Problem Set 2

1 Competitive and uncompetitive inhibition (12 points)

a. Reversible enzyme inhibitors can bind enzymes reversibly, and slowing down or halting enzymatic reactions. If an inhibitor occupies the active site of an enzyme and prevents binding of the substrate to the enzyme, we call it a competitive inhibitor. In the presence of a competitive inhibitor, the Michaelis-Menten equation becomes

\[ V_0 = \frac{V_{\text{max}}S_0}{\alpha K_M + S_0} \]

What is \( \alpha \) in terms of \( K_I \) and \( I_0 \)? How does a competitive inhibitor affect the maximal turnover rate and the substrate concentration at which half that rate is reached? (Begin with a definition of total enzyme as \( [E]_{\text{tot}} = [E] + [ES] + [EI] \) and use the derivation of the Michaelis-Menten equation as a guide. Assume \( S_0, I_0 \gg [E]_{\text{tot}} \)).

\[ E + S \rightleftharpoons ES \rightarrow E + P \]

\[ E + I \rightarrow EI \]

\[ K_i \]

Problem 1a-d Image and problem courtesy of Alexander van Oudenaarden. Used with permission.

b. Unlike a competitive inhibitor, an uncompetitive inhibitor binds only to the Enzyme-Substrate (ES) complex, at a site distinct from the substrate active site. In the presence of an uncompetitive inhibitor, the Michaelis-Menten equation is altered to

\[ V_0 = \frac{V_{\text{max}}S_0}{K_M + \beta S_0} \]

What is \( \beta \) in terms of \( K'_I \) and \( I_0 \)? How does a competitive inhibitor affect the maximal turnover rate and the substrate concentration at which half that rate is reached?

\[ E + S \rightleftharpoons ES \rightarrow E + P \]

\[ E + I \rightarrow ESI \]

\[ K'_i \]

\[ ESI \]
c. A \textit{mixed} inhibitor binds at a site distinct from the substrate active site, but it binds to either \( ES \) or the enzyme by itself. Assume detailed balance for the reversible reactions. How is the Michaelis-Menten equation modified in the presence of a mixed inhibitor?

\begin{center}
\includegraphics[width=0.5\textwidth]{reaction_diagram}
\end{center}

d. From an inspection of the curves below, determine the nature of the inhibition by acetazolamide on the carbonic anhydrase reaction, \( H_2CO_3 \rightleftharpoons H_2O + CO_2 \).

\begin{center}
\includegraphics[width=0.5\textwidth]{inhibition_curves}
\end{center}

2 \textbf{Stability analysis of dynamical systems\textsuperscript{1} (10 points)}

a. For each of the following 1D dynamical systems, identify the fixed points and use a graphical argument \( (f(x) = \frac{dx}{dt} \text{ vs. } x) \) to determine the stability. In addition, plot the potential potential function \( V(x) \) \( (f(x) = -\frac{dV}{dx}) \). \( r \) can be positive, zero or negative. Discuss all three scenarios.

1. \( \frac{dx}{dt} = rx - x^2 \)
2. \( \frac{dx}{dt} = rx - x^3 \)

b. (Strogatz 6.4) Consider the following 2D dynamical systems (two species competing for the same limited resource, for example, “rabbits” vs. “sheep”), where \( x, y \geq 0 \). Find the fixed points, investigate their stability, draw the nullclines, and sketch plausible phase portraits. Indicate the basins of attraction of any stable fixed points.

1. \( \frac{dx}{dt} = x(3 - x - y), \quad \frac{dy}{dt} = y(2 - x - y) \)
2. \( \frac{dx}{dt} = x(3 - 2x - y), \quad \frac{dy}{dt} = y(2 - x - y) \)

\textsuperscript{1}Chapter 2 & Chapter 6, “Nonlinear dynamics and chaos” by Steven Strogatz.
3 Positive feedback and bistability (12 points)

a. Suppose the protein $X$ is a transcriptional activator, and a promoter $D$ which is activated by the binding of $X$. If the downstream gene happens to code for $X$ itself, the resulting positive feedback can lead to bistability. We will consider an scenario such as that described in problem 3 of the first problem set in which there are two binding sites and binding is strongly cooperative.

