

Xml2jupyter: Mapping parameters between XML and Jupyter widgets

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Software

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Summary

Jupyter Notebooks (Kluyver et al. 2016) 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 (hopefully in some standard format) to provide parameter values. Xml2jupyter is a Python package that bridges this gap. 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. Users modify parameter values via the widgets and the values are written to the XML configuration file. Xml2jupyter has been tested using the PhysiCell (Ghaffarizadeh et al. 2018) simulation software and will be used by students for classroom and research projects.

A PhysiCell configuration file defines model-specific user parameters in XML. Each parameter element consists of its name with attributes, defining its data *type* and *units* (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.

To illustrate, we show the following contrived XML example, containing each of the four allowed data types:

When we map this into Jupyter widgets, we have the following that shows the rendered widgets (left). The middle snapshot shows the result after the user does a single delta step (up) on the radius (note the step size of 10) and the right snapshot after the user single steps the threads (note the step size of 1).



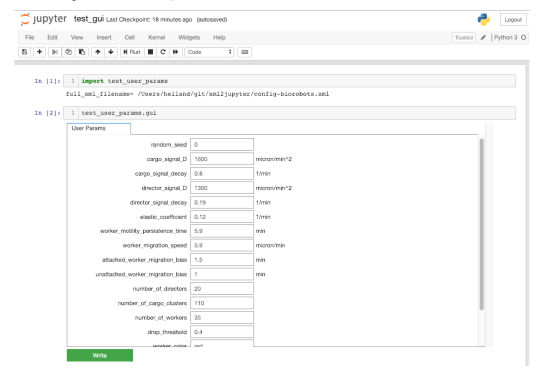


In another example, we use an XML configuration file from the *biorobots* sample project included with PhysiCell:

A workflow to demonstrate this project is the following:

• python xml2jupyter.py <config_file>.xml

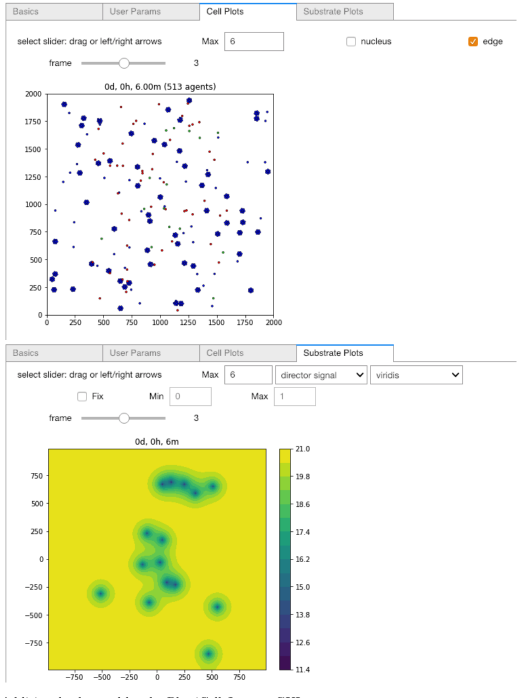
A user would execute a Python script, xml2jupyter.py, that parses this XML and generates a Python module containing the Jupyter widgets, together with methods to populate their values from the XML and write their values back to the XML. This can be displayed as a very simple GUI in a Jupyter notebook to provide the mapping to/from the XML parameter values, as shown below:





PhysiCell Jupyter GUI

Our ultimate goal is to generate a functional GUI for PhysiCell users. Xml2jupyter provides one important piece of this - dynamically generating widgets for custom user parameters for a model. With the addition of static components (tabs) of the GUI, a user can also visualize output results from simulations. This additional functionality requires modules not available in the Python standard library, e.g., Matplotlib (Hunter 2007) to display plots, and SciPy to parse PhysiCell output data. We provide instructions for installing these additional dependencies on github link.



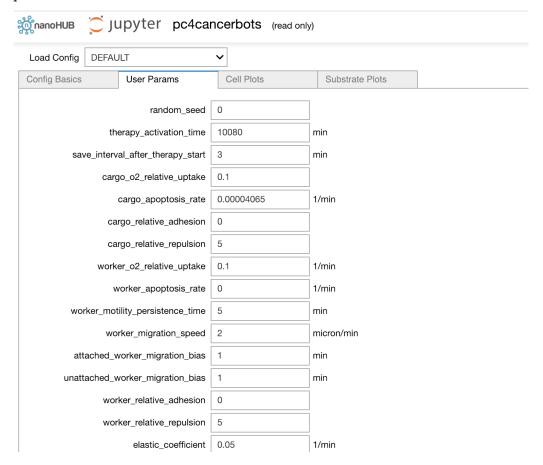
Additional tabs used by the PhysiCell Jupyter GUI.



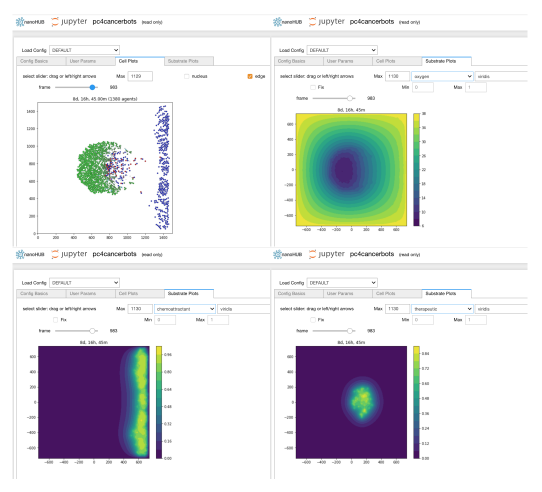
Extensions and Discussion

We hope others will be inspired to extend the idea of this project to other text-based configuration files. XML is only one of several data-interchange formats. It just happens to be the one of interest to us for PhysiCell.

Xml2jupyter has helped us port PhysiCell-related Jupyter tools to nanoHUB (Krishna et al. 2013), a scientific cloud for nanoscience education and research that includes running interactive or batch simulations. For example, we show screen shots from our pc4cancerbots tool running on nanoHUB, where the *User Params* tab has been generated using the xml2jupyter.py script. Readers can create an account on nanoHUB and run pc4cancerbots for themselves.







We welcome suggestions and, better yet, contributions (via pull requests) to xml2jupyter. For example, currently, we simply arrange the generated widgets in a single column. It may be preferable to allow a matrix-style layout, using additional attributes in the XML. Also, it might be nice to provide more informative tooltip text for widgets. # Acknowledgements

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