PathVisio-Validator: a rule-based validation plugin for graphical pathway notations

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ABSTRACT

Purpose: The PathVisio-Validator plugin aims to simplify the task of producing biological pathway diagrams that follow graphical standardized notations, such as Molecular Interaction Maps or the Systems Biology Graphical Notation. This plugin assists in the creation of pathway diagrams by ensuring correct usage of a notation, and thereby reducing ambiguity when diagrams are shared among biologists. Rulesets, needed in the validation process, can be generated for any graphical notation that a developer desires, using either Schematron or Groovy. The plugin also provides support for filtering validation results, validating on a subset of rules, and distinguishing errors and warnings.

Availability: The PathVisio-Validator plugin works with versions of PathVisio 2.0.11 and later on Windows, Mac OS X and Linux. The plugin along with the instructions, example rulesets for Groovy and Schematron, and Java source code can be downloaded at http://pathvisio.org/wiki/PathVisioValidatorHelp. The software is developed under the open-source Apache 2.0 License and is freely available for both commercial and academic use.

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1 INTRODUCTION

Biological researchers use pathway diagrams as the medium to relay information about interactions among biological entities. Graphical notations like the Molecular Interaction Map (MIM) notation and the Systems Biology Graphical Notation (SBGN) promote the creation of unambiguous biological pathway diagrams through the use of well-defined rules for syntax. The specifications for these notations are long and technical, and this is a barrier to the wider usage of these notations by researchers. This issue presents a clear need not only for tools that simplify the usage of these graphical notations, but also tools that guide users in their correct usage.

PathVisio is an open-source Java-based tool for creating and editing biological pathways and linking diagram elements to external bioinformatics databases (van Iersel et al., 2008). By itself, PathVisio does not compel users to create diagrams using any particular notation; support for specific notations is added through the use of plugins, such as the MIM plugin (Luna et al., 2011b) or the SBGN plugin, which is under development (http://www.pathvisio.org/wiki/SbgnPluginHelp). Given this open nature of PathVisio, we have developed a validation framework similar to what developers are accustomed to in source code editors that report syntax errors. This accommodates the validation of pathway diagrams against any notation that PathVisio can draw with just the click of a button.

There are several projects related to standardized formats for computational biology that have associated validators, including well-known formats like BioPAX (Demir et al., 2010) and SBML (Hucka et al., 2003); there is also validation capability in BCML, a SBGN-compliant format (Beltrame et al., 2011), and SBGN-ED (Czauderna et al., 2010), an SBGN diagram editor. A key difference between SBGN-ED and the plugin presented here is the extensible support for other notations in the latter, which is especially important for the PathVisio community since its users may be using various graphical notations for diverse purposes. A second difference is that rulesets (specifically, Schematron rulesets) created for use with this plugin can be reused in other software projects. The use of Schematron is a feature that this plugin shares with some generic XML editors with validation capability, such as XMLSpy (http://xml-tool.com/ValidatorBuddy.htm).

2 IMPLEMENTATION

The validation of pathways using the plugin is performed based on a ruleset (a collection of rules). The plugin allows users to use their own custom-created rulesets or standard rulesets by notation developers. Described below are the main features of the PathVisio-Validator plugin and its user interface:

Plugin interface and current notation support: the interface for the plugin exists under the ‘Validator’ tab of PathVisio as shown in Figure 1. At the bottom of the tab are buttons for loading a ruleset and validating the current diagram. In order to validate a pathway diagram, the user simply needs to choose a ruleset and press the ‘Validate’ button. Validation messages for the current pathway diagram will then appear in the side panel as shown in Figure 1.
Validation against a Groovy ruleset, consumes less time than its Validation Report Language. This validation report is parsed and is completed. currently in development as a separate project. Validation support for biological pathway diagrams. SBGN support for PathVisio is the WikiPathways database (http://wikipathways.org), a repository also provide a ruleset with rules used for the curation of diagrams on plugin supports the MIM notation through a Schematron ruleset. We can ignore particular messages or types of messages. Currently, the or both, using a drop-down menu. Users also have an option to select with all the rules in a ruleset or only those under a specific group, rulesets may be put into groups. Thus, a user can validate a diagram with all the rules in a ruleset or only those under a specific group, thereby making it more selective. Schematron supports rule groups using the ‘phase’ XML tag with one or more ‘active’ rules as child XML nodes. Groovy supports rule groups through a specific method that defines rule groups and their associated rules.

3 CONCLUSION

The PathVisio-Validator plugin has been developed to enhance the quality of pathways created in PathVisio by assisting biologists in drawing pathways according to specific graphical notations. Validation of this nature allows researchers to be more confident that their diagrams will not be ambiguous to readers. The extensible nature of the rulesets should allow it to adapt to preferences of PathVisio users, and the support for different ruleset formats (i.e. Schematron and Groovy) should allow users to create rulesets based on their priorities. Moreover, this plugin will also encourage the users of PathVisio, especially the beginners to create pathways based on specific notations, such as MIM, helping to promote the adoption of such standards. In the future, we plan to integrate this validation framework into WikiPathways to provide automated diagram validation (Pico et al., 2008), so that all uploaded/updated pathways are validated against common rules. Presented here is a narrow use case of the validation of graphical notations, but the ideas presented here can be extended to other biological standards written in XML-based formats. These ideas are of use in the wider standards community in providing validation support in multiple programming languages.

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