

Operator 0.8 released: a new Sechemtic user script



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Semweb, Chemistry

Abstract

Mike released Operator 0.8, which picks up RDF (RDFa en eRDF) from HTML pages, and adds actions to it. I blogged earlier about the beta and wrote a script for it for chemical RDFa . At this moment, Chemical blogspace and RDF for Molecular Space (see this blog) are using chemical RDFa to semantically markup molecular information.

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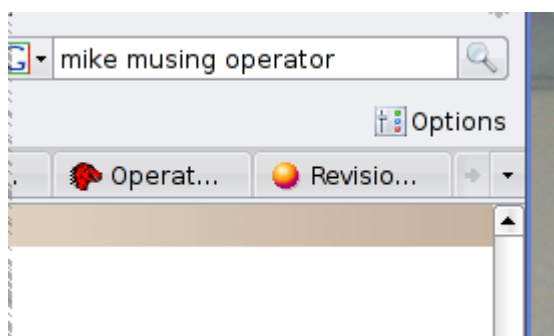
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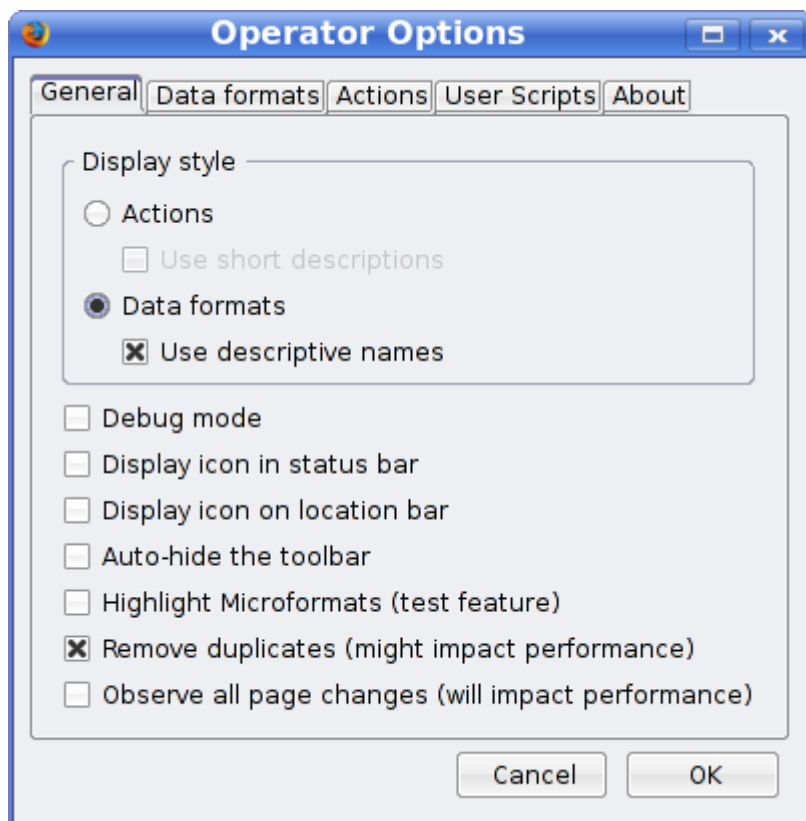
Mike released [Operator 0.8](#), which picks up RDF (RDFa en eRDF) from HTML pages, and adds actions to it. I [blogged earlier about the beta](#) and wrote a script for it for [chemical RDFa](#) . At this moment, [Chemical blogspace](#) and [RDF for Molecular Space](#) (see [this blog](#)) are using chemical RDFa to semantically markup molecular information.

The new Operator release ([download](#)) has one notable API change: it now uses “RDF” as key for semantic information; the add-on now supports eRDF too. So, when installing or updating to version 0.8, you also need to update the Sechemtic user script to [version 1.1 or better](#).

Installing Operator scripts is a bit more work than Greasemonkey userscripts. Save the script to your home directory, or any other place you can easily find on the hard disk. After installing the Operator add-on, click the *Options* button:

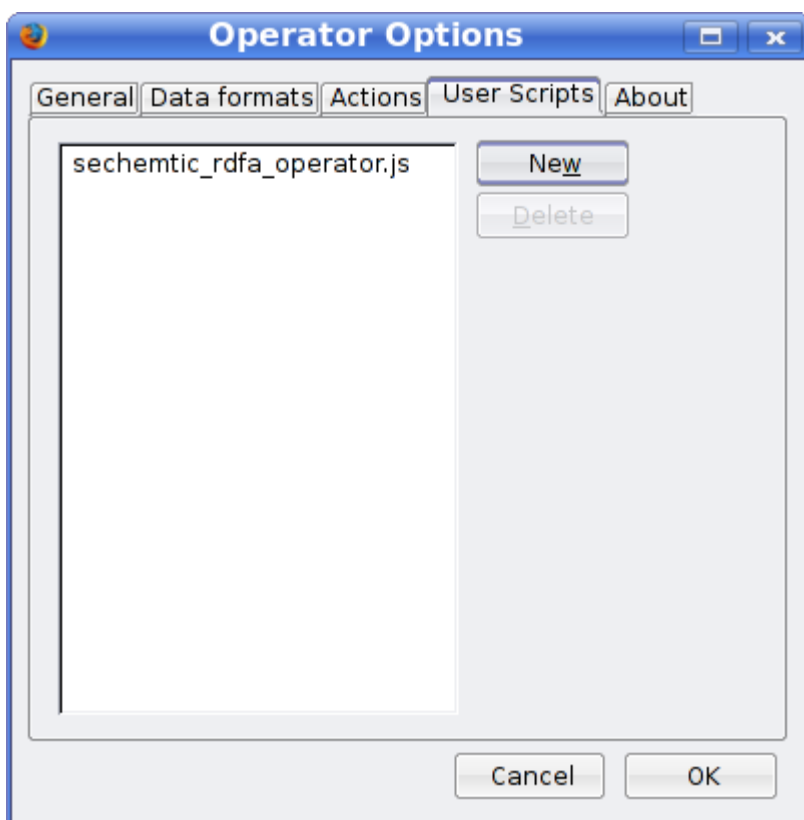


For the RDFa script to work, you need to make sure that the *Display style* is set to *Data formats*:



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Then you can go to the *User Scripts* tab, and use the *New* button to add the script you downloaded and saved to your hard disk earlier:



Then, after rebooting Firefox (looks like MS-Windows :()), you can go to Chemical blogspace and [look up molecules](#), and see output like that described in [RDFa Operator in action on Cb](#).