

## 1 Problem 1

To see the effect of parameters on the steady state mRNA model, the original values were decreased and increased by 20%. Table One reports the "increasing" effects as the "decreasing" effect was more or less equal and opposite. Problem Set 1 includes all sources and rationales for original values. Note that  $k_A$  was assumed to be zero so changing this value wouldn't have an effect although increasing abortion would probably lower mRNA concentrations. Also note that increasing the level of change to a 50% difference didn't make the plots that didn't change start to change meaning they are not sensitive to these parameters. Finally, gene length  $L$  was not recorded because it has the predicted inverse effect of  $k_{Ej}$  due to the inverse proportion between them. Appendix A has every plot produced by CHEME5440HW2.m that motivates the descriptions in Table One.

**Table One: Effect of Key Parameters on mRNA Profile**

<i>Parameter</i>	<i>Symbol</i>	<i>Effect when Increased</i>
Elongation Rate	$k_E$	Increases mRNA concentration at all levels although more so once induced at high inducer concentration
Degradation rate	$k_X^d$	Decreases mRNA concentration at all levels although more so once induced at higher inducer concentration
Gene Concentration	$G$	No effect
Rate of Open Complex Formation	$k_I$	Increases mRNA concentration at all levels although almost all once induced at higher inducer concentrations
Rate of Closed Complex Formation	$k_+$	No effect
Rate of Closed Complex Dissociation	$k_-$	No effect
Saturation Constant	$K_X$	No effect
Specific Growth Rate	$\mu$	Only decreases mRNA concentration once induced or higher inducer concentration
Cooperativity Constant	$n$	Only decreases mRNA at the regions not at kinetic limit or lower to middle inducer concentrations
RNA Polymerase Concentration	$R_{XT}$	Increases mRNA concentration at all levels although more so once induced at higher inducer concentration
Time Constant	$\tau$	Decreases mRNA concentration at all levels although more so once induced at higher inducer concentrations
Weight Without Inducer	$W_1$	Only increases mRNA at the regions not induced or low inducer concentrations
Weight With Inducer	$W_2$	Only increases mRNA in the middle regions when the step is occurring but before kinetic limit
Affinity Constant	$K$	Only decreases mRNA in the middle regions when the step is occurring but before kinetic limit
Hill Function	$f_I$	Only decreases mRNA at the regions not at kinetic limit or lower to middle inducer concentrations

## 2 Problem 2 Part A

Starting with transcription, we can write an mRNA balance where the dilution term  $B^{-1}B'$  is already substituted as specific growth rate:

$$\frac{dm_1}{dt} = r_{X1}u_{X1} - k_{X1}^d m_1 - \mu m_1$$

$$\frac{dm_2}{dt} = r_{X2}u_{X2} - k_{X2}^d m_2 - \mu m_2$$

$$\frac{dm_3}{dt} = r_{X3}u_{X3} - k_{X3}^d m_3 - \mu m_3$$

Where

$$r_{Xj} = k_{EXj} R_{XT} \frac{G_j}{K_{Xj} \tau_{Xj} + (\tau_{Xj} + 1) G_j}$$

And

$$u_{X1} = \frac{W_{RT1} + W_I f_I}{1 + W_{RT1} + W_I f_I}$$

$$u_{X2} = \frac{W_{RT2} + W_{12} f_{12} + W_{32} f_{32}}{1 + W_{RT2} + W_{12} f_{12} + W_{32} f_{32}}$$

$$u_{X3} = \frac{W_{RT3} + W_{13} f_{13} + W_{23} f_{23}}{1 + W_{RT3} + W_{13} f_{13} + W_{23} f_{23}}$$

Where each  $f_{ij}$  is described as:

$$f_I = \frac{I^n}{K^n + I^n}$$

$$f_{12} = \frac{p_1^n}{K^n + p_1^n}$$

$$f_{13} = \frac{p_1^n}{K^n + p_1^n}$$

$$f_{23} = \frac{p_2^n}{K^n + p_2^n}$$

$$f_{32} = \frac{p_3^n}{K^n + p_3^n}$$

Now translation of each gene can be described via analogy as:

$$\frac{dp_1}{dt} = r_{L1}u_{L1} - k_{L1}^d p_1 - \mu p_1$$

$$\frac{dp_2}{dt} = r_{L2}u_{L2} - k_{L2}^d p_2 - \mu p_2$$

$$\frac{dp_3}{dt} = r_{L3}u_{L3} - k_{L3}^d p_3 - \mu p_3$$

Where:

$$r_{Lj} = k_{ELj} R_{LT} \frac{m_j}{K_{Lj} \tau_{Lj} + (\tau_{Lj} + 1) m_j}$$

And the control terms is at 1 because the system is assumed to be operating at the kinetic limit:

$$u_{Lj} = W_j = 1$$

Now in order to write this in matrix form we define  $\mathbf{x}$  as the 6x1 mRNA/protein vector,  $\mathbf{A}$  as the consumption matrix,  $\mathbf{S}$  as the 6x6 stoichiometric matrix (here, read identity), and  $\mathbf{r}$  as the 6x1 overall reaction rate vector:

$$\frac{d\mathbf{x}}{dt} = \mathbf{A}\mathbf{x} + \mathbf{S}\mathbf{r}$$

$$\frac{d}{dt} \begin{pmatrix} m_1 \\ m_2 \\ m_3 \\ p_1 \\ p_2 \\ p_3 \end{pmatrix} = \begin{pmatrix} -(\mu + k_{X1}^d) & 0 & 0 & 0 & 0 & 0 \\ 0 & -(\mu + k_{X2}^d) & 0 & 0 & 0 & 0 \\ 0 & 0 & -(\mu + k_{X3}^d) & 0 & 0 & 0 \\ 0 & 0 & 0 & -(\mu + k_{L1}^d) & 0 & 0 \\ 0 & 0 & 0 & 0 & -(\mu + k_{L2}^d) & 0 \\ 0 & 0 & 0 & 0 & 0 & -(\mu + k_{L3}^d) \end{pmatrix} \begin{pmatrix} m_1 \\ m_2 \\ m_3 \\ p_1 \\ p_2 \\ p_3 \end{pmatrix} + \begin{pmatrix} 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} r_{X1}u_{X1} \\ r_{X2}u_{X2} \\ r_{X3}u_{X3} \\ r_{L1}u_{L1} \\ r_{L2}u_{L2} \\ r_{L3}u_{L3} \end{pmatrix}$$

