# Dynamic Model of Heavy Metal Detection Biosensor

Minghui Yin, Sherry Dongqi Bao Tianjin University

October 7, 2018

#### **Abstract**

### 1 Introduction

Modeling is a powerful tool in synthetic biology. It can provide us with an important engineering approach to characterize our pathways quantitatively and predict their performance, thus help us test and modify our design.

Through the dynamic model, we hope to gain insights of the characteristics of our whole circuit's dynamics.

### 2 Method

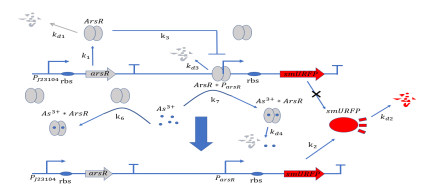


Figure 1: Schematic diagram of plasmid1

At the beginning, on the plasmid#1, the promoter  $P_{arsR}$  isn't bound with ArsR, thus it is active. ArsR and smURFP are transcribed and translated under the control of the promoters  $P_{arsR_u}$  and  $P_{arsR_d}$ , with subscript u and d representing upstream and downstream separately. The subscript l of smURFP in the equation means

leaky expression without the expression of  $As^{3+}$ . As ArsR is expressed gradually, it will bind with the promoter  $P_{arsR}$  and make it inactive. [pola2018novel]

$$P_{\text{J23104}} \xrightarrow{k_1} P_{\text{J23104}} + \text{ArsR} \tag{1}$$

$$P_{\text{arsR}_d} \xrightarrow{k_2} P_{\text{arsR}_d} + \text{smURFP}_l$$
 (2)

$$ArsR + P_{arsR} = \frac{k_3}{k_{-3}} ArsR^* P_{arsR}$$
 (3)

On the plasmid#2, the fusion protein of dCas9 and RNAP(RNA polymerase) are produced after transcription and translation, and sgRNA is produced after transcription.

$$P_{\text{tet}} \xrightarrow{k_4} P_{\text{tet}} + \text{dCas}9^* \text{RNAP}$$
 (4)

$$P_{\text{tet}} \xrightarrow{k_5} P_{\text{tet}} + \text{sgRNA}$$
 (5)

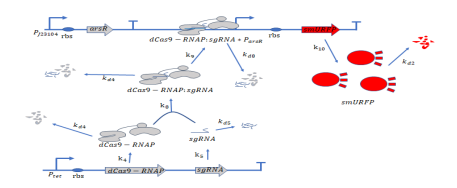


Figure 2: Schematic diagram of dCas9/RNAP

dCas9(\*RNAP) can bind with its target DNA sequence without cutting, which is at the upstream of the promoter  $P_{arsR_d}$ . Simultaneously, dCas9 can lead RNAP to bind with the promoter  $P_{arsR_d}$  and enhance the transcription of smURFP. However, because the promoter  $P_{arsR_d}$  has already bound with ArsR, as a result, RNAP can't bind with the promoter  $P_{arsR_d}$  [bikard2013programmable].

However, at the presence of  $As^{3+}$ , it can bind with ArsR, then dissociate ArsR and  $P_{arsR_d}$ , which makes the combination of RNAP and  $P_{arsR_d}$  possible.

(Declaration: 
$$[dCas9/RNAP] = [dCas9] = [RNAP]$$
;  $[P_{arsR_d}] = [P_{arsR_u}] = 0.5[P_{arsR}]$ )

$$ArsR + As^{3+} \xrightarrow{k_6} As^{3+*} ArsR$$
 (6)

$$ArsR^*P_{arsR} + As^{3+} \underset{k_{-7}}{\longleftarrow} As^{3+*}ArsR$$
 (7)

$$dCas9*RNAP + sgRNA \xrightarrow{k_8} dCas9*RNAP:sgRNA$$
 (8)

$$\mathsf{dCas}9^*\mathsf{RNAP}:\mathsf{sgRNA} + P_{\mathsf{arsR_d}} \xrightarrow[k_{-9}]{k_9} \mathsf{dCas}9^*\mathsf{RNAP}:\mathsf{sgRNA}^*\mathsf{P}_{\mathsf{arsR_d}} \tag{9}$$

$$P_{\mathsf{arsR}_\mathsf{d}} \xrightarrow{k_{10}} P_{\mathsf{arsR}_\mathsf{d}} + \mathsf{smURFP} \tag{10}$$

