## Mini-Project in Mathematical and Computational Modeling

École Polytechnique Fédérale de Lausanne, Switzerland

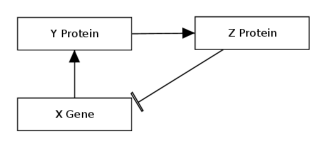
Florian + Dariush

## Introduction

Introduction to the article goes here Introduction to the article goes here

The Model

## Part A - One-Cell Model



(a) One-Cell Model

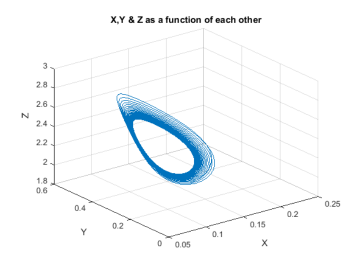
The gene mRNA X codes for protein Y which, in turn, activates transcriptional inhibitor Z. The resulting model behaves as a three-variable oscillator.

$$\begin{split} \frac{\delta X}{\delta t} &= v_1 \frac{K_1^n}{K_1^n + Z^n} - v_2 \frac{X}{K_2 + X} \\ \frac{\delta Y}{\delta t} &= k_3 X - v_4 \frac{Y}{K_4 + Y} \\ \frac{\delta Z}{\delta t} &= k_5 Y - v_6 \frac{Z}{K_6 + Z} \end{split}$$

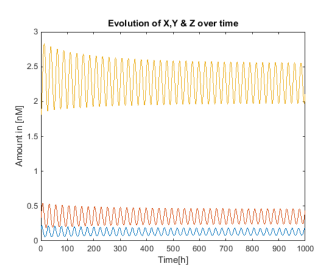
translation rate of X $v_1$ degradation rate of X $v_2$ degradation rate of Y $v_4$ degradation rate of Z $v_6$ transcription rate of Y $k_3$ transcription rate of Z $k_5$ 

Michaelis constant of X $K_1$ Michaelis constant of Y $K_4$ 

Michaelis constant of Z



(a) Trajectories The limit cycle is reached as the variations of X(t), Y(t) and Z(t) become fixed: The trajectories converge, non-lineary (the distance between similar trajectories aren't regular) towards an hundred hours. The signal are not in phase but have the same, ellipse (where the blue stripes accumulate)



(b) Frequency spectrum The amplitude of the three variations stabilize after a few regular, frequencies.

Figure 3:

Trajectories of X(t), Y(t) and Z(t) with initial conditions:  $X_0=0.16,\,Y_0=0.33,\,Z_0=1.8$  [nM] We observe on both graphs that Z(t) has the bigger amplitude of variation whereas X(t) and Y(t) have small amplitudes. Additionally, the convergence towards a single loop in (a) indicate that the frequencies of the signals are equal; this is illustrated as well in (b)

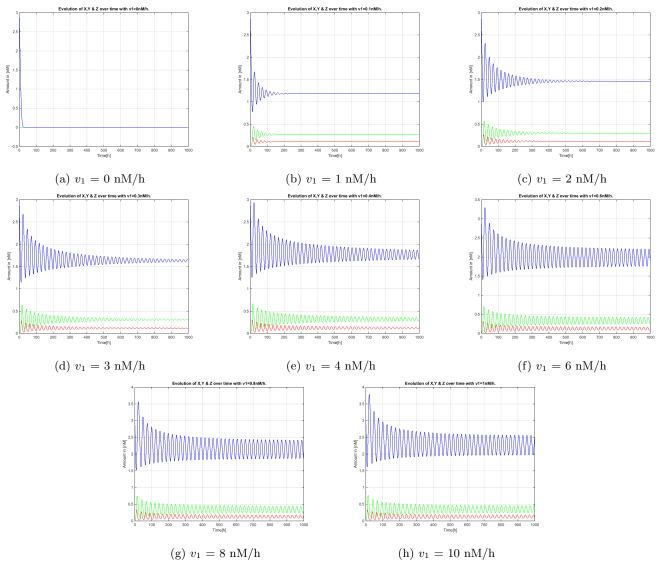


Figure 4: X(t), Y(t) and Z(t) with initial conditions  $X_0 = 0.16$ ,  $Y_0 = 0.33$ ,  $Z_0 = 1.8$  [nM] The first signal to fade is Y(t) and its oscillatory stability predicts stability of the system. We also observe that the signals converge towards null or the limit cycle in a non-linear fashion. At the opposite, it is rather difficult to predict the threshold value of  $v_1$  using those plots?