### 3 Problem 2 Part B

In order to solve this set of ODEs, we have to define all the parameters. Table Two defines them

**Table Two: Relevant Parameters**

Parameter Name	Symbol	Value	Units	Source
Gene concentration	$G_{1,2,3}$	200	plasmids/cell	Given
Gene concentration	$G_{1,2,3}$	0.972	$\mu\text{mol}/gDW$	See Note 1 Below
Gene length	$L_{1,2,3}$	1200,2400,600	nucleotides	Given
<i>E.coli</i> TX Elongation Rate	$e_X$	42	nt/s	Gotta[1]
TX Rate of Elongation	$k_{EX1,2,3}$	0.0350,0.01750.07	$s^{-1}$	See Note 2 Below
RNAP Concentration	$R_{XT}$	8000	molecules/cell	Bremer[2]
RNAP Concentration	$R_{XT}$	38.88	$\mu\text{mol}/gDW$	See Note 1 Below
TX Rate of Open Complex Formation	$k_I$	1/42	$s^{-1}$	McClure[3]
TX Rate of Closed Complex Formation	$k_+$	5	$\mu M^{-1}s^{-1}$	See Note 3 Below
TX Rate of Close Complex Dissociation	$k_-$	0.1	$s^{-1}$	See Note 3 Below
TX Rate of Abortive Initiation	$k_A$	0	$s^{-1}$	$k_A \ll k_{I} \text{ or } k_E$
Ribosome Concentration	$R_{LT}$	20,100	ribosomes/cell	Nilsson[4]
Ribosome Concentration	$R_{LT}$	97.68	$\mu\text{mol}/gDW$	See Note 1 Below
<i>E. coli</i> Translation Elongation Rate	$e_L$	14.5	aa/s	Dalbrow[5]
TX Rate of Elongation	$k_{EL1,2,3}$	0.0362,0.0181,0.0725	$s^{-1}$	See Note 2 Below
TL Rate of Open Complex Formation	$k_{LI}$	1/15	$s^{-1}$	1/Time from Siwiak[6]
TL Rate of Closed Complex Formation	$k_{L+}$	5	$\mu M^{-1}s^{-1}$	See Note 3 Below
TL Rate of Close Complex Dissociation	$k_{L-}$	0.1	$s^{-1}$	See Note 3 Below
TL Rate of Abortive Initiation	$k_{LA}$	0	$s^{-1}$	$k_{LA} \ll k_{LI} \text{ or } k_{EL}$
Bacterial Protein Half-Life	$t_{0.5}^p$	72000	s	Koch[7]
mRNA half-life	$t_{0.5}^m$	120	s	Li[8]
Doubling Time of <i>E.coli</i>	$t_d$	1800	s	Given
Weight of Inducer on 1	$W_I$	100	Unitless	See Note 4 Below
Weight of 1 on 2	$W_{12}$	100	Unitless	See Note 4 Below
Weight of 1 on 3	$W_{13}$	100	Unitless	See Note 4 Below
Weight of 2 on 3	$W_{23}$	0 or 10	Unitless	See Note 4 Below
Weight of 3 on 2	$W_{32}$	10	Unitless	See Note 4 Below
Basal Weight of 1	$W_{RT1}$	0.00001	Unitless	See Note 4 Below
Basal Weight of 2	$W_{RT2}$	0.00001	Unitless	See Note 4 Below
Basal Weight of 2	$W_{RT3}$	0.00001	Unitless	See Note 4 Below
Affinity Constant	$K$	0.3	$mM$	See Note 4 Below
Cooperativity Constant	$n$	2	Unitless	See Note 4 Below

**Note 1**

Molecules per cell were converted to  $\mu\text{mol}/g\text{DW}$  Avogadro's Number, 30% dry weight given, the volume of a cell as  $6.7\text{e-}10\text{L}/\text{cell}$  (Wang[9]), and the weight of the wet cell at  $1.7g/L$  (Glazyrina[10]).

**Note 2**

$$k_{Ej} = \langle k_E \rangle \frac{L}{L_j}$$

Where  $\langle k_E \rangle$  is the average elongation rate and  $L, L_j$  refer to the characteristic gene length (average here) and the current gene length (1200, 2400, 600 nt or 1/3 of that for protein). This simplifies to the elongation rate  $e_X$  (42 nt/s) divided by the gene length for example for transcription:

$$k_{EX1} = e_X / L_1 = 42 / 1200 = 0.035 s^{-1}$$

This same method is used for protein  $k_{ELj}$  from  $e_L$ .

