s(x_s, x_f) \\ \mathbf{r}_f(x_s, x_f) \end{bmatrix} \tag{31}
\]

where the slow variables, \( x_s \), participate only in slow reactions, and the fast variables, \( x_f \), are governed by both fast and slow reactions. The \( \mathbf{0} \) matrix in Eq. 31 has \( N_s \) rows and \( M_f \) columns, and the stoichiometry matrix is equal to \( \mathbf{L} \cdot [S_{\text{slow}} S_{\text{fast}}] \).

In terms of the preceding examples, the association and dissociation of enzyme-substrate complexes are chosen as the fast reactions, \( x_s \) corresponds to the total variables defined by TQSSA (denoted with hats in our examples), and \( x_f \) corresponds to the original complexes in the system. Any conservation relations in the system show up as all zero rows of \( S_s \) and can be eliminated from the differential equation. By construction there are no linear combinations of species that are governed only by slow reactions among the \( N_f = N - N_s \) “fast” variables.

In the deterministic formulation of TQSSA, the derivatives of the fast variables are set equal to zero and the resulting equations solved for the quasi-steady-state values of the fast variables in terms of the slow variables. This in turn allows for the simulation of the slow subsystem with the values for the fast variables replaced by their quasi-steady-state values.

To see how this idea transfers to the stochastic setting, consider the chemical master equation for the entire system (dropping the bold face):
\[
\frac{\partial P(x_i, x_j; t)}{\partial t} = \sum_{\text{slow}} a_i(x_i - S_{ij}, x_j - S_{df}, t) P(x_i - S_{ij}, x_j - S_{df}, t) + \sum_{\text{fast}} a_i(x_i, x_j - S_{df}, t) P(x_i, x_j - S_{df}, t) - \sum_{\text{slow}} a_i(x_i, x_j) P(x_i, x_j; t) - \sum_{\text{fast}} a_i(x_i, x_j) P(x_i, x_j; t),
\]

(32)

where \( a_i \) is the propensity of the \( i \)th reaction. Writing \( P(x_i, x_j; t) = P(x_i|x_j; t) \times P(x_j; t) \) in the above and summing over the fast species gives

\[
\frac{\partial P(x_i; t)}{\partial t} = \sum_{x_j} \left\{ \sum_{\text{slow}} a_i(x_i - S_{ij}, x_j - S_{df}, t) P(x_i - S_{ij}, x_j - S_{df}, t) P(x_i - S_{ij}; t) + \sum_{\text{fast}} a_i(x_i, x_j - S_{df}, t) P(x_i, x_j - S_{df}, t) P(x_i; t) - \sum_{\text{slow}} a_i(x_i, x_j) P(x_i, x_j; t) - \sum_{\text{fast}} a_i(x_i, x_j) P(x_i, x_j; t) \right\},
\]

(33)

Defining

\[
\langle a_i(x_i, x_j)|x_i; t \rangle = \sum_{x_j} a_i(x_i, x_j) P(x_i|x_j; t),
\]

(34)

and recognizing that \( \langle a_i(x_i, x_j)|x_i; t \rangle = \langle a_i(x_i, x_j - \alpha)|x_i; t \rangle \), allows us to write

\[
\frac{\partial P(x_i; t)}{\partial t} = \sum_{x_j} \langle a_i(x_i - S_{ij}, x_j; t)|x_i - S_{ij}; t \rangle P(x_i - S_{ij}; t) + \sum_{x_j} \langle a_i(x_i, x_j)|x_i; t \rangle P(x_i; t) - \sum_{x_j} \langle a_i(x_i, x_j)|x_i; t \rangle P(x_i; t).
\]

(35)

This can be recognized as the master equation of the system consisting of just the slow reactions involving the species \( x_i \), having stoichiometric matrix \( S_i \) and propensities \( \langle a_i(x_i, x_j)|x_i; t \rangle \).

If the original slow reaction rates depend at most linearly on the fast variables \( x_f \), which will be the case if no slow variable is the result of a bimolecular reaction where both reactants are fast variables, then \( \langle a_i(x_i, x_j)|x_i; t \rangle \) will depend only on the slow variables and the means of the fast variables given the values of the slow variables. (This was the major restriction in Goutsias (10).) We make the further assumption that the differential equations governing the system provide accurate information concerning the means of the variables. Then, assuming that the fast variables go to steady state quickly compared to the evolution of the slow variables (the standard TQSSA assumption), we can set the derivatives of the fast variables to zero, yielding

\[
S_{df} \frac{\partial P}{\partial x_f} (x_i, x_f) + S_{df} \frac{\partial P}{\partial x_f} (x_i, x_f) = 0.
\]

(36)

Equation 36 provides \( N_f \) coupled equations in \( N \) unknowns. Given the assumption of mass-action kinetics, these equations will be at most second order in the variables, \( x_i \) and \( x_f \). In most cases, we can solve these equations for the \( N_f \) fast variables in terms of the slow variables, providing us with the means of the fast variables needed to compute the rates of the slow reactions of the reduced stochastic system. Clearly, as pointed out by Goutsias (17), the solution of the deterministic chemical reaction equations can differ from the mean values of the solution of the stochastic reaction equations when the molecular populations approach zero. Although not a concern for the problems considered in this article, this potential limitation must be kept in mind for the general case.