\[ \nu_1 \] be the rate of expression from DNA bound by two $X$ molecules ($DX_AX_B$), and $\nu_0 < \nu_1$ the rate of expression from free DNA ($D$); note that these rates of expression are assumed to have units of concentration per unit time. Use the results of problems 2 and 3 of the first problem set to show for an organism that grows at rate $\gamma$, where the degradation of $X$ is negligible compared to dilution, the time-evolution of the concentration $x \equiv [X]$ may be written in the form:

\[
\frac{dx}{dt} = \frac{\nu_0 + \nu_1 K_1 K_2 x^2}{1 + K_1 K_2 x^2} - \gamma x
\]

where $K_i (i = 1, 2)$ denote association constants. (Note that in the first problem set $K$s stand for dissociation constants)

1. Steady state solutions occur at those values of $x$ at which the rate of generation, $f(x) = \frac{\nu_0 + \nu_1 K_1 K_2 x^2}{1 + K_1 K_2 x^2}$, and degradation, $g(x) = \gamma x$, are equal. How many steady states can be in this system? Sketch 5 qualitatively different positions of the function $f(x)$ relative to the function $g(x)$. Label those values of $x$ for which $f(x) > g(x)$ with a rightward arrow, and those for which $f(x) < g(x)$ with a leftward arrow. Which solutions are stable?

b. For two genes, positive feedback can arise from mutual inhibition. Consider a toggle switch modeled by the equations (Box 1, T.S.Gardner et al, Nature 2000)

\[
\frac{du}{dt} = \frac{\alpha}{1 + v^2} - u
\]

\[
\frac{dv}{dt} = \frac{\alpha}{1 + u^2} - v
\]

Which values of $\alpha$ give the system bistability? Justify your answer.

4 Kinetic Proofreading

Many cellular processes, such as transcription of a gene, translation of mRNA, or even the recognition of an antibody by a T-cell require high fidelity. During translation, for example, the ribosomes
are attempting to perform their enzymatic function on a particular substrate, but many other substrates in the cell look similar so it is difficult to perform the enzymatic function accurately. Yet, translation errors occur at typical rates as low as $\sim 10^{-4}$. In this problem, we will look at a mechanism that is employed to achieve such high accuracy. This was first suggested by J. J. Hopfield (see ref at end).

Let $c$ be a recognition site, to which two substrates can bind, $C$, the correct and $D$, the wrong one. The correct complex then produces the expected product:

$$
C + c \xleftrightarrow{k_c} Ce \rightarrow \text{correct product} \quad K_C = k_c/k_c
$$

$$
D + c \xleftrightarrow{k_d} De \rightarrow \text{error product} \quad K_D = k_d/k_d \quad [1]
$$

The rate of product formation from the complex ($W$) is determined by the strength of the covalent bonds between, for example, two amino acids, and is approximately the same for both the products.

Fidelity is quantified in terms of the error fraction $f$, which is the ratio of rate of production of wrong product to correct product.

a. Assuming that the substrate concentrations in the cell are maintained at a constant (and equal) value for $C$, $D$, what is the error fraction? In the limit of $W << k_c$, the error fraction is minimized. What is this minimum value, $f_0$? (Note that being in the same medium, the concentration of the recognition sites, $c$, is the same for both reactions)

Now consider adding an additional intermediate state of the complex in the above reaction:

$$
C + c \xleftrightarrow{k_c} Cc \xrightarrow{m'} Ce^* \xrightarrow{w} \text{product} \quad m = 0, l_c = 0, l_c' = 0
$$

b. What is the error rate in this process? Detailed balance puts a restriction on the values that $m, m'$ can take. What is this constraint? It turns out that under this constraint, $f \geq f_0$, so this might not seem like a good method to increase fidelity.

Next assume steps 2, 3 in the above reaction are one way steps, i.e. $m = 0, l_c = 0$. The reaction steps can be made strongly directed like this by coupling them with other strongly driven reactions, such as ATP hydrolysis, so that they are nearly one way.

c. If $m' < k_c$, so that you can assume reaction 1 is in quasi-equilibrium by itself, and with $m = 0, l_c' = 0$, show that the error fraction now is $\approx f_0^2$. (The ratio of off rates for the original and modified complex is the same, since it involves the same substrate falling off.)

Adding multiple steps like this can greatly increase the specificity of the reaction. With $n$ one way steps, the error fraction can be lowered to $f_0^n$. However, at each of these steps, the cell has to consume energy (for ATP hydrolysis), so this is a costly function. Another mechanism involves introducing a delay between complex formation and product precipitation. Since the dissociation rate of the wrong substrate is higher, this delay allows more time for the wrong substrate to fall off. See Uri Alon proofreadign chapter for details.

Ref: Kinetic Proofreading: A New Mechanism for Reducing Errors in Biosynthetic Processes Requiring High Specificity, J. J. Hopfield, PNAS 71(10), 4135-4139