**Note 3**

The values of  $k_+$  and  $k_-$  were derived from McClure's[3] formula and value for the slope of a tau plot in *E.Coli* D promoters:

$$McClureSlope = 1.04 \mu M * s = \frac{k_- + k_I}{k_+ k_I}$$

$$1.04 k_+ (1/42) = k_- + 1/42$$

We can arbitrarily assign  $k_+ = 5 \mu M^{-1} s^{-1}$  which leads to  $k_- = 0.1 s^{-1}$ . This is valid because when these values are used later in calculating  $K_{Xj}$ , the ratio of  $k_-/k_+$  or some dissociation constant  $k_D$  is restored. This is best described below:

$$McClureSlope = \frac{k_- + k_I}{k_+ k_I}$$

$$K_{Xj} = \frac{k_- + k_I}{k_+}$$

Thus:

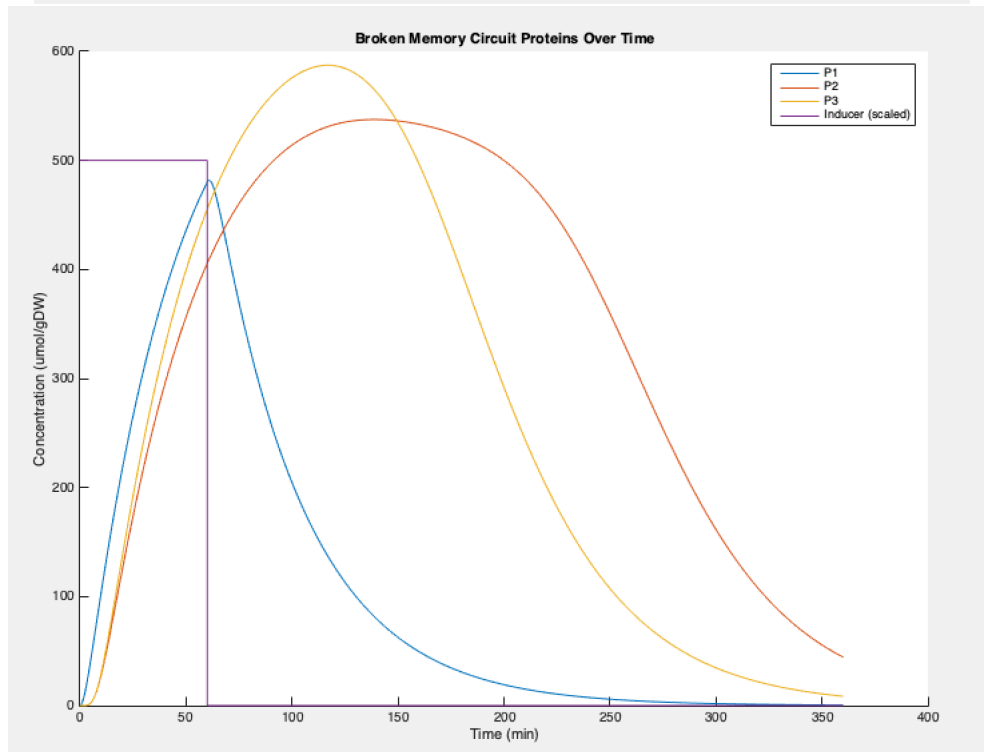
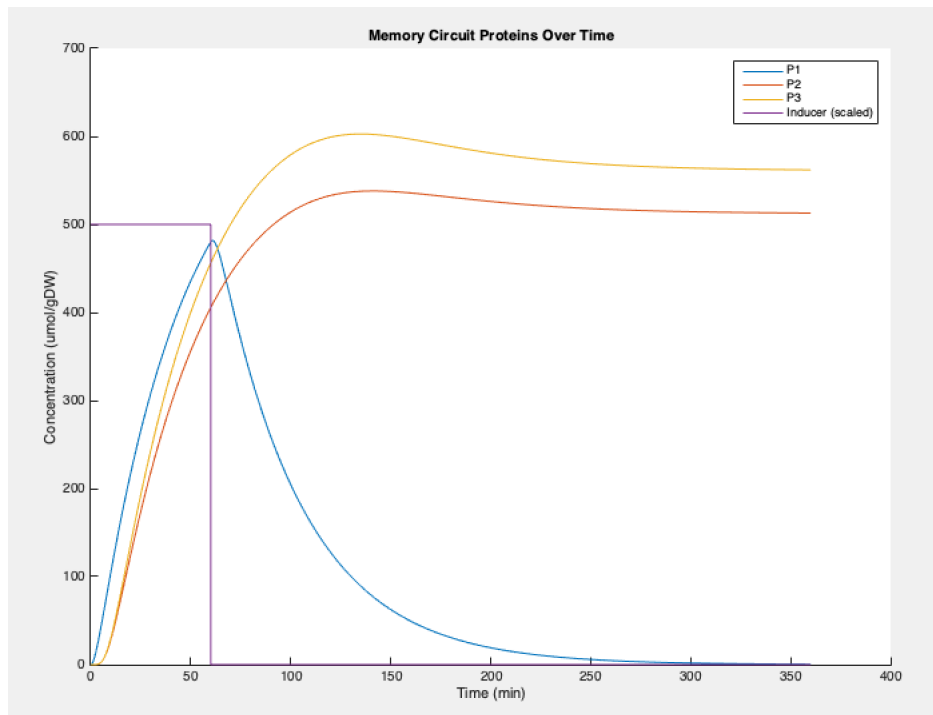
$$K_{Xj} = McClureSlope * k_I$$

Plugging into either equation for  $K_{Xj}$  returns the same value. It also happens that  $K_D$  or  $k_-/k_+$  for *E.Coli* with the lac promoter is  $550 nM$  (Bintu[11]) which is only one order of magnitude off our arbitrary assignment for transcription ( $20 nM$ ). The values for binding between a gene and RNAP and the values for a ribosome binding to mRNA were assumed to be the same. This is because the values are not easily measurable and have not been analyzed in McClure's fashion for translation. In other words,  $k_- = k_{L-}$  and  $k_+ = k_{L+}$ . It is reasonable to assume they are on the same order of magnitude as some sort of enzyme is binding a large polymer substrate.

**Note 4**

The weights, cooperativity constants, and saturation constants mostly follow the trends or values given in Problem Set 1. They are on similar orders of magnitude. The basal rates were assumed to approach zero which is reasonable to say as the inducers should have a larger effect.  $n$  was raised to 2 to increase the effect of the Hill function step and produce more descriptive plots. The effect of proteins 2 and 3 on each other were decreased below protein 1's effect so that the readjustment on steady state could be observed as protein 1 dies off in a more timely manner (under 360 minutes). It should be noted that  $W_{23}$  is changed to be 0 when the circuit is broken.