COUPLED GK SWITCHES (BISTABILITY)

We now apply this general approach to coupled GK switches (Fig. 7 a). To reactions shown in Eqs. 18 and 19, we add a second GK switch, which converts unphosphorylated \( E \) to a phosphorylated form \( \text{EP} \) and back. The conversion of \( E \) to \( \text{EP} \) is catalyzed by \( S \), and the dephosphorylation reaction of \( E \) is catalyzed by the phosphatase, \( F \). In this network, \( S \) and \( E \) act as mutual antagonists. This network describes the transition from the \( G_2 \) phase to mitosis (the M phase) in the eukaryotic cell cycle, where \( S = \text{MPF} \) (M-phase promoting factor, a dimer of Cdc2 and cyclin B) and \( E = \text{Wee1} \) (a kinase that...
phosphorylates and inactivates Cdc2). Novak and Tyson (18) used this module in a mathematical model for the G2-to-M transition during early embryonic cell cycles of *Xenopus*. Using GK rate laws, like Eq. 20, they showed that the concentration of S may exhibit two stable steady-state values, \(|S| \approx 0\) and \(|S| \approx S_T\), for intermediate values of \(S_T\). That is to say, the control system (Fig. 7 a) can exhibit bistability for \(S_T\) in a certain interval \((S_1 < S_T < S_2)\). Ciliberto et al. (14) showed that bistability is maintained after unpacking the reaction network into elementary steps, provided that \(S_p\) has some ability to bind E but limited ability to convert E to \(E_p\).

To apply the TQSSA to the resulting network, they introduced the definitions

\[
\dot{\hat{S}} = S + [E:S] + [S:E]
\]

\[
\dot{\hat{E}} = E + [E:S] + [S:E] + [S_p:E]
\]

\[
\dot{S_T} = S + S_p + [E:S] + [S:E] + [D:S_p] + [S_p:E]
\]

\[
\dot{D_T} = D + [D:S_p]
\]

\[
\dot{F_T} = F + [F:E_p].
\]

(37)

In terms of these variables, the TQSSA equations are

\[
\frac{d\dot{\hat{S}}}{dt} = k_1^p[D:S_p] - k_2^p[E:S]
\]

\[
\frac{d\dot{\hat{E}}}{dt} = k_2^p[F:E_p] - k_2^p[S:E] - k_2^p[S_p:E]
\]

(38)

and

\[
[D:S_p]^2 - (D_T + K_{md} + S_T - \hat{\hat{S}} - [S_p:E])[(D + [D:S_p] + D_T[S_T - \hat{\hat{S}} - [S_p:E]) = 0
\]

\[
\]

\[
\]

\[
[S_p:E]^2 - (S_T - \hat{\hat{S}} - [D:S_p] + K_{mb} + \dot{\hat{E}} - [E:S] - [S:E])[S_p:E] + (S_T - \hat{\hat{S}} - [D:S_p])(\dot{\hat{E}} - [E:S] - [S:E]) = 0
\]

\[
[F:E_p]^2 - (F_T + K_{mb} + E_T - \dot{\hat{E}})[F:E_p] + F_T(E_T - \dot{\hat{E}}) = 0
\]

(39)

In Fig. 9, we plot a sample trajectory of \(\dot{\hat{S}}\) in the bistable regime \((S_T = 95)\); we initialize the system with \(\hat{\hat{S}} = 8, \hat{\hat{S}}_p = 87\) molecules. The upper panel shows a trajectory calculated from a full Gillespie simulation and the lower panel shows a stochastic TQSSA simulation. It appears from this example that the residence time in the upper steady state is much shorter in the stochastic TQSSA calculation than in the full Gillespie calculation. To investigate the apparent discrepancy, we calculate the residence time distributions of lower and upper steady states in both full Gillespie and stochastic TQSSA simulations. We initialize the system at the lower steady state \((\hat{\hat{S}} = 8)\) and record the time \(t_0\) when \(\hat{\hat{S}}(t)\) reaches the first time its upper steady-state value \((\hat{\hat{S}}(t_0) = 58)\). Then we follow the trajectory until \(\hat{\hat{S}}(t)\) next reaches its lower steady-state value \((\hat{\hat{S}}(t_1) = 8)\). Continuing this process, we record a sequence of times \(t_0, t_1, t_2 \ldots\) when \(\hat{\hat{S}}(t)\) flips back