We then take degradation into account:

$$\operatorname{ArsR} \xrightarrow{k_{d1}} \emptyset \tag{11}$$

$$smURFP \xrightarrow{k_{d2}} \emptyset$$
 (12)

$$ArsR^*P_{arsR} \xrightarrow{k_{d3}} P_{arsR} \tag{13}$$

$$As^{3+*}ArsR \xrightarrow{k_{d4}} As^{3+}$$
 (14)

$$dCas9*RNAP \xrightarrow{k_{d5}} \emptyset$$
 (15)

$$sgRNA \xrightarrow{k_{d6}} \emptyset$$
 (16)

dCas9\*RNAP:sgRNA 
$$\xrightarrow{k_{d7}}$$
 Ø (17)

dCas9\*RNAP:sgRNA\*
$$P_{arsR} \xrightarrow{k_{d8}} P_{arsR}$$
 (18)

We can now write each reaction into ODEs (not following the order above). We will use v as an arbitrary variable, simply there to denote a rate of reaction. The "r0x" notation will be used later in the programming to remove clutter from the code. The several complexes involved:ArsR\*P $_{arsR}$ , As $^{3+*}$ ArsR, dCas9\*RNAP, dCas9\*RNAP:sgRNA\*P $_{arsR}$ , are respectively abbreviated as  $cplx_1$ ,  $cplx_2$ ,  $cplx_3$ ,  $cplx_4$ ,  $cplx_5$ .

(1)Production of protein ArsR (r01):

$$v = k_1[P_{J23104}]$$

(2) Degration of ArsR protein (r02):

$$v = -k_{d1}[ArsR]$$

Note that there should be a minus sign. This is because it will be decreasing the concentration of the ArsR protein . We can now put them together, to form an ordinary differential equation (ODE) for the rate of change in ArsR protein concentration.

$$\frac{d[ArsR]}{dt} = k_1[P_{arsR}] - k_{d1}[P_{arsR}][ArsR] \tag{1}$$

(3)Production of protein smURFP under unbound promoter (r03):

$$v = k_2[P_{arsR}]$$

(4)Production of protein smURFP under dCas9\*RNAP:sgRNA-bound promoter (r04):

$$v = k_{10}[cplx_5]$$

(5) Degration of smURFP protein (r05):

$$v = -k_{d2}[smURFP]$$

These two come together to form an ODE for the rate of change in smuRFP protein concentration:

$$\frac{d[smURFP]}{dt} = k_2[P_{arsR}] + k_{10}[cplx_5] - k_{d2}[smuRFP]$$
 (2)

(6) Combination of ArsR protein and  $P_{arsR}$  promoter(r06):

$$v = k_3 [ArsR][P_a rsR]$$

(7)Combination of  $As^{3+}$  and ArsR protein bound with  $P_{arsR}$  promoter , also dissociation of ArsR protein and  $P_{arsR}$  promoter(r07):

$$v = k_7 [As^{3+}][cplx_1]$$

(8) Degration of ArsR protein in the  $ArsR*P_{arsR}$  complex(r08):

$$v = -k_{d3}[cplx_1]$$

Similarly, these three form the ODE for the rate of change in  $ArsR*P_{arsR}$  concentration. :

$$\frac{d[cplx_1]}{dt} = k_3[ArsR][P_{arsR}] - k_7[As^{3+}][[cplx_1] - k_{d3}[cplx_1]$$
 (3)

Note that, in this ODE, there should be a minus sign in front of  $k_7$ . This is because it will be decreasing the concentration of  $ArsR*P_{arsR}$  complex. (9)Production of dCas9-RNAP fusion protein (r09):

$$v = k_4[P_{tet}]$$

(10) Degration of dCas9-RNAP fusion protein(r010):

$$v = k_{d5}[cplx_3]$$

(11)Combination of dCas9-RNAP fusion protein and it's sqRNA(r011):

$$v = k_8[cplx_3][sgRNA]$$

These three form the ODE for the rate of change in dCas9-RNAP concentration.

$$\frac{d[cplx_3]}{dt} = k_4[P_{tet}] - k_8[cplx_3][sgRNA] - k_{d5}[cplx_3] \tag{4}$$