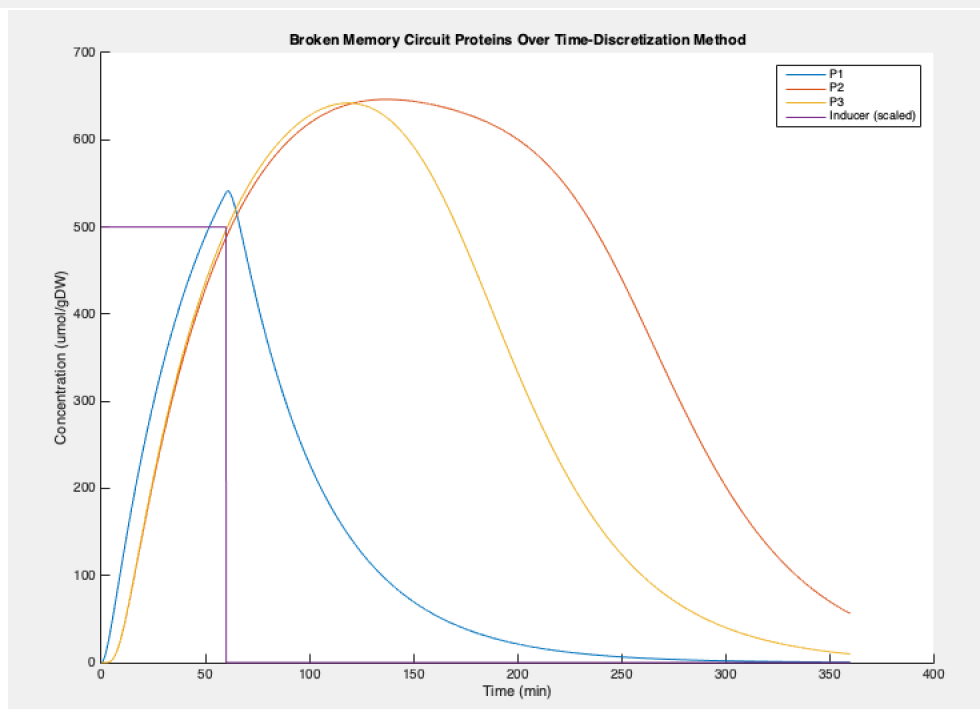
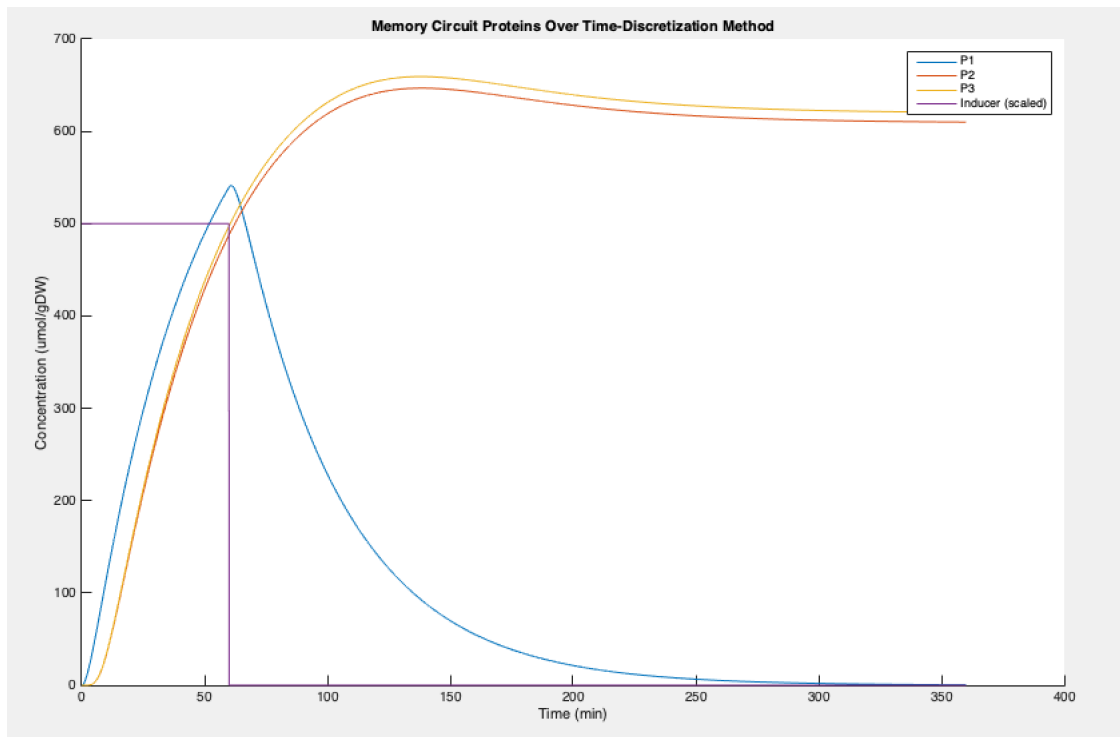
**The solutions** for a memory circuit and broken circuit are plotted as follows. The broken circuit simply removes the weight or effect that protein 2 has on the translation of protein 3 ( $W_{23} = 0$ ). Note that the inducer is scaled. HW2Run.m plots these values using ode45 calling function CHEME5440HW2B.m. The values for  $W_{23}$  were changed manually for each plot. Characteristic memory and steady state adjustments are shown