### TABLE 1 Parameter values for coupled GK switches

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Substrate</th>
<th>Letter,</th>
<th>(k_1)</th>
<th>(K_m)</th>
<th>(L_T)</th>
</tr>
</thead>
<tbody>
<tr>
<td>D, E</td>
<td>S</td>
<td>b, c</td>
<td>0.625</td>
<td>17.6</td>
<td>1600</td>
</tr>
<tr>
<td>E, E</td>
<td>S_p</td>
<td>d, e</td>
<td>0.0001125</td>
<td>0.816</td>
<td>160</td>
</tr>
</tbody>
</table>

The association rate constants \(k_1\) are in units molecule\(^{-1}\) min\(^{-1}\), and dissociation \(k_2\) and catalytic \(k_3\) rate constants are min\(^{-1}\). Units for Michaelis-Menten constants \(K_m\) are molecules. \(L_T\) is the total number of molecules of enzymes D, E, and F.

In the same way as before, we can obtain a master equation for the slow reactions:

\[
\dot{\hat{S}}_p = \frac{k_1^p(\hat{\hat{S}}_p)}{k_2^p(\hat{\hat{S}}_p)} \hat{\hat{S}}
\]

\[
\dot{\hat{E}}_p = \frac{k_2^p(\hat{\hat{S}}_p)}{k_2^p(\hat{\hat{S}}_p)} \hat{\hat{E}}_p
\]

(40)

with propensities based on the expected values of the numbers of enzyme-substrate complexes. We calculate the expected values of the complexes by solving the coupled algebraic equations (Eq. 39). In Fig. 8, we compare the steady-state probability distribution of \(\dot{\hat{S}}\) obtained from full Gillespie simulations of this reaction network with stochastic TQSSA simulations of Eq. 40. We calculate the steady-state probability distributions for different values of \(S_T\). In the monostable zone \((S_T < 88)\), the prediction of stochastic TQSSA is good (Fig. 8 a). As \(S_T\) increases into the bistable zone, the prediction of stochastic TQSSA is still fine (Fig. 8 b), although the relative intensities of the two peaks in the bistable zone differ from the “exact” result of full Gillespie simulations. In the upper monostable region \((S_T > 99)\), the probability distribution calculated from the stochastic TQSSA is a little shifted from the exact result (Fig. 8 c). In the deterministic case, the system settles to a particular steady state depending on initial conditions. In the stochastic case, the system jumps back and forth between the alternative stable steady states.
and forth between its upper and lower steady states. We define \((t_1 - t_0)\) as a sample residence time in the upper steady state, \((t_2 - t_1)\) as a sample residence time in the lower steady state, and so on. From a long simulation, we compute many sample residence times for both steady states. In Fig. 10, we plot cumulative distributions of these residence times. The top panel compares the cumulative probability distribution for residence times in the upper steady state from full Gillespie and stochastic TQSSA simulations (Fig. 10, a and b, respectively). The dashed lines are single exponential fits to the cumulative distribution. The bottom panel is the same for the lower steady state. Though the stochastic TQSSA calculation produces an exponential decay distribution, the mean residence times for the upper \((T_U = 42.15 \times 10^3 \text{ min})\) and lower \((T_L = 5.55 \times 10^3 \text{ min})\) steady states do not agree with the full Gillespie simulation \((T_U = 129.96 \times 10^3 \text{ min}; T_L = 20.84 \times 10^3 \text{ min})\).

Suspecting that these divergent results might be attributable to poor timescale separation (see the Supplementary Material, Data S1) in our parameter set (Table 1), we repeated the simulations for a new parameter set with large rate constants for association and dissociation of enzyme-substrate complexes, while keeping the rate constant for enzymatic conversion of substrate to product unchanged (see the Supplementary Material, Table S6, Data S1). In this case, we find that stochastic TQSSA simulations are in excellent agreement with full Gillespie simulations with respect to steady-state probability distributions (see the Supplementary Material, Fig. S4, a-c, Data S1) and mean residence times in the lower and upper steady states.