Note that, in this ODE, there should be a minus sign in front of  $k_8$ . This is because it will be decreasing the concentration of dCas9-RNAP complex. (12)Production of sqRNA (r012):

$$v = k_5[P_{tet}]$$

(13) Degration of sqRNA(r013)

$$v = k_{d6}[sgRNA]$$

These two plus r010 come together to form an ODE for the rate of change in sqRNA concentration:

$$\frac{d[sgRNA]}{dt} = k_5[P_{tet}] - k_8[cplx_3][sgRNA] - k_{d6}[sgRNA]$$
 (5)

Similarly, in this ODE,there should be a minus sign in front of  $k_8$ . This is because it will be decreasing the concentration of sgRNA complex. (14)Combination of  $As^{3+}$  and ArsR protein (r014):

$$v = k_6 [As^{3+}][ArsR]$$

(15)Degration of ArsR protein in the  $As^{3+}*ArsR$  complex(r015):

$$v = -k_{d6}[As^{3+} * ArsR]$$

These two plus r06 come together to form an ODE for the rate of change in  $As^{3+}*ArsR$  concentration and an ODE for the rate of change in  $As^{3+}$  concentration:

$$\frac{d[cplx_2]}{dt} = k_6[As^{3+}][ArsR] + k_7[As^{3+}][cplx_1] - k_{d4}[cplx_2]$$
 (6)

Note that, in this ODE, there should be no minus sign in front of  $k_7$ , which is different from ODE(3). This is because it will be increasing the concentration of  $As^{3+}*ArsR$  complex.

$$\frac{d[As^{3+}]}{dt} = -k_6[As^{3+}][ArsR] - k_7[As^{3+}][cplx_1]$$
 (7)

(16)Combination of dCas9-RNAP:sqRNA complex and  $P_{arsR}$  promoter (r016):

$$v = k_9[cplx_4][P_{arsR}]$$

(17) Degration of dCas9-RNAP:sqRNA complex (r017):

$$v = -k_{d7}[cplx_4]$$

These two plus r010 come together to form an ODE for the rate of change in dCas9-RNAP:sgRNA complex concentration:

$$\frac{d[cplx_4]}{dt} = k_8[cplx_3][sgRNA] - k_9[cplx_4][P_{arsR}] - k_{d7}[cplx_4]$$
 (8)

Similarly, in this ODE, there should be no minus sign in front of  $k_8$ , which is different from ODE(4). This is because it will be decreasing the concentration of dCas9-RNAP:sgRNA complex.

(18))Degration of dCas9-RNAP:sgRNA complex in  $dCas9-RNAP:sgRNA*P_{arsR}$  compex(r018):

$$v = -k_{d8}[cplx_5]$$

This reaction and r016 come together to form an ODE for the rate of change in  $dCas9-RNAP:sgRNA*P_{arsR}$  complex concentration:

$$\frac{d[cplx_5]}{dt} = k_9[cplx_4][P_{arsR}] - k_{d8}[cplx_5] \tag{9}$$

At last, the ODEs for the rate of change in three promoter:

$$\frac{d[P_{J23104}]}{dt} = 0 ag{10}$$

$$\frac{d[P_{ArsR}]}{dt} = 0 {(11)}$$

$$\frac{d[P_{tet}]}{dt} = 0 (12)$$

Altogether, we have 8 ODEs:

$$\frac{d[ArsR]}{dt} = k_1[P_{arsR}] - k_{d1}[P_{arsR}][ArsR] \tag{1}$$

$$\frac{d[smURFP]}{dt} = k_2[P_{arsR}] + k_{10}[cplx_5] - k_{d2}[smuRFP]$$
 (2)

$$\frac{d[cplx_1]}{dt} = k_3[ArsR][P_{arsR}] - k_7[As^{3+}][[cplx_1] - k_{d3}[cplx_1]$$
 (3)

$$\frac{d[cplx_3]}{dt} = k_4[P_{tet}] - k_8[cplx_3][sgRNA] - k_{d5}[cplx_3] \tag{4}$$

$$\frac{d[sgRNA]}{dt} = k_5[P_{tet}] - k_8[cplx_3][sgRNA] - k_{d6}[sgRNA]$$
 (5)

$$\frac{d[cplx_2]}{dt} = k_6[As^{3+}][ArsR] + k_7[As^{3+}][cplx_1] - k_{d4}[cplx_2]$$
 (6)

$$\frac{d[As^{3+}]}{dt} = -k_6[As^{3+}][ArsR] - k_7[As^{3+}][cplx_1]$$
 (7)