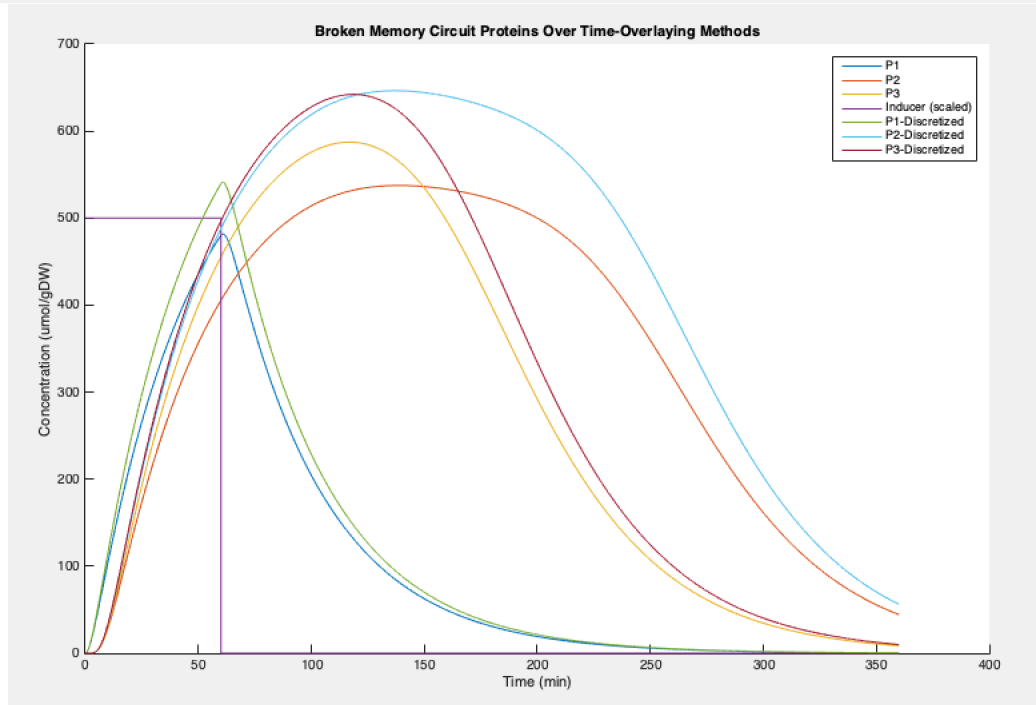
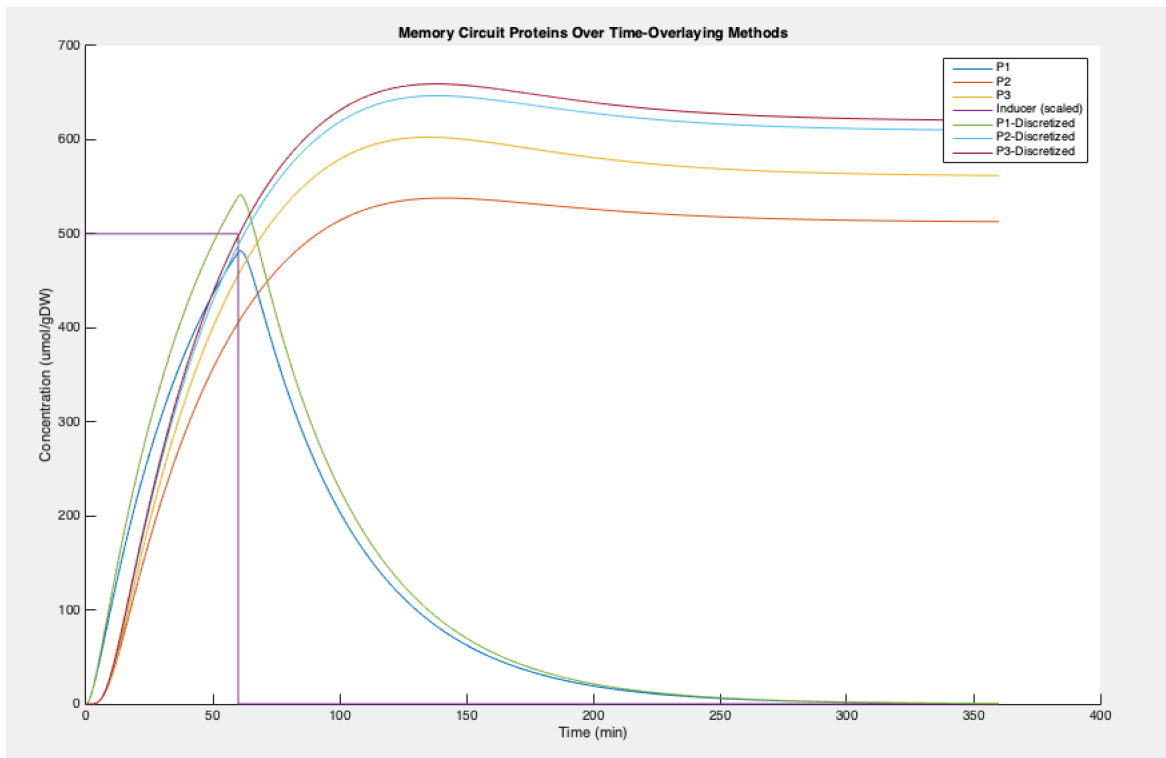


