

A quick start guide v1.1

September 03 2016

Department of Applied Mathematics and Computer Science Section of Embedded Systems Engineering Bio-Design Automation Group Richard Petersens Plads, Lyngby 3200 Technical University of Denmark

http://bda.compute.dtu.dk







Contents

About D-VASim	
 Analyzing the SBML model using D-VASim 1.1. Launching D-VASim 1.2. Generating a Virtual Laboratory Instrument 	4 4
2. Virtual Experimentation for Stochastic Simulations 2.1. Simulation Settings 2.2. Additional features of Stochastic Simulation in D-VASim	7 8
3. Virtual Experimentation for ODE Simulations	
4. General Software Options and Tips	18
References	20

About D-VASim

D-VASim stands for Dynamic Virtual Analyzer and Simulator. It is developed to analyze and simulate the genetic logic circuit models developed in the Systems Biology Markup Language (SBML) [1]. Practically, D-VASim can be used to simulate any bio-model available in SBML L3v1 (Level 3 version 1) core; however, it is solely designed for the simulations of genetic logic circuits, where user can apply external signals on the model during runtime.

D-VASim is developed on graphical programming language platform called LabVIEW (Laboratory Virtual Instrument Engineering Workbench) [2]. Besides graphical programming, textual programming language, like JAVA, is also used to integrate JSBML [3] library in it.

In this short document, you will learn how to use D-VASim (v1.1) to perform virtual laboratory experiments. This requires you to have presynthesized bio-model, in the form of SBML file, you want to test. This SBML can be obtained by creating a bio-model using any other bio-tool and generating its SBML L3v1 format.

Stochastic simulation algorithm (SSA) has been implemented to perform stochastic simulations of SBML models. Furthermore, D-VASim is also capable of simulating the deterministic behavior of a bio model by solving ordinary differential equations.

Latest version of D-VASim can be downloaded from <u>http://bda.compute.dtu.dk/downloads/</u>. The example SBML models, included in the download package, are developed at University of Utah by Myer's group [4]. The video demo of D-VASim v1.1 can be seen at <u>http://bda.compute.dtu.dk/user-manuals/</u>.

Conventions

	The following conventions appear in this document: This icon denotes a tip, which notifies you to advisory information.
	This icon denotes an alert, which notifies you to important information.
bold	Bold text denotes items that you must select or click or enter the value in the software, such as open file option or running the simulation button or entering the value of simulation speed etc.
i talic	Italic text denotes the name of a folder or a file path.
bold and italic	Bold and italic text denotes the name of a file.

1. Analyzing the SBML model using D-VASim

In this section, you will learn how to import and analyze the SBML model of a genetic circuit in D-VASim. In this document, we will use the SBML model of a genetic AND gate enclosed in the *Sample_SBML_Models* folder in the download package.

1.1. Launching D-VASim

The front interface of D-VASim, shown in Figure 1, will appear when you double-click on *D-VASim.app* (for MAC OS) or *D-VASim.exe* (for Windows OS).

		/ A C : T		D-VASim (Dyi		-					[
-www.sim	D-1	ASIM: L	Oynamic '	virtual /	Analyz	zer and	Simul	ator	Techni	cal Univers	ity of Denmark
elect SBML Mod	del (.xml)										
1							Obtain pa	rameters f	from Cello UCF		
odel Informati	on										Generate SSA VI
ID			Substance Units	Length Units	Extent Unit	ts Convers	ion Factor M	odel Comp	onents		
Nama			Time Units	Naluura Iluina	Anna Maita						enerate ODE VI
Name			Time Units	Volume Units	Area Units						
				,	,		· · · ·				
pecies Par	ameters	Reaction	s Events	Functions	Unit	s Com	partments	Initia	l Assignments	Rules	Constraints
	0 Name	Compartment	Initial Amount	Initial Conce	ntration	Units C)nlv Susbtan	ce Units	Boundary Cond	ition Cons	tant Conversio
· .		Compartment	Initial Amount	Initial Conce	entration l	Units C)nly Susbtan	ce Units	Boundary Cond	ition Cons	tant Conversio
		Compartment	Initial Amount	Initial Conce	ntration	Units C	nly Susbtan	ce Units	Boundary Cond	ition Cons	tant Conversio
		Compartment	Initial Amount	Initial Conce	entration	Units C	only Susbtan	ce Units	Boundary Cond	ition Cons	tant Conversio
· .		Compartment	Initial Amount	Initial Conce	ntration	Units C	Only Susbtan	ce Units	Boundary Cond	ition Cons	tant Conversio
		Compartment	Initial Amount	Initial Conce	ntration l	Units C	only Susbtan	ce Units	Boundary Cond	ition Cons	tant Conversio
· .		Compartment	Initial Amount	Initial Conce	entration	Units C	Only Susbtan	ce Units	Boundary Cond	ition Cons	tant Conversio
· .		Compartment	Initial Amount	Initial Conce	entration	Units C	Dnly Susbtan	ce Units	Boundary Cond	ition Cons	tant Conversio
iD		Compartment	Initial Amount	Initial Conce	entration	Units C	Only Susbtan	ce Units	Boundary Cond	ition Cons	tant Conversio
· .		Compartment	Initial Amount	Initial Conce	entration	Units C	Dniy Susbtan	ce Units	Boundary Cond	ition Cons	tant Conversio
· .		Compartment	Initial Amount	Initial Conce	entration	Units C	Only Susbtan	ce Units	Boundary Cond	ition Cons	tant Conversio
· .		Compartment	Initial Amount	Initial Conce	entration I	Units C	Only Susbtan	ce Units	Boundary Cond	ition Cons	tant Conversio
· .		Compartment	Initial Amount	Initial Conce	entration I	Units C	Only Susbtan	ce Units	Boundary Cond	ition Cons	tant Conversio
· .		Compartment	Initial Amount	Initial Conce	entration I	Units C	Only Susbtan	ce Units	Boundary Cond	ition Cons 	tant Conversio