**DISCUSSION**

The stochastic TQSSA approach differs from tau-leaping and chemical Langevin approaches in that it does not simulate the fast reactions. In tau-leaping (3–6), the number of fast-reaction events that occur during each time interval is estimated using a Poisson random variable, whereas Haseltine and Rawlings (8) simulate the fast reactions using chemical Langevin equations. These methods can work for enzyme-catalyzed reactions, but tau-leaping can be slow for stiff systems and the accuracy of the chemical Langevin equation can be a problem when the number of molecules of reacting species becomes small.
TQSSA values of the fast species. Furthermore, assuming that the reactions, not species, according to timescale. His approach is using Gillespie’s stochastic simulation algorithm will accurately reproduce the noise in the reaction system while proximation (QSSA). Thus, simulating the reduced reaction obtained using the quasi-steady-state approximation. The labeled lines represent full Gillespie steady state. The labeled lines represent full Gillespie and stochastic TQSSA (b). Dashed lines are best-fitting exponential curves to the cumulative probability distributions.

FIGURE 10 Bistable switch. Plot of cumulative probability \( p_t \) of residence times \( t_R \) for \( s_T = 95 \). (Upper) Upper steady state. (Lower) Lower steady state. The labeled lines represent full Gillespie (a), and stochastic TQSSA (b). Dashed lines are best-fitting exponential curves to the cumulative probability distributions.

Other approaches rely on the separation of timescales that often occurs in enzyme-catalyzed reactions. Rao and Arkin (9) separate the species according to whether they evolve on fast or slow timescales and show how to obtain a master equation for the slow species that depends on the average values of the fast species. Furthermore, assuming that the timescale separation allows the fast species to equilibrate quickly, they show that in the Michaelis-Menten case the master equation for the slow species corresponds to the reduced reaction obtained using the quasi-steady-state approximation (QSSA). Thus, simulating the reduced reaction using Gillespie’s stochastic simulation algorithm will accurately reproduce the noise in the reaction system while avoiding the bottleneck created by the fast reactions.

Goutsias (10) takes a similar approach, but separates the reactions, not species, according to timescale. His approach is more general than Rao and Arkin in terms of how he determines the average values of the fast species, and for the Michaelis-Menten case his approach is not limited by the conditions under which QSSA is valid. In the general case, his approach requires the solution of a system of quadratic equations subject to a linear inequality constraint after each occurrence of a slow reaction.

Cao et al. (11,12) also identify fast- and slow-reaction channels, but propose a framework using the steady-state probability function for the fast species to calculate the moments of the fast species. Recognizing that this calculation will not always be feasible, they explore several approximate methods, the simplest of which involves using equilibrium values from the associated reaction-rate equations.

The TQSSA approach differs from the quasi-equilibrium assumption of Goutsias (10) and Cao (11,12) in that it allows the slow-timescale reactions to impact the steady state of the fast reactions. For the case of large timescale separation, all three approaches will give comparable results. This case is nice theoretically since the average values of the complexes decay quickly to their steady-state values after the firing of a slow reaction, and the master equation governing the slow reactions using the steady-state average propensities is a good approximation to the actual master equation that has time-varying coefficients. In cases with less timescale separation, the time-varying coefficients of the actual master equation do not settle out between firings of the slow reactions. In these cases, the TQSSA approach can still provide a good approximation, as shown in our examples.

For the Goldbeter-Koshland ultrasensitive switch, our stochastic results (means and standard deviations) for the slow variables using TQSSA are all but identical to the results obtained by simulating the full system using Gillespie’s stochastic simulation algorithm. In addition, the means for the complexes, which are not directly simulated, are very accurate in all cases, and the standard deviations are accurate except in cases where the free enzyme populations are reduced to only a few molecules. In these problem cases, the inaccuracies in the standard deviations are of the order of 1 molecule. This accuracy for variables that are not directly simulated is surprising. It is due largely to the fact that the standard deviation is composed of components from both the fast and slow reactions, and the fast (unsimulated) component is very small (see Fig. 1 b).

Furthermore, for slow variables, the TQSSA technique accurately captures the autocorrelation function of fluctuations about a steady state. For the complexes, free enzyme, and free substrate variables that are not directly simulated, the autocorrelation functions can consist of multiple exponential components, and our approach will find only the slowest components of the autocorrelation functions. This is to be expected, since all information about fast transients is lost in the TQSSA approach.

Our final example of a coupled Goldbeter-Koshland switch exhibiting bistability is a particularly sensitive test of the accuracy of the stochastic TQSSA, since small changes in stochastic trajectories near the unstable steady state can change the stable steady state that is approached. For the parameter values considered here (Table 1), there is little
timescale separation between complex formation and product formation. Even so, TQSSA produces reasonably accurate probability distributions, as seen in Fig. 8, a–c, although the residence time distribution is inaccurate (Fig. 10).

In summary, our results show that stochastic simulations based on the TQSSA can be implemented in a straightforward, algorithmic manner to efficiently calculate accurate trajectories for the slow variables in a multiple-timescale system of chemical reactions.

SUPPLEMENTARY MATERIAL
To view all of the supplemental files associated with this article, visit www.biophysj.org.

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