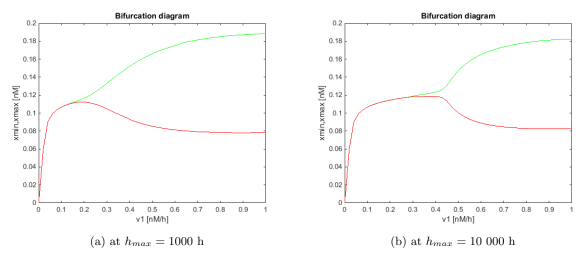


Figure 5: Bifurcation Diagram:  $X_{min}$  and  $X_{max}$  plotted at time intervals [9/10; 1] of  $h_{max}$ A limit cycle might be reached when  $X_{min} \neq X_{max}$ . However, the system needs to be run for enough time for the cycle to be reached, as the (a) suggests. (b) illustrates the non-linear convergence of the system; also the threshold for  $v_1$  seems to be around 4.5

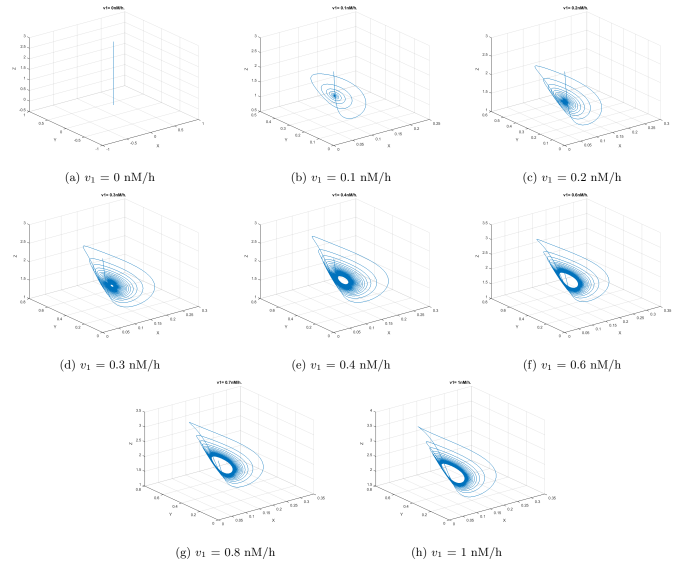


Figure 6: Trajectories when varying  $v_1$  with initial conditions  $X_0 = 0.16$ ,  $Y_0 = 0.33$ ,  $Z_0 = 1.8$  [nM]  $v_1$  has to reach a certain value for X(t) to be able to compensate its inhibition by Z(t) and therefore for the system to reach a limit cycle. We observe that this value is slightly greater than 4nM/h, as the trajectories still converge to null in (e); there is an 'eye', even though it is smaller than in (f) and (g), since the timescale is not big enough to let the system dissipate completely.

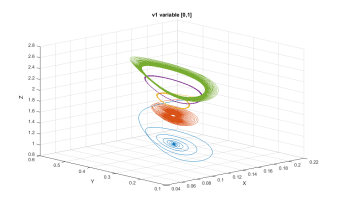
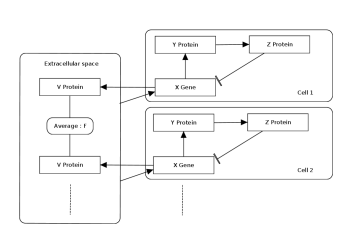


Figure 7: Superimposed trajectories at late timepoints with initial conditions  $X_0 = 0.16$ ,  $Y_0 = 0.33$ ,  $Z_0 = 1.8$  [nM] and  $v_1 = 0.1/0.3/0.5/0.7/0.9$  nM/h. We observe here that Z(t) tends to reach greater concentration stability with increasing  $v_1$ .