Figure 1. Front interface of D-VASim.

Complete the following steps to import the SBML model of a genetic AND gate circuit.

- 1. Click **open file** button on the option "Select SBML Model (.xml)".
- 2. Navigate to the file *and_RB.xml* placed under the application directory./*D-VASim/Sample_SBML_Models/*.

This will run the JSBML library and display the SBML components as shown in Figure 2.

- \ _\Sim	D-\	/ASim: D	ynamic \	Virtual A	Analy	zer a	nd Simu	lator	Technical U	niversity o	f Denmark	
elect SBML Mo												
웹 Macintosh HD:U	sers:hasa	n:Documents:Gene	tic Circuits by Mye	ers:and_RB.xml			Obtain	parameters	from Cello UCF		rate SSA V	
lodel Informati	on									Gene	rate SSA v	
ID			Substance Units	Length Units	Extent U	nits Cor	version Factor	Model Comp	onents			
and			mole					listOfUnitDe				
Name			Time Units	Volume Units	Area Uni	ts		listOfCompa listOfSpecie		Generate ODE VI		
Created by iBioSi	m flatten r	outine		litre				listOfParam				
pecies Pa	rameters	Reactions	Events	Functions	Ur	nits (Compartments	Initia	al Assignments	Rules	Constraints	
	9											
ID	Name	Compartment	Initial Amount	Initial Conce	ntration	Units	Only Susbt	ance Units	Boundary Condition	Constant	Conversio	
CI	-	Cell	Initial Amount	Initial Conce	ntration	Units	Only Susbt	ance Units	Boundary Condition	false	Conversio	
CI GFP	-	Cell Cell	0.0 0.0	NaN NaN	ntration	Units	-	ance Units	•	false false	null null	
CI GFP Lacl	-	Cell Cell Cell	0.0 0.0 0.0	NaN NaN NaN	ntration	Units	true true true	ance Units	false false true	false false false	null null null	
CI GFP Lacl P1	-	Cell Cell Cell Cell	0.0 0.0 0.0 4.0	NaN NaN NaN NaN	ntration	Units	true true true true	ance Units	false false true false	false false false false	null null null null	
CI GFP Lacl P1 P2	-	Cell Cell Cell Cell Cell Cell	0.0 0.0 0.0 4.0 2.0	NaN NaN NaN NaN NaN	ntration	Units	true true true true true	ance Units	false false true false false	false false false false false	null null null null null	
Cl GFP Lacl P1 P2 P3	-	Cell Cell Cell Cell Cell Cell Cell	0.0 0.0 4.0 2.0 2.0	NaN NaN NaN NaN NaN NaN	ntration	Units	true true true true true true	ance Units	false false true false false false	false false false false false false	null null null null null null	
CI GFP Lacl P1 P2 P3 P3_mRNA	-	Cell Cell Cell Cell Cell Cell Cell Cell	0.0 0.0 4.0 2.0 2.0 0.0	NaN NaN NaN NaN NaN NaN NaN	ntration	Units	true true true true true true true true	ance Units	false false true false false false false	false false false false false false false	null null null null null null null	
CI GFP Lacl P1 P2 P3_mRNA Promoter_GFP	-	Cell Cell	0.0 0.0 4.0 2.0 2.0 0.0 2.0	NaN NaN NaN NaN NaN NaN NaN	ntration	Units	true true true true true true true true	ance Units	false false true false false false false false false	false false false false false false false false	null null null null null null null null	
Cl GFP Lacl P1 P2 P3 P3_mRNA	-	Cell Cell Cell Cell Cell Cell Cell Cell	0.0 0.0 4.0 2.0 2.0 0.0	NaN NaN NaN NaN NaN NaN NaN	ntration	Units	true true true true true true true true	ance Units	false false true false false false false	false false false false false false false	null null null null null null null	
Cl GFP Lacl P1 P2 P3 P3_mRNA Promoter_GFP	-	Cell Cell	0.0 0.0 4.0 2.0 2.0 0.0 2.0	NaN NaN NaN NaN NaN NaN NaN	ntration	Units	true true true true true true true true	ance Units	false false true false false false false false false	false false false false false false false false	null null null null null null null null	
