

xml2jupyter: Mapping parameters between XML and Jupyter widgets

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Software

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Summary

Jupyter Notebooks (Kluyver et al. 2016, Perkel (2018)) provide executable documents (in a variety of programming languages) that can be run in a web browser. When a notebook contains graphical widgets, it becomes an easy-to-use graphical user interface (GUI). Many scientific simulation packages use text-based configuration files to provide parameter values and run at the command line without a graphical interface. Manually editing these files to explore how different values affect a simulation can be burdensome for technical users, and impossible to use for those with other scientific backgrounds. xml2jupyter is a Python package that addresses these scientific bottlenecks. It provides a mapping between configuration files, formatted in the Extensible Markup Language (XML), and Jupyter widgets. Widgets are automatically generated from the XML file and these can, optionally, be incorporated into a larger GUI for a simulation package, and optionally hosted on cloud resources. Users modify parameter values via the widgets, and the values are written to the XML configuration file which is input to the simulation's command-line interface. xml2jupyter has been tested using PhysiCell (Ghaffarizadeh et al. 2018), an open source, agent-based simulator for biology, and it is being used by students for classroom and research projects. In addition, we use xml2jupyter to help create Jupyter GUIs for PhysiCell-related applications running on nanoHUB (Madhavan et al. 2013).

A PhysiCell configuration file defines model-specific <user_parameters> in XML. Each parameter element consists of its name with attributes, defining its data type, units (optional), description (optional), whether the widget should be hidden (optional), and the parameter's default value. The attributes will determine the appearance and behavior of the Jupyter widget. For numeric widgets (the most common type for PhysiCell), xml2jupyter will calculate a delta step size as a function of the default value and this step size will be used by the widget's graphical increment/decrement feature.

To illustrate, we show the following simple XML example, containing each of the four (currently) supported data types and the various attributes:



radius	250	micron	initial tumor radius
threads	8		
fix_persistence			

Figure 1: Simple example of XML parameters as Jupyter widgets.

When we map this into Jupyter widgets, we obtain the rendered results in Figure 1. Notice the color parameter is not displayed since we specified it should be hidden in the XML. The name of the other parameters, their values, and attributes, if present, are displayed in rows (as disabled Jupyter button widgets). Using alternating row colors ("zebra stripes") helps visually match associated fields and avoid changing the wrong parameter value. For numeric widgets (type "int" or "double"), we compute a delta step value based on the magnitude (log) of the initial value. For example, the radius widget will have a step value of 10, whereas threads will have a step value of 1.

For a more realistic example, consider the config_biorobots.xml configuration file (found in the config_samples directory). The XML elements in the <user_parameters> block include the (optional) description attribute which briefly describes the parameter and is displayed in another widget. To demonstrate xml2jupyter on this XML file, one would: 1) clone or download the repository, 2) copy the XML configuration file to the root directory, and 3) run the xml2jupyter.py script, providing the XML file as an argument.

```
$ cp config_samples/config_biorobots.xml .
$ python xml2jupyter.py config_biorobots.xml
```

The xml2jupyter.py script parses the XML and generates a Python module, user_params.py, containing the Jupyter widgets, together with methods to populate their values from the XML and write their values back to the XML. To "validate" the widgets were generated correctly, one could, minimally, open user_params.py in an editor and inspect it.

But to actually see the widgets rendered in a notebook, we provide a simple test:

```
$ python xml2jupyter.py config_biorobots.xml test_user_params.py
$ jupyter notebook test_gui.ipynb
```

This should display a minimal notebook in your browser and, after selecting Run all in the Cell menu, you should see the notebook shown in Figure 2.

PhysiCell Jupyter GUI

Our ultimate goal is to generate a fully functional GUI for PhysiCell users. xml2jupyter provides one important piece of this - dynamically generating widgets for custom user parameters for a model. By adding additional Python modules to provide additional components (tabs) of the GUI that are common to all PhysiCell models, a user can configure, run, and visualize output from a simulation. Two tabs that provide visualization of output files are shown in Figure 3 with results from the biorobots simulation. Note that some of the required modules are not available in the Python standard library, e.g., Matplotlib (Hunter 2007) and SciPy (Oliphant 2007). We provide instructions for installing these additional dependencies in the repository README.



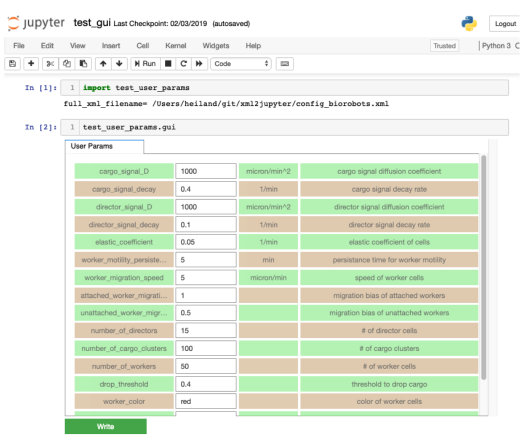


Figure 2: The biorobots parameters rendered as Jupyter widgets.

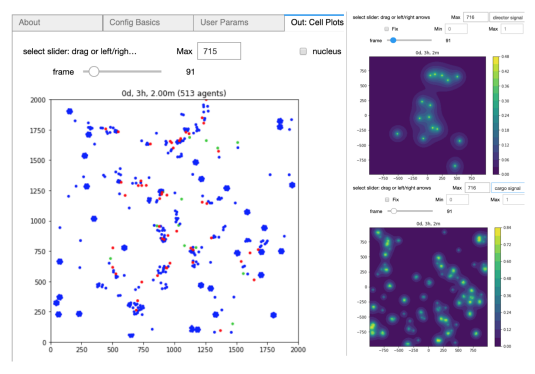


Figure 3: Plotting the biorobots (cells; left) and signals (substrates; right).