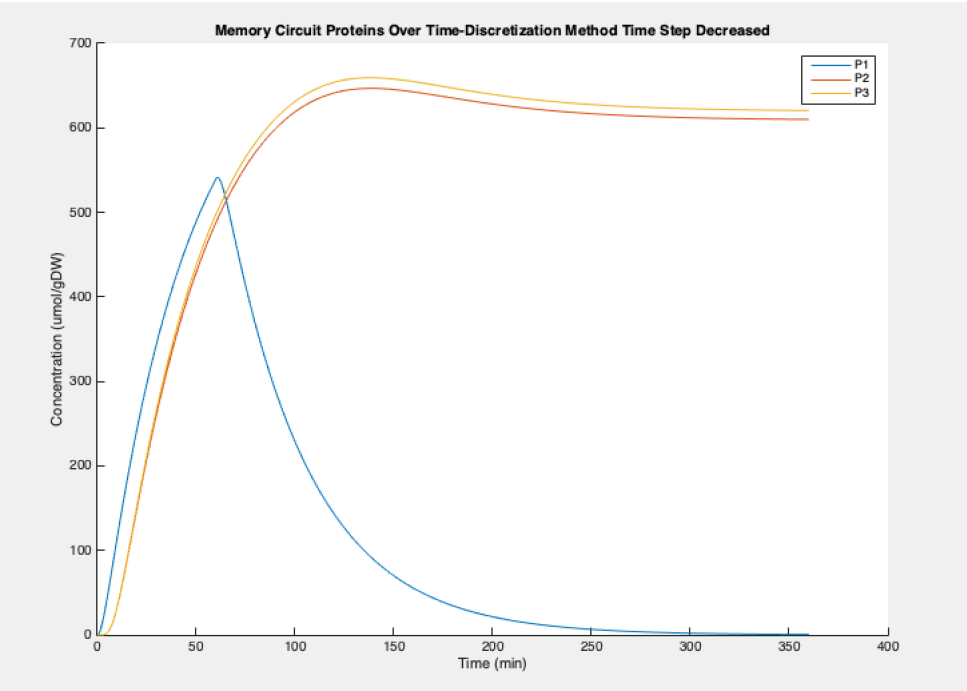
## 4 Problem 2 Part C

Using the given discretization, CHEME5440HW2C.m plots the protein profile over time. It calls the function `f.m` that simply calculates the Hill Functions ( $f_I$ ). The plots for the discretization method, discretization method overlayed with the numerical solution, and discretization method with decreased time step (0.01 min) are given below. It is notable that the discretization gives similar resolution in the kinetics where the values do not jump

drastically and follow the same patterns as the numerical ODE solver. The discretization magnitudes are slightly higher in general, which probably has to do with some effect being less strong in the discretization (such as dilution or degradation). The discretization method also has protein 3 drop below protein 2 earlier in the broken circuit. Note that decreasing the time step does not change the discretization solution.