Cl GFP Lacl P1 P2 P3 P3_mRNA Promoter_GFP	-	Cell Cell	0.0 0.0 4.0 2.0 2.0 0.0 2.0	NaN NaN NaN NaN NaN NaN NaN	ntration	Units	true true true true true true true true	ance Units	false false true false false false false false false	false false false false false false false false	null null null null null null null null	
Cl GFP Lacl P1 P2 P3 P3_mRNA Promoter_GFP	-	Cell Cell	0.0 0.0 4.0 2.0 2.0 0.0 2.0	NaN NaN NaN NaN NaN NaN NaN	ntration	Units	true true true true true true true true	ance Units	false false true false false false false false false	false false false false false false false false	null null null null null null null null	
Cl GFP Lacl P1 P2 P3 P3_mRNA Promoter_GFP	-	Cell Cell	0.0 0.0 4.0 2.0 2.0 0.0 2.0	NaN NaN NaN NaN NaN NaN NaN	ntration	Units	true true true true true true true true	ance Units	false false true false false false false false false	false false false false false false false false	null null null null null null null null	
Cl GFP Lacl P1 P2 P3 P3_mRNA Promoter_GFP	-	Cell Cell	0.0 0.0 4.0 2.0 2.0 0.0 2.0	NaN NaN NaN NaN NaN NaN NaN	ntration	Units	true true true true true true true true	ance Units	false false true false false false false false false	false false false false false false false false	null null null null null null null null	
Cl GFP Lacl P1 P2 P3 P3_mRNA Promoter_GFP	-	Cell Cell	0.0 0.0 4.0 2.0 2.0 0.0 2.0	NaN NaN NaN NaN NaN NaN NaN	ntration	Units	true true true true true true true true	ance Units	false false true false false false false false false	false false false false false false false false	null null null null null null null null	

Figure 2. D-VASim showing the SBML components of a genetic AND gate model.

All the SBML components of a model can be analyzed here in a clean tabular form. For example, The **Specie** tab shown in Figure 2 depicts the total number of specie used in the genetic AND gate model along with their corresponding details. Similarly, the **Reactions** tab shown in Figure 3 contains the information related to the reaction kinetics of a model. D-VASim also generates the ordinary differential equations of a model, which can be seen in Figure 3. These equations are used to simulate the deterministic behavior of SBML model.

Species Parameters	Reactions	Events	Fun	ctions U	nits (Compartme	ents 🗍 li	Initial Assignments		Rules	Constraints
Total Reactions 7 Reaction Info											
Reaction ID	Name	Reversible	Fast	Compartmen	t Reactan	ts Count	Reactants	R-Stoich	Constant	Products Co	ount Produ
Degradation_GFP		false	false	Cell	1		GFP	1.0	true	0	
Degradation_Cl		false	false	Cell	1		CI	1.0	true	0	
Degradation_TetR		false	false	Cell	1		TetR	1.0	true	0	
Degradation_Lacl		false	false	Cell	1		Lacl	1.0	true	0	T
•											F
Reaction Math											
Reaction ID	Math										
Degradation_GFP	kd*GFF	2									
Degradation_Cl	kd*Cl										
Degradation_TetR	kd*Tet	:R									
Degradation_Lacl	kd*Lac	:									T
•											F
Ordinary Differential Equations											
-(kd*Cl)+10.0*(ko_P1*ng_P1*	Ko_P1*RNAP/	(1+Ko_P1*F	RNAP+	(Kr_Lacl_P1*La	cl)^nc_Lac	_P1))+10.0)*(ko_P2*ng	P2*Ko P	2*RNAP/(1+	Ko_P2*RNAP	+(Kr_TetR_P
-(kd*GFP)+10.0*(ko_Promoter											
•											Þ

Figure 3. Reactions tab showing the reaction kinetics of a genetic AND gate model.

