## BIOE147 - Fall 2013 - PS2

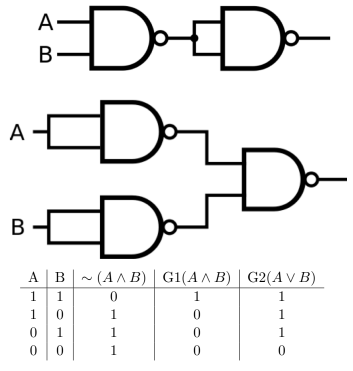
#### Zackery Field

October 20, 2013

## 1. Logic Gates and Biology

The NAND gate is often the only one used in electronic circuits. One of the main reasons for this is that the NAND gate is functionally complete, which means that all other logic gates can be represented by only NAND gates.

(a.) Write a truth table for the following NAND gate circuits and show what gate they represent:



- (b.) Design a biological system that performs the function of a NAND gate and the two gates above. Use only transcriptional regulation and use only the specific biological parts covered in the course.
  - **AND:** In order to implement the AND architecture we can using a two repressors. Let  $p_1, p_2$  be two constitutive promoters and  $p_3, p_4$  be two repressible promoters,  $r_3, r_4$  be the repressor protein of  $p_3, p_4$ , and  $s_3, s_4$  be the two small molecule repression inhibitors.  $p_1$  promotes the transcription of  $r_3$ , and  $p_2$  promotes the transcription of  $r_4$ . The reporter R is being repressed by both  $p_3$  and  $p_4$ . It is assumed that the repression of either  $p_3$  or

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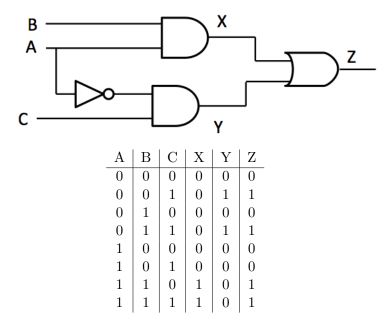
 $p_4$  is enough to repress the transcription of R. So, only the presence of both  $s_3$  and  $s_4$  will activate transcription of the reporter. Options for repressible systems are LacO/LacI and cl-ts. The first gate in the figure is the equivalent of an AND gate.

- **NAND:** The NAND gate can be created by implementing an AND gate and then simply flipping the output using a repressor. Starting with the AND gate above, make the output be another repressor protein  $r_5$  (instead of the reporter R). This repressor protein will act on on  $p_5$  and then repress the transcription of R. This will ensure that R is only repressed when both  $s_3$  and  $s_4$  are present. Since LacI/LacO and cl-ts were recommended for the AND gate, tetR can be used as the  $p_5/r_5$  system.
- OR: The second gate in the figure is an OR gate. To construct this gate you can make two orthogonal circuits that both regulate the production of the same reporter R (GFP). Each of these circuits will act in the same way. Let p be the repressor, r be the repressor protein that modulates p, and s be the small molecule inhibitor that inactives r. The repressor proteins for each of the circuits will be transcribed constitutively. Therefore, the presence of either the s for one circuit or the other (or both) will activate the transcription of the reporter R.
- (c.) Could we construct the two gates above using only the NAND gate in biological systems? Is it feasible to utilize the NAND only architecture in biological systems?

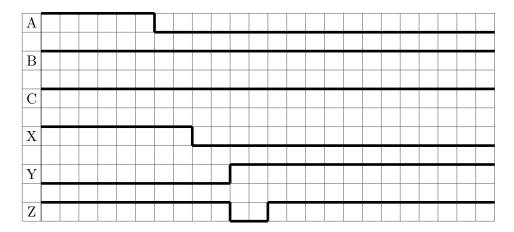
With the materials that we have been restricted to thusfar, no. If all parts known were available it may be feasible to construct the two example circuits, but beyond that it would not be feasible to construct larger systems with NAND only architecture. A key reason that NAND only architecture works in EE applications is that you can construct a many orthogonal NAND gates with ease, while in biology you do not have that luxury.