## Part B - Multiple Cells Model

 $v_1$ 



(a) Multiple Cells Model

The gene X codes for protein Y which, in turn, activates transcriptional inhibitor Z. In addition, gene X activates a positive feedback loop through the mean concentration of extracellular protein V

$$\begin{split} \frac{\delta X}{\delta t} &= v_1 \frac{K_1^n}{K_1^n + Z^n} - v_2 \frac{X}{K_2 + X} + v_c \frac{KF}{K_c + KF} \\ \frac{\delta Y}{\delta t} &= k_3 X - v_4 \frac{Y}{K_4 + Y} \\ \frac{\delta Z}{\delta t} &= k_5 Y - v_6 \frac{Z}{K_6 + Z} \\ \frac{\delta V_i}{\delta t} &= k_7 X_i - v_8 \frac{V_i}{K_8 + V_i} \\ \end{split}$$
 where  $F = \frac{1}{N} \sum_{i=1}^N V_i$ 

translation rate of Xdegradation rate of X $v_2$ degradation rate of Y $v_4$ degradation rate of Z $v_6$ degradation rate of V $v_8$  $k_3$ transcription rate of Y $k_5$ transcription rate of Z $k_7$ transcription rate of V

transcription rate of X $k_1$  $K_1$ Michaelis constant of X $K_4$ Michaelis constant of Y $K_6$ Michaelis constant of Z $K_8$ Michaelis constant of V $K_c$ Michaelis constant of X by F $v_c$ Activation rate of X by FKCoupling Constant

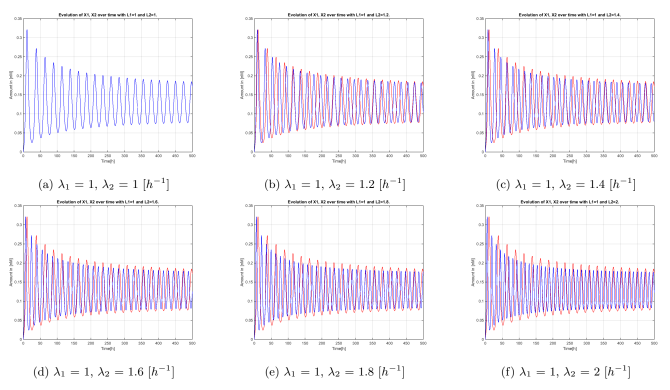


Figure 10:  $X_1(t)$  and  $X_2(t)$  trajectories in a two-cells Model with K=0 ( $\leftrightarrow$  no coupling) Figure (a) has both signals perfectly aligned. We observe no synchronisation, as excepted.

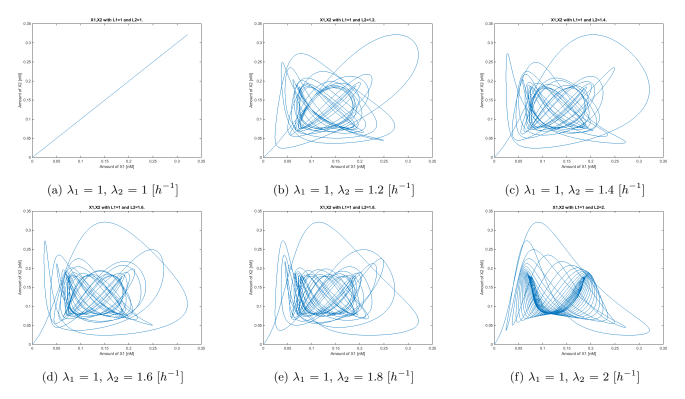


Figure 11:  $X_1$  and  $X_2$  trajectories with varying  $\lambda_i$  in a two-cells Model with K=0Figure (a), the control, makes perfect sense since the two cells have the same period, hence the exact same signal. With unequal periods, the limit cycles of both cells aren't in phase and form these '8' patterns. The fluctuations at the beginning of trajectories come from the inner adjustment of the cells (see Figure 6). The box that appears represent the variation of  $X_1$  and  $X_2$  and therefore directly gives us their minimal and maximal values. (f) gives us weird shit ?what can we say bout (f) ?

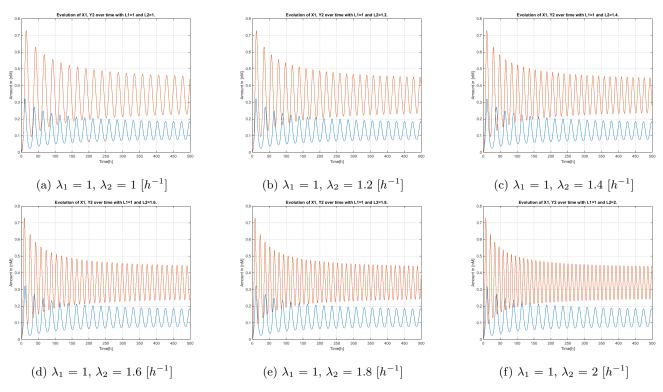


Figure 12:  $X_1(t)$  and  $Y_2(t)$  trajectories in a two-cells Model with K=0 We observe no synchronisation, as excepted. nothing else to say? :(