1.2. Generating a Virtual Laboratory Instrument

Once the components of SBML model are analyzed, a virtual environment for the interactive simulation can be generated simply by clicking on any of the two options, **Generate SSA VI** or **Generate ODE VI** shown at the top right corner of a tool. Now click **Generate SSA VI** button to generate a virtual instrument (VI) for stochastic simulations of a genetic AND gate model. This will bring up a virtual instrument shown in Figure 4.

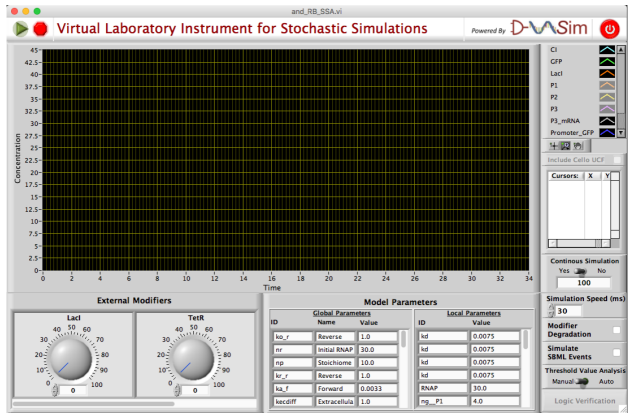


Figure 4. Virtual Instrument specifically for a genetic AND gate model.

Any instrument for stochastic simulation generated by D-VASim for the first time does neither reveal the names of graph legend nor the modifier controls, but only the parameters information. Pressing the **RUN** button (\triangleright shown at the top left corner) starts the simulation and makes the graph legends and modifiers visible.

2. Virtual Experimentation for Stochastic Simulations

The virtual instrument (shown in Figure 4) looks similar to a physical instrument, which can be used to interact with the model by tuning the input concentrations with the control knobs and observing the effects graphically.

Once the simulation is started after pressing the button, user can increase or decrease the concentration of input (or external modifier) specie and observe its effects on the model during run time. This runtime interaction with the model gives user an insight of being in the lab performing wet-lab experiments.

Figure 5 shows the screenshot of the stochastic simulation of a genetic AND gate taken randomly after ~2150 time units has lapsed. This figure signifies that unlike SBML events, which are predefined in the SBML file, user can interact with the model and change the concentration of input specie to any level and at any instant of time. In case if the concentration of input specie are required to be triggered to any specific level instantly, the numeric displays located beneath the control knobs can be used. As same as the user interact with the model by varying the input concentration using control knobs, the parameters can also be varied and their effects can be observed graphically during runtime.

When a **STOP** button (located besides the run button at the top left corner) is pressed, screenshot of a graphical window is captured and saved inside *Sim_Results* folder located in the default application directory.

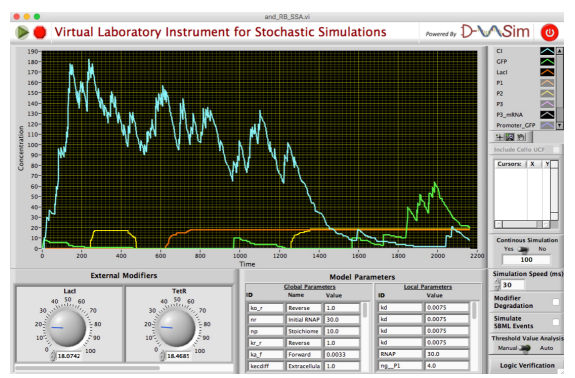


Figure 5. Screen capture of the stochastic simulation of a genetic AND gate.

Each image is saved by the following naming convention; "<modelname>_SSA_Screenshot.bmp", where <model-name> is the name of a model. For example, the screenshot of a graphical window of a genetic AND gate model is saved as *and_RB_SSA_Screenshot.bmp*. The simulation traces of the entire session are also saved in the *Sim_Results* folder with the convention "<model-name>_SSADatalog.xls". Pressing the **RUN** button again starts the simulation from the beginning.

Alert – When the **STOP** button is pressed, the captured screenshot and the data log file overwrites any previously saved files of the same model.

Tip – The properties of graphical plots, including colors, can be changed by a left mouse click on the color of graph legend and selecting the desired menu.

2.1. Simulation Settings

The simulation settings shown at the right bottom corner contains the following six simulation options.