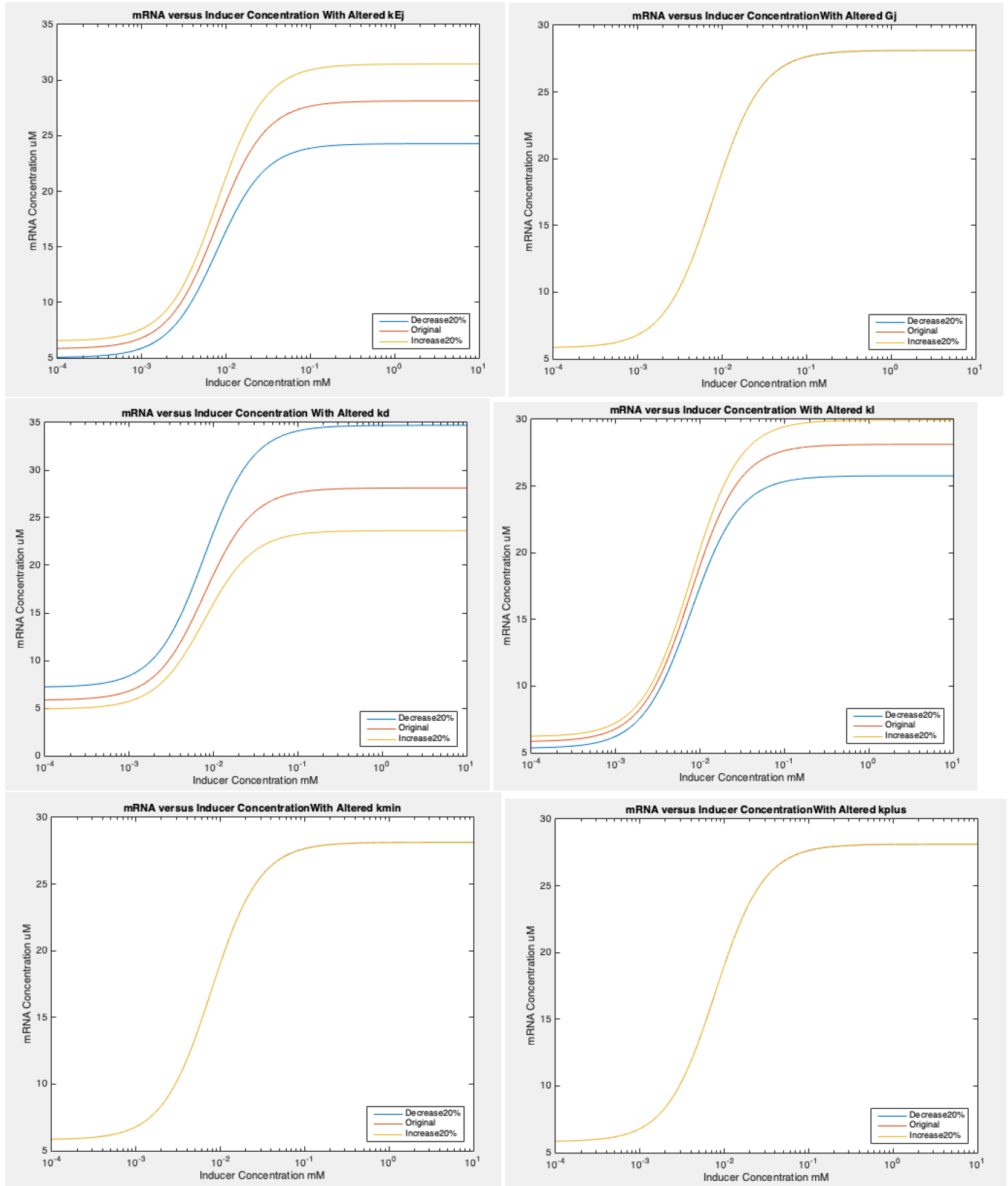


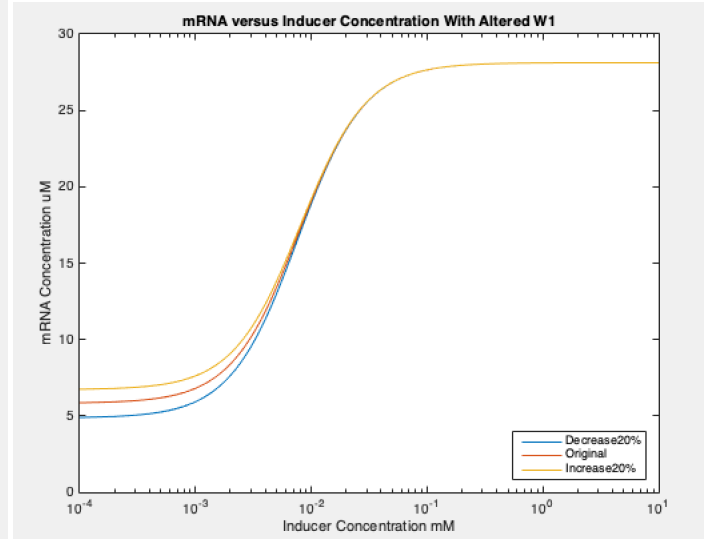
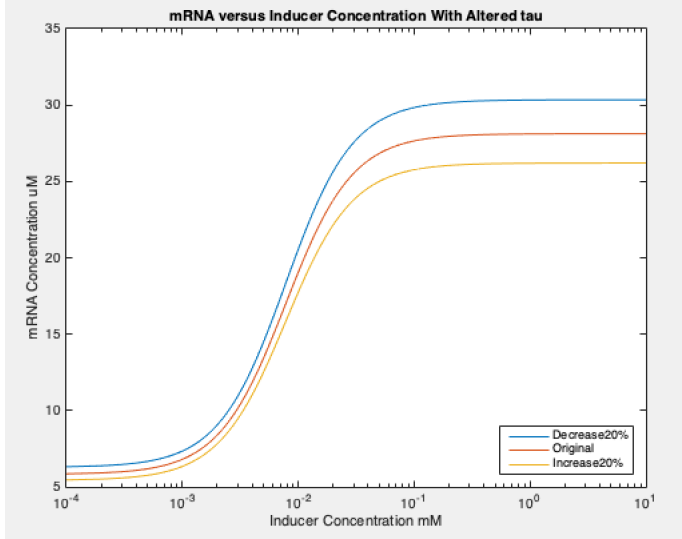
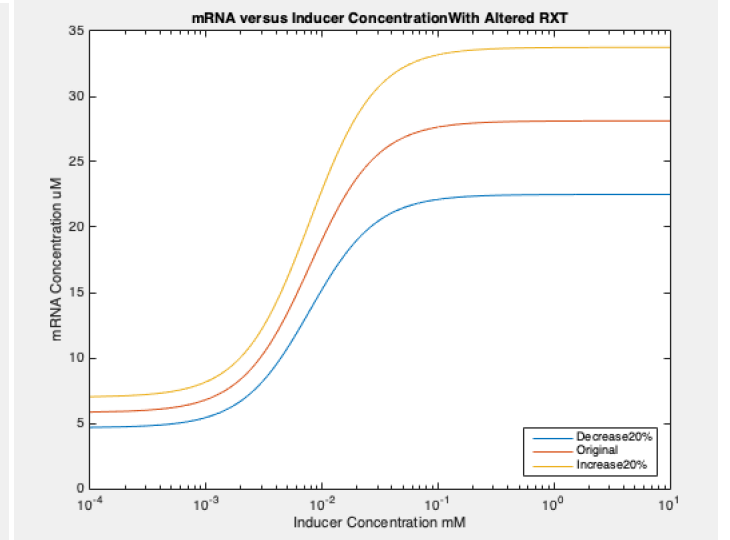
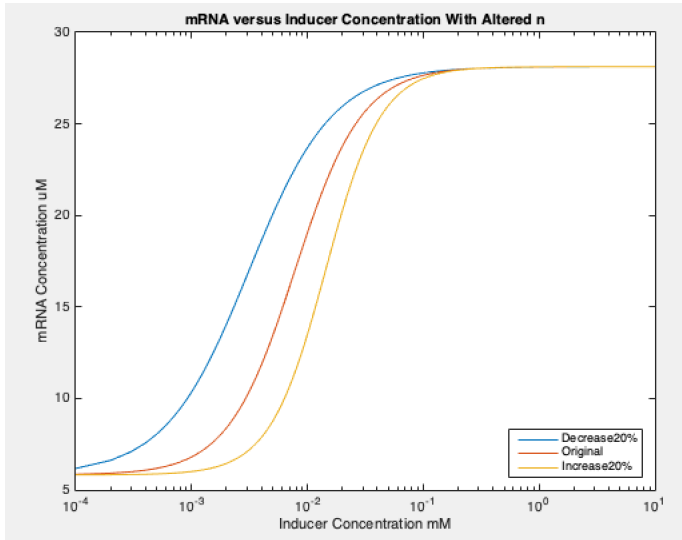
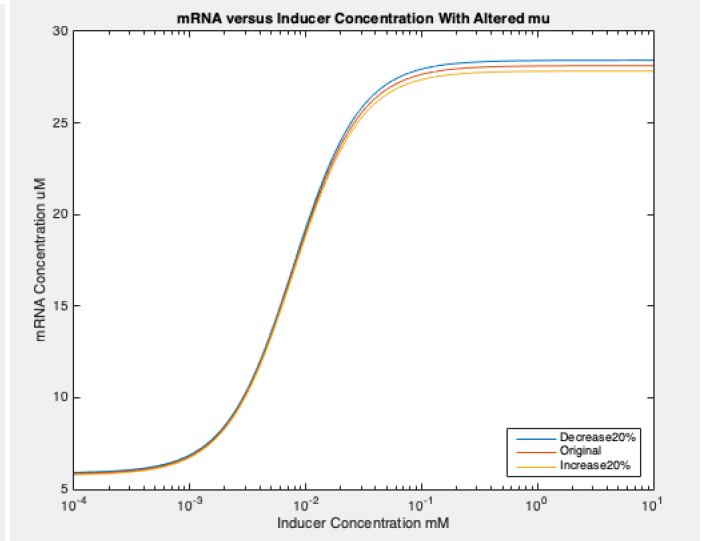
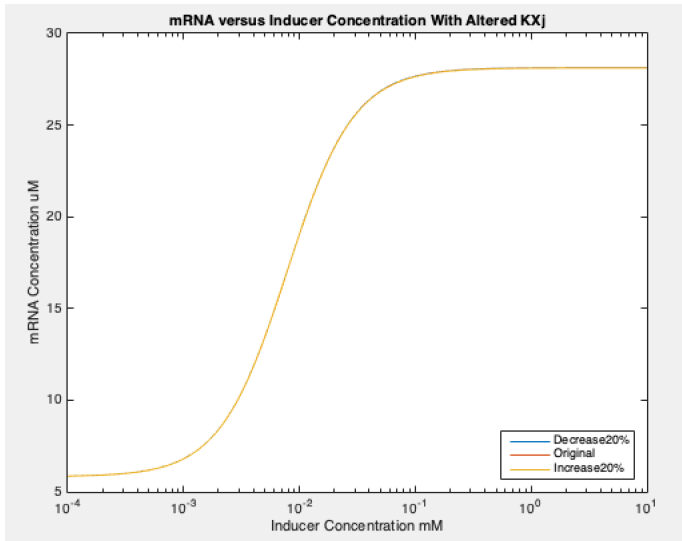