$$\frac{d[cplx_4]}{dt} = k_8[cplx_3][sgRNA] - k_9[cplx_4][P_{arsR}] - k_{d7}[cplx_4]$$
 (8)

$$\frac{d[cplx_5]}{dt} = k_9[cplx_4][P_{arsR}] - k_{d8}[cplx_5] \tag{9}$$

$$\frac{d[P_{J23104}]}{dt} = 0 ag{10}$$

$$\frac{d[P_{ArsR}]}{dt} = 0 {(11)}$$

$$\frac{d[P_{tet}]}{dt} = 0 ag{12}$$

Table 1: Parameters

Rate constants	Value	Units	Reference
k1	1.999e-5	1/s	Berset et al.
k2	3.312e-6	1/s	Berset et al.
k3	3.3e7	1/M	Berset et al.
k4	1.995e-5	1/s	Estimated to be the same as in comparison to k1
k5	3.312e-6	1/s	Estimated to be the same as in comparison to k2
k6	1.66e7	1/M	Berset et al.
k7	1.26e4	1/s	Berset et al.
k8	1.6e-2	1/s	2017igem Munich
k9	1.66e-5	1/s	2017igem Munich
k10	4e-5	1/s	Estimated to be slow in comparison to k2
kd1	3.07e-3	1/s	Berset et al.
kd2	1e-5	1/s	Berset et al.
kd3	1e-3	1/s	Berset et al.
kd4	1.53e-3	1/s	Berset et al.
kd5	2e-2	1/s	Estimated to be fast in comparison to kd1
kd6	7.62e-3	1/s	Estimated according to Berset et al.
kd7	1e-2	1/s	Estimated to be slow in comparison to kd5
kd8	1e-1	1/s	Estimated to be slow in comparison to kd7

### 2.1 simulation

SimBiology toolbox provides functions for modeling, simulating, and analyzing biochemical pathways on basis of the powerful computing engine of Matlab.

COPASI is freeware developed with collaboration of VBI and EMLR. It provides almost the same functions as SimBiology, though not quite powerful. But compared with SimBiology, it provides a friendly user interface for model analysis, such as parameter estimation, and parameter scan.

Through the figure, we can see that the smURFP fluorescence gradually increased and then reached a steady state after a period of time in the presence of arsenic ions.

#### 2.2 sensitivity

Parameter sensitivity analysis is used to identify which parameters are more critical in effecting the output of the pathway, and help to gain a deeper insight of the structure and the function of the system. The sensitivity is calculated in this way:

$$sensitivity \frac{\Delta output}{\Delta parameter} \tag{19}$$

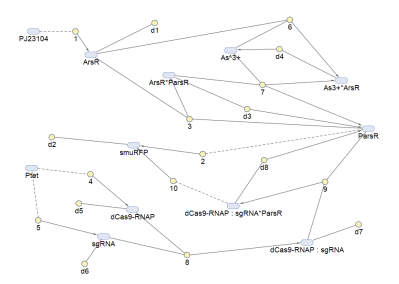


Figure 3: reaction map generated from the reaction set above using  $\mathsf{SimBiology}$   $\mathsf{Toolbox}$ 

The Matlab scripts used for sensitivity analysis see attached.

## References

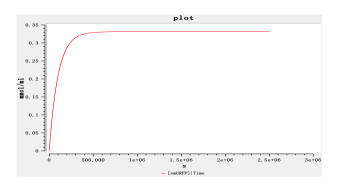


Figure 4: Schematic diagram of smURFP fluorescence by COPASI  $\,$ 

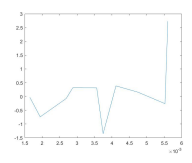


Figure 5: sensitivity of k1

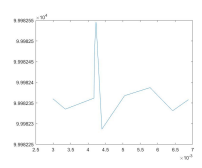


Figure 6: sensitivity of k2

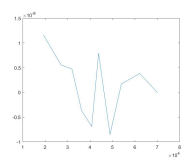


Figure 7: sensitivity of k3

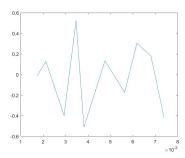


Figure 8: sensitivity of k4

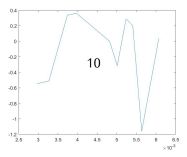


Figure 9: sensitivity of k5