Continous Simulation
Yes 🌨 No
100







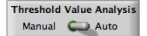


perform continuous simulation or for a specific interval of time. If switched to "No", the box below can be used to specify the simulation time for which user wants to run the simulation. The default value of this switch is "Yes" meaning that the continuous simulation is always performed unless specified.
2 Simulation Speed: This option allows you to speed up or slow

1. **Continuous Simulation:** User may specify here if they want to

- 2. **Simulation Speed**: This option allows you to speed up or slow down the simulation by entering the numerical value in milliseconds.
- 3. **Modifier Degradation**: When checked, perform more realistic simulation by allowing the input concentrations to degrade over time.
- 4. **Simulate SBML Event**: When checked, simulate the events described in SBML file.

Tip – The triggering time and the assignment value of SBML events can be changed during run time until the time of event occurs. For example, if the triggering time of event is 1000 time units, the trigger time and the assignment value can be changed before 1000 time units has lapsed.



5. Threshold Value Analysis:

Manual – In this mode, VI assume that the threshold value of input specie and the propagation delays are already known. Auto – When this mode is selected, VI estimates the threshold value based on the setting tab shown in Figure 6 below.

• **Start at** – Specify the initial concentration of input specie from which the analysis is desired to be started. It is recommended to keep this value 0 for correct results.





• **Increment of** – This control allow you to set up the concentration value, which the tool increments during each iteration for checking threshold concentration.

For example: If you set

Start at: 0

Increment of: 5

Stop at: 15

Tool will check the threshold concentration for 4 different levels of concentration. i.e. $0 \rightarrow 5 \rightarrow 10 \rightarrow 15$, for all possible input combinations.

O O Thre	shold Value	Settings
Input concentra	ation values	
Start at	Increment of	of Stop at
Assumed Time	e Delay	Settling Time
No. of iteration	<u>15</u>	Name of Output GFP
Verification RT	-	
<u>% Acceptance</u> 90	of consistenc	Υ
	ОК	1

Figure 6. Settings dialogue for automatic threshold value analysis.

- **Stop at** Specify the concentration of inputs at which the threshold analysis should be stopped.
- Assumed Time Delay Specify initially assumed time delay.
- No. of iterations For how many times a user wants to verify if the estimated threshold value is consistent.
- Verification RT For how long a user wants to verify the estimated threshold value for each no. of iterations mentioned above.
- Settling Time It is also an assumed value, which specifies the time required by a model to get stable. During verification process, D-VASim triggers the concentration of inputs to the estimated threshold level once this time value has elapsed. Thus the verification starts once the input is triggered to its estimated threshold value and therefore it should run for enough amount of time to measure the propagation delay as well as the correctness of an estimated threshold value. Depending on the complexity of input-

output stage, the value of "Verification RT" is suggested to be at least 3-4 times higher than the value of *Settling Time*.

- Name of Output D-VASim now allows user to perform the threshold value and propagation delay analysis either for an entire circuit or for the intermediate circuit components. In this field, user can specify the name of output specie. The timing analysis is performed between the input specie, selected as control knobs (see section 2.2) and the output specie specified in this field.
- % Acceptance of consistency Specify the minimum percentage of threshold value consistency a user want. If the chosen value is 90, the tool give the results only when the consistency of estimated threshold value is more than or equal to 90%.

Alert – It is an indication of wrong parameter selection if either tool takes too long to estimate the results or produce incorrect results. In this case user should consider increasing the value of "Verification RT" further higher (beyond 3-4 times) than the value of "Settling Time". Though assuming large values of these parameters, including "Assumed Time Delay", may increase the estimation time, but gives the better estimation.

When the tool finish estimating the threshold value with the desired minimum percentage of consistency, the results are displayed in the dialogue box as shown in Figure 7. The value of estimated propagation delay is the average value over all "No. of iterations" specified in the settings dialogue (Figure 6). The value enclosed in braces after the average estimated propagation delay specifies the standard deviation.

💿 💿 🔿 Threshold Analysis Results
Results of Threshold Value Analysis
Estimated Threshold Value
Estimated Propagation Delay 581 (±75.11)
Output Consistency (%) 100
ОК

Figure 7. Dialogue box indicating the estimated results of threshold value and propagation delay analysis.

The results are stored in the current application directory with the following naming convention; "<model-name>_SSA-TP-Analysis.txt", where <model-name> is the name of a model. For the case of genetic AND gate, the results will be saved as *and_RB_SSA-TP-Analysis.txt*.