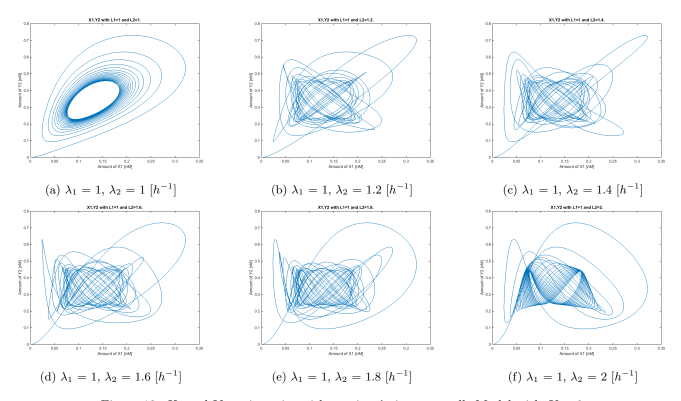


Figure 13:  $X_1$  and  $Y_2$  trajectories with varying  $\lambda_i$  in a two-cells Model with K=0 Once again, both cells tend to reach their limit cycles without any kind of interaction. The same observations as in Figure 11 can be made.

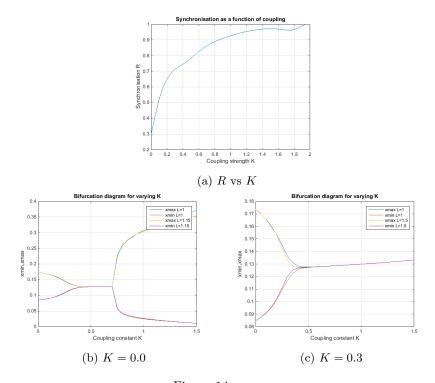


Figure 14: raraara

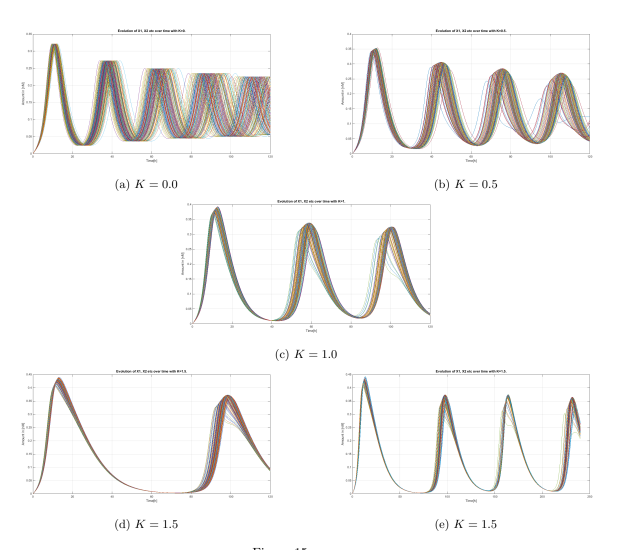
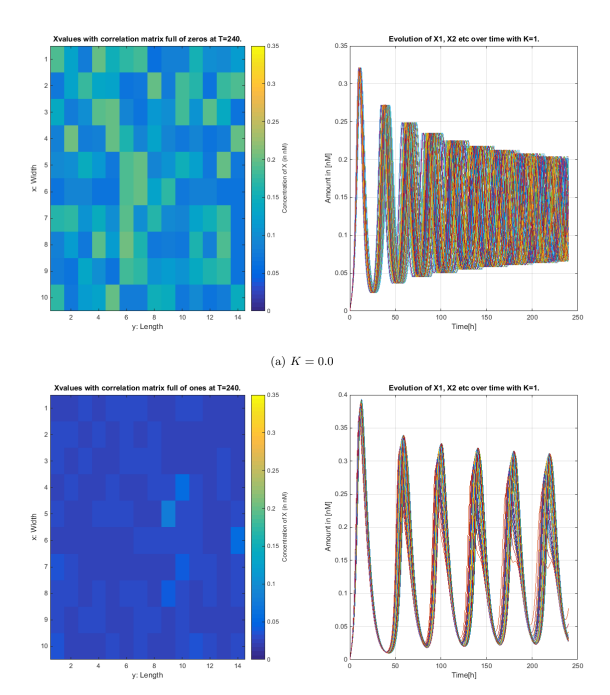


Figure 15: raraara



(b) K = 0.3

Figure 16: raraara