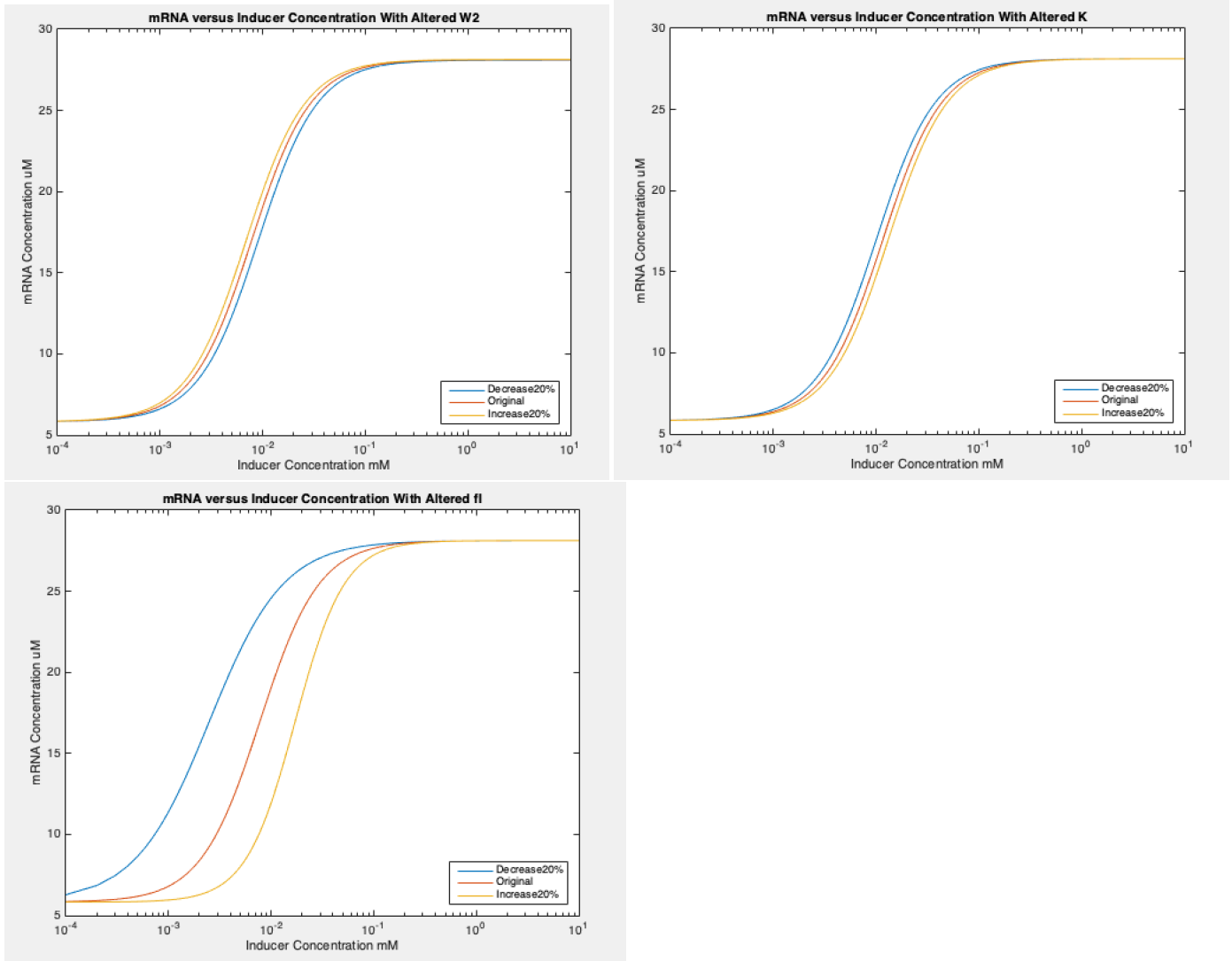




## 5 Appendix A







## References

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