Logic Verification

Logic Verification – Obtain the Boolean logic exist in the model. Clicking this button prompts the user to input the threshold values of input and output specie, as shown in Figure 8. When automatic threshold value analysis is performed, this box is auto-populated with the estimated threshold value. This option is disabled when the simulation is stopped.

Alert – Make sure to apply all the possible input combinations each after the correct propagation delays. The results of boolean logic analysis will otherwise be incorrect.

Alert – Though, D-VASim v1.1 supports the logic verification of n-input genetic logic circuits, but the logic verification algorithm is tested for up to 2-input genetic circuits. It is because the models for more complex circuits are not available.

🔴 🕘 🔘 Th	reshold Values
Logic Ve	rification Settings
Select Outp GFP Enter Three	shold Values
GFP	10
Lacl	10
TetR	10
	ОК

Figure 8. Dialogue box to set up the threshold values for Boolean logic analysis.

When all the threshold values are entered, a new window (Figure 9) pops up indicating the logic function, in terms of a Boolean expression, extracted from the genetic logic circuit model. This window also contains the entire simulation data as analog data points and their corresponding digital data according to the defined threshold values. If all the input combinations are applied correctly during the simulation, a message in green text (Figure 9 (a)) will appear showing the percentage fitness of the boolean expression in the acquired simulation data. If all the possible input combinations are not applied, a boolean expression may appear with the message in red text (Figure 9 (b)) indicating that all input combinations are not applied correctly. Boolean expression box may also appear blank when all the input combinations are not applied correctly. A correct set of input combinations is the one in which all the possible input combinations are applied each after the minimum propagation delay.

000	Boole	ean Logic	Results				
Logic Expression	ression Simulation Data						
Lacl.TetR	Time	GFP Analog	GFP Digitial	Lacl Analog	Lacl Digitial	TetR Analog	TetR 🔺 Digitial
	0.00	0.0	0	0.0	0	0.0	0
	8.62	0.0	0	0.0	0	0.0	0
Fitness = 99.88 %	22.16	0.0	0	0.0	0	0.0	0
Tuless = 33.00 %	25.50	0.0	0	0.0	0	0.0	0
	27.27	0.0	0	0.0	0	0.0	0
	28.65	0.0	0	0.0	0	0.0	0
Close	34.83	0.0	0	0.0	0	0.0	0 🔻
			-		-		· ·

(a)

000	Boole	ean Logic	Results				
Logic Expression	Simulatio	n Data					
Lacl.TetR	Time	GFP Analog	GFP Digitial	Lacl Analog	Lacl Digitial	TetR Analog	TetR J Digitial
	0.00	0.0	0	0.0	0	0.0	0
J	10.64	0.0	0	0.0	0	0.0	0
All input combinations	20.94	0.0	0	0.0	0	0.0	0
are not applied correctly!	22.34	0.0	0	0.0	0	0.0	0
are not apprica correctly.	22.94	0.0	0	0.0	0	0.0	0
	29.06	0.0	0	0.0	0	0.0	0
Close	32.19	0.0	0	0.0	0	0.0	0
		·	i _		i _		· - L
		(b)					

Figure 9. Boolean logic extracted from the analog simulation data along with its corresponding digital data. (a) Percentage fitness appears when all the input combinations are properly applied.

(b) Error message appears when all the input combinations are not applied correctly before trying to estimate the boolean logic of a circuit.

2.2. Additional features of Stochastic Simulation in D-VASim

In D-VASim v1.1, two new features have been introduced, which allows user to:

- 1) Create control knobs of specie manually. This not only helps user to vary the concentration of selected specie during runtime, but also perform the timing analysis between these specie and the specified output.
- 2) View the mixed signal waveforms (digital and analog) for better timing analysis.

The control knobs for the input specie can be created by dropping down the **Options** menu and selecting **External Inputs** as shown in Figure 10(a). When **External Inputs** is selected, a pop-up window is appeared as shown in Figure 10(b). The user can drag the specie name from left-hand box to right-hand box in order to create its control knob. In Figure 10(b), LacI and TetR specie are selected whose control knobs are created as shown in Figure 4. These control knobs can only be created when the simulation is not running.

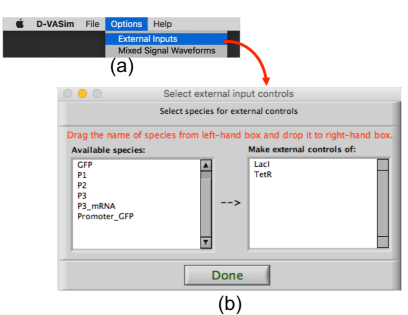


Figure 10. Process of creating control knobs for specie. (a) Select **External Inputs** from **Options** dropdown menu. (b) Drag the name of specie from left-hand to right-hand box to create its control knob.