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Load Config DEFAULT v pc4cancerbots									
About	Config Basics	User Para	ıms	Out: Cell Plots	Out: Substrate Plots				
			10000						
	therapy_activation_time		10080		min	time at which worker and cargo cells are injected			
number_of_injected_cells			500			number of injected cells for therapy			
	worker_fraction		0.1			fraction of injected cells that are workers			
	save_interval_after_therapy_start		60		min	save (output) interval after therapy begins			
	cargo_o2_relative_uptake		0.1			cargo cell oxygen uptake rate (relative to cancer cells)			
	cargo_apoptosis_rate		0.00004065	5	1/min	apoptosis rate for cargo cells			
	cargo_relative_adhesion		0			cargo cell adhesion (relative to cancer cells)			
	cargo_relative_repulsion		5			cargo cell repulsion (relative to cancer cells)			
	worker_o2_relative_uptake		0.1			worker cell oxygen uptake rate (relative to cancer cells)			
	worker_apoptosis_rate		0		1/min	apoptosis rate for workers			
	worker_relative_adhesion		0			worker cell adhesion (relative to cancer cells)			
	worker_relative_repulsion		5			worker cell repulsion (relative to cancer cells)			
	worker_motility_persistence_time		5		min	motility persistence time for workers			
	worker_migration_speed		2		micron/min	migration speed for workers			
	attached_worker_migration_bias		0.5			migration bias for attached workers			
u	unattached_worker_migration_bias		0.5			migration bias for unattached workers			
mo	otility_shutdown_detection_thresho	d	0.001			worker stops if chemoattractant < threshold			
	elastic_coefficient		0.05		1/min	elastic coefficient for focal adhesion			
	cargo_release_o2_threshold		10		mmHg	oxygen threshold for releasing cargo			
	max_elastic_displacement		50		micron	maximum distance to maintain focal adhesion			
	damage_rate		0.03333		1/min	damage rate during therapy			
	repair_rate		0.004167		1/min	repair rate during therapy			
	drug_death_rate		0.004167		1/min	maximum drug-induced death rate			
	tumor_radius		200		micron	initial tumor radius			

Figure 4: The cancer biorobots parameters in a nanoHUB Jupyter application.

Extensions and Discussion

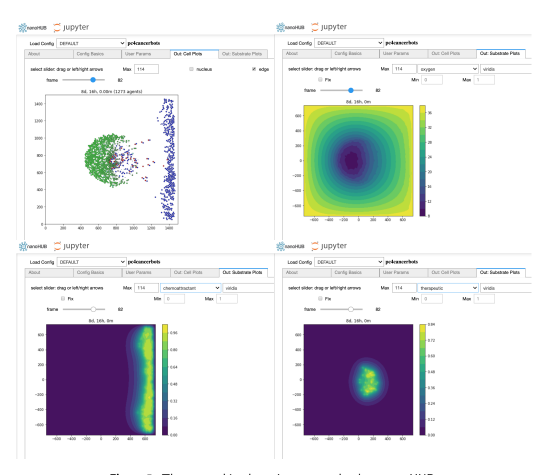
We hope others will be inspired to extend the core idea of this project to other text-based configuration files. XML is only one of several data-interchange formats, and while we created this tool for XML-based configurations based on needs to create GUIs for PhysiCell projects, the approach should be more broadly applicable to these other formats. And while the additional Python modules that provide visualization are also tailored to PhysiCell output, they can serve as templates for other scientific applications whose input and output file formats and provide similar functionality.

xml2jupyter has helped us port PhysiCell-related Jupyter tools to nanoHUB, a scientific cloud for nanoscience education and research that includes running interactive simulations in a browser. For example, Figure 4 shows the xml2jupyter-generated *User Params* tab in our pc4cancerbots tool running on nanoHUB. Figure 5 shows the cells (upper-left) and three different substrate plots for this same tool. This particular model and simulation is described in this video.

Other PhysiCell-related nanoHUB tools that have been created using xml2jupyter include pc4heterogen, pcISA, and pc4cancerimmune. Readers can create a free account on nanoHUB and run these simulations for themselves. We encourage students to use xml2jupyter to create their own nanoHUB tools of PhysiCell models that 1) can be run and evaluated by the instructor, 2) can be shared with others, and 3) become part of a student's living portfolio. (Another repository, https://github.com/rheiland/tool4nanobio, provides instructions and scripts to help generate a full GUI from an existing PhysiCell model.)

We welcome suggestions and contributions to xml2jupyter. For example, currently, we arrange the generated parameter widgets vertically, one row per parameter. This is an appropriate layout for an educational setting. But if a GUI will be used by researchers who are already familiar with the parameters, it may be preferable to generate a more compact layout of widgets, e.g., in a matrix with only the parameter names and values.





 $\textbf{Figure 5:} \ \ \textbf{The cancer biorobots Jupyter notebook on nanoHUB}.$



Moreover, it may be useful to provide additional control over styling and placement by a separate style.xml or similar file, or by an external cascading style sheet. We will explore these options in the future, with the aim of separating as much GUI specification and styling from the original scientific application as possible. Such a decoupling would make it easier for scientific developers to continue refining their scientific codes without worrying about impact on the GUI, and without undue encumbrance by non-scientific annotations.

Also, we currently provide just 2-D visualizations of (spatial) data. In the near future, we will provide visualizations of 3-D models and welcome suggestions from the community.

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