## 2. Time Delays in Circuits

(a.) Fill out the truth table and timing diagram. Assume all gates have a delay of one time unit.



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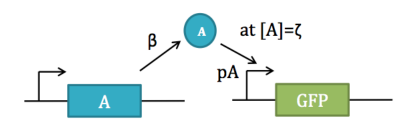
(b.) What happened to output Z when A switched signals?

There was a time delay of one time step when X turned to the 0-state, and a time step of two when Y turned to the 1-state. This meant that there was one time unit where both X and Y were in the 0-state, and therefore for the time step following, Z was also in the 0-state. If the system is designed for X and Y to fluctuate while Z maintains the 1-state invariant then the time delay observed that puts Z in the 0-state would cause issue with whatever behaved with the expectation that Z would always be in the 1-state.

(c.) Redesign the circuit above to eliminate the problem caused by the delay.

An addition of a buffer gate between A and the X gate would stall the A signal for another time step as Y switched from the 0-state to the 1-state. This would eliminate the one time step gap where Z is in the 0-state.

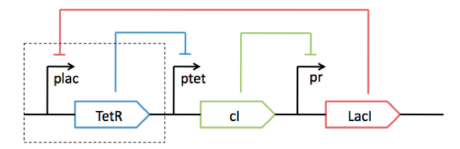
(d.) If there is no A in the system at time 0 and its expression is turned on, find how much time it would take for the cell to begin expressing GFP.



$$\frac{d[A]}{dt} = \beta - \alpha$$
$$[A] = \exp^{(\beta - \alpha)t}$$
$$t = \frac{\ln \zeta}{\beta - \alpha}$$

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## 3. Cooperative binding and part definitions



First consider the plac-TetR transcription unit that is repressed by a source of LacI.

(a.) Find an expression for the concentration of promoter-repressor (dimerized) complex at equilibrium in terms of the constants and the concentration of LacI.

$$\frac{d(plac_1)}{dt} = k_{\text{On}_{plac_1}}[plac_0] - k_{\text{Off}_{plac_1}}[plac_1]$$

$$\frac{d(plac_2)}{dt} = k_{\text{On}_{plac_1}}[plac_1] - k_{\text{Off}_{plac_2}}[plac_2]$$

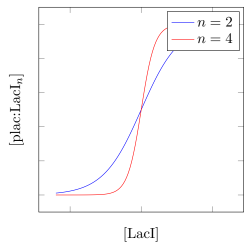
$$[plac_1] =$$

$$[plac_2] =$$

(b.) Under what conditions does the above expression contain a Hill-like term?

The above expression will have a Hill-like term whenever there is n-merization of the repressor proteins when they interact with the promoter.

With these conditions, plot transfer curves for [plac:LacI<sub>n</sub>] as a function of [LacI] when the Hill coefficient is n=2, and n=4.



In what circuits might n = 4 be more useful than n = 2 and vice verse?

n=4 would be more useful in any circuit where you need more digital response, and you have less concern on the effect of repressor protein production on cell fitness. On the other hand, n=2 has its place in a system where you want to see a change in the system occur at very small concentration of repressor protein.

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Now consider the whole circuit. Assume that repressor-promoter binding is much faster than the other reactions and use the same assumptions made in deriving the Hill equation from above

- (c.) Write differential equations for each species.
- (d.) Break the system into 1,3, and 18(?) parts. For each view of the system, briefly summarize the function of each part and list the inputs and outputs.
- (e.) Say you want to hook up a reporter to the circuit by introducting another plasmid with plac driving GFP. Qualitatively, how might the repressilator be affected? How might you minimize the interaction between the repressilator/reporter?

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# 4. Simulation Using Parts and Compositors

(a.) Simulate the repressilator using the SimPartComposition scripts. Model the repressor-promoter binding as you did in deriving the Hill form. It may be helpful to use the following parameters listed below.

(b.) Are oscillations more likely for high or low values of the following: n,  $k_{ts}$ , and  $k_{tl}$ ?