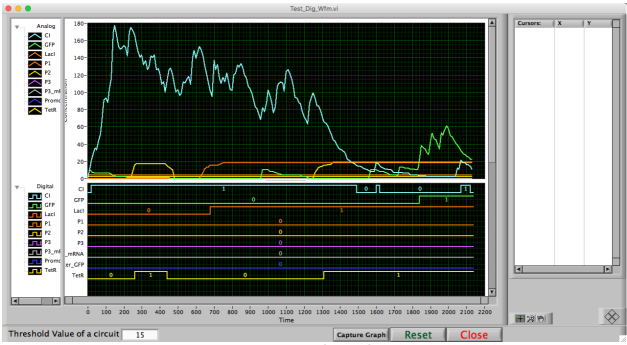


Figure 11. Mixed signal waveforms of genetic AND gate.

Similarly, after performing interactive stochastic simulation of a model in the D-VASim virtual laboratory environment, the mixed signal waveforms for timing analysis can be viewed by selecting **Mixed Signal Waveforms** option from the **Options** dropdown menu. Figure 11 shows the window of mixed signal waveforms generated for the simulation traces of genetic AND gate shown in Figure 5. This figure shows the analog and digital traces, in parallel, for genetic AND gate. The digital curves are generated based on the estimated threshold value shown in Figure 7. This value is automatically extracted if the auto threshold value analysis is performed, otherwise user can enter the threshold value and **Reset** the curves. **Capture Graph** button takes the screen-shot of waveforms and save it inside *Sim_Results* folder under the default application directory. Each image is saved by the file name *MixedSignalWaveforms.bmp*. The blank vertical box shown at the right-hand side in Figure 11 allows user to create cursors (see section 4 for more details about how to create cursors).



Alert – When the Capture Graph button is pressed, the captured screenshot overwrites any previously saved files of the same model.

Tip – The properties of graphical plots, including colors, can be changed by a left mouse click on the color of graph legend and selecting the desired menu.

3. Virtual Experimentation for ODE Simulations

Similar to generating virtual instrument for stochastic simulations, the virtual instrument for deterministic simulations can also be generated by clicking on **Generate ODE VI** button depicting in Figure 2.

When the button is pressed, the deterministic simulation results of a genetic AND gate for 6000 time units are appeared as shown in Figure 12. As same as in the case of stochastic simulation, user can change the concentration of external input specie using control knobs for deterministic simulation also. However, every single change in the input concentration or parameters results in the execution of selected simulation algorithm for the defined interval of time and plots the new set of curves.

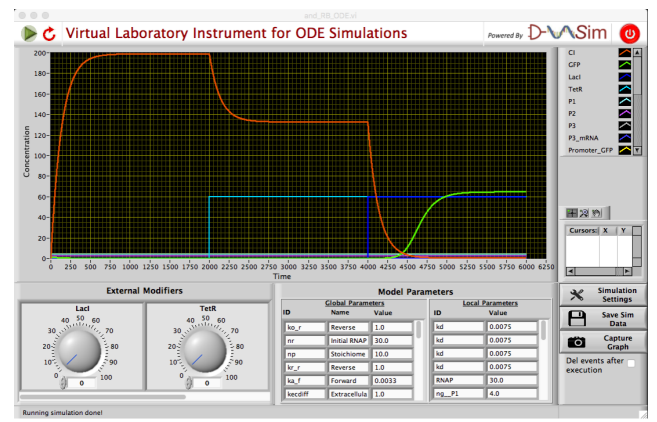


Figure 12. Screen capture of the deterministic simulation of a genetic AND gate.

The horizontal *status bar* is shown at the bottom in Figure 12. It shows the status of the actions being performed in this VI for ODE simulations. For example, pressing the run button shows the status of simulation as shown in Figure 13 below. The progress bar shown at right-hand side depicts the percentage of results currently being obtained. The plot is visible once the progress bar reaches to 100 %.

```
Running simulation ...
```

65%

Figure 13. Screen capture of the status bar showing that the VI is running simulation. The progress bar shown at right-hand side indicates that the 65% of simulation data is obtained.

When a **RESET** button (\checkmark) button is pressed, screenshot of a graph window and the simulation data is captured and saved under *Sim_Results* folder located in the default application directory. Each image is saved by the following naming convention; "<modelname>_ODE_Screenshot.bmp", where <model-name> is the name of a model. For example, the screenshot of a graphical window of a genetic AND gate model is saved as *and_RB_ODE_Screenshot.bmp*. The simulation traces of the entire session are saved with the naming convention; "<model-name>_ODEDatalog.xls". Pressing the **RUN** button again starts the simulation from the beginning.

Alternatively, the simulation data and the graphical plots can be stored by pressing the buttons shown in Figure 14(a) and 14(b) respectively.



Figure 14. Available options in ODE VI to (a) Save simulation data. (b) Capture graph window.

Alert – When the **RESET** or **Save Sim Data** or **Capture Graph** button is pressed, the captured screenshot and the data log file overwrites any previously saved files of the same model.



Tip – The properties of graphical plots, including colors, can be changed by a left mouse click on the color of graph legend and selecting the desired menu.

3.1. Simulation Settings

The **Simulation Settings** button (shown in Figure 15(a)) helps user to set the desired simulation parameters. A settings window, shown in Figure 15(b), is popped-up when user clicks on the **Simulation Settings** button.

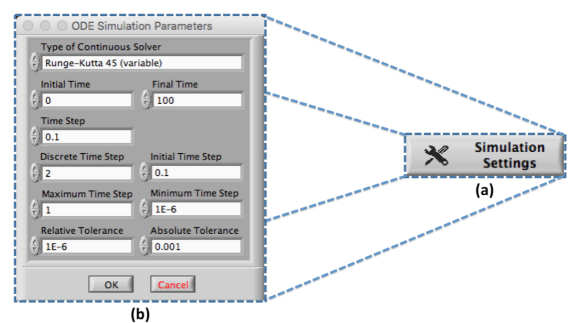


Figure 15. ODE simulation properties (a) Simulation settings option. (b) Window of simulation settings.

D-VASim VIs for ODE simulation supports the following ten continuous solvers:

- 1. Runge-Kutta 1 (Euler)
- 2. Runge-Kutta 2
- 3. Runge-Kutta 3
- 4. Runge-Kutta 4
- 5. Runge-Kutta 23 (variable)
- 6. Runge-Kutta 45 (variable)
- 7. BDF (variable)
- 8. Adams-Moulton (variable)
- 9. Rosenbrock (variable)
- 10. Discreet States Only

The parameters shown in Figure 15(b) are set as the default parameters. There is one other option available at the right bottom of ODE VI, which is described below

Del events after execution **Del events after execution:** When marked checked, the events are deleted internally after executing once. This helps increasing the simulation speed by avoid checking the executed events repeatedly.



Tip – If the desired simulation results are not obtained, try simulating the model again by checking or unchecking the option "Del events after execution".

4. General Software Options and Tips

Following are some general tips and options for both stochastic and ODE virtual instruments generated by D-VASim.



Tip – Hover mouse over simulation options to pop up the tip strip with a short description.

Tip – Pressing keys CMD+SHIFT+H (on MAC OS) and CTRL+H (on Windows OS) open the context help.

Tip – User can right click on the graph window to get the runtime shortcut menu shown in Figure 16 below.

Copy Data
Description and Tip Visible Items
Clear Graph Create Annotation
Delete All Annotations
/ AutoScale X
/ AutoScale Y
/ Smooth Updates
Autosize Plot Legend
Optional Plane
Export •

Figure 16. Runtime shortcut menu of graph window.



Graph Palette: This option can be used to zoom-in, zoom-out, expand graph etc.

Cursors:	X	Y	
•		F	

Cursor Legend: With the help of this box, cursors can be set to track the values of desired specie. For example, single-plot cursor can be set to monitor GFP specie by **right clicking** in the Cursors window and then selecting **Single-Plot** from **Create Cursors** option, as depicted in Figure 17(a). Once the cursor is created, it can be snap to GFP specie by **right clicking** on the created cursor and then go to **Snap To** option and select **GFP**. These steps are shown in Figure 17(b).



Figure 17. Steps of creating cursors on specie.

Finally, similar to any physical instrument, virtual instruments powered by D-VASim also have emergency shut down button (⁽¹⁾) shown at the top right corner of a virtual instrument. This button can be used to stop and shut down the instrument instantly.

References

- [1]. Systems Biology Markup Language, <u>http://sbml.org/Main_Page</u>.
- [2]. National Instruments LabVIEW, <u>http://www.ni.com/labview/</u>.
- [3]. Dräger A, Rodriguez N, Dumousseau M, Dörr A, Wrzodek C, Le Novère N, Zell A, and Hucka M. JSBML: a flexible Java library for working with SBML. *Bioinformatics* (2011), 27(15): 2167–2168.
- [4]. Chris J. Myers. Engineering Genetic Circuits. Chapman & Hall/CRC Press, July 2009.