

Scripting JChemPaint

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Abstract

Today and tomorrow, Stefan, Gilleain, Arvid and I are having a JChemPaint Developers Workshop in Uppsala, to sprint the development of JChemPaint3, for which Niels layed out the foundation already a long time ago.

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Today and tomorrow, Stefan, [Gilleain](#), Arvid and I are having a [JChemPaint Developers Workshop](#) in Uppsala, to sprint the development of JChemPaint3, for which [Niels](#) laid out the foundation already a long time ago.

Gilleain and Arvid are merging their branches into a [single code base](#), while Stefan is working on the [Swing application and applet](#). The Bioclipse SWT-based widget is being developed for [Bioclipse2](#).

The new design separates widget/graphics toolkit specifics from the chemical drawing and editing logic. Regarding the editing functionality, this basically comes down to have a semantically meaningful edit API. This allows us to convert both Swing and SWT mouse events into things like `addAtom("C", atom)`, which would add a carbon to an already existing *atom*. However, without too much phantasy, it allows adding a scripting language. This is what I have been working on. Right now, the following API is available from the Bioclipse2 JavaScript console (via the *jcp* namespace, in random order):

- `ICDKMolecule jcp.getModel()`
- `IAtom getClosestAtom(Point2d)`
- `setModel(ICDKMolecule)` (*for really fancy things*)
- `removeAtom(IAtom)`
- `IBond getClosestBond(Point2d)`
- `updateView()` (*all edit command issue this automatically*)
- `addAtom(String,Point2d)`
- `addAtom(String,IAtom)` (*which works out coordinates automatically*)
- `Point2d newPoint2d(double,double)`
- `updateImplicitHydrogenCounts()`
- `moveTo(IAtom, Point2d)`
- `setSymbol(IAtom,String)`
- `setCharge(IAtom,int)`
- `setMassNumber(IAtom,int)`
- `addBond(IAtom,IAtom)`
- `moveTo(IBond,Point2d)`
- `setOrder(IBond,IBond.Order)`
- `setWedgeType(IBond,int)`
- `IBond.Order getOrder(int)`
- `zap()` (*sort of `sudo rm -Rf /*`*)
- `cleanup()` (*calculate 2D coordinates from scratch*)
- `addRing(IAtom,int)`
- `addPhenyl(IAtom)`

This API (many more method will follow) is not really aimed at the end user, who will simply point and click. The goal of this scripting language is, at least at this moment, to test the underlying implementation using Bioclipse. Future applications, however, may include simple scripts which use some logic to convert the editor content. For example, replacing a t-butyl

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fragment into a pseudo atom “t-Bu”. The key thing to remember, is that this will allow Bioclipse to have non-CDK-based programs act on the JChemPaint editor content (e.g. using `getModel()` and `setModel(ICDKMolecule)`). More on that later.

A simple script could look like: Or, as screenshot:

The screenshot displays the Bioclipse application window. The main canvas shows a chemical structure of a molecule with two benzene rings connected by a methylene group, which is further connected to a carbonyl group. The left sidebar, titled 'Bioclipse Explorer', shows a file tree with folders like 'CDK Demo Scripts', 'DemoData', 'pcore-test', and 'PeptideMimetics'. The 'PeptideMimetics' folder is expanded, showing files such as '10.cml', '3.cml', 'spectrum.10.cml', 'spectrum.3.cml', 'submission.10.cml', 'submission.3.cml', 'submission.4.cml', 'submission.5.cml', 'thesis.bib', and 'typing.js'. The bottom panel contains a 'JavaScript Console' with the following code and output:

```
js> oxygen = jcp.getModel().getAtomContainer().getAtom(11)
Atom(16046778, S:O, H:0, 2D:[(274.0576230702553, 125.52238289382183)],
AtomType(16046778, FC:O, NC:O, EV:0, Isotope(16046778, MN:16, Element(16046778,
S:O, ID:a12))))

js> jcp.addAtom("C", oxygen)
js> newCarbon = jcp.getModel().getAtomContainer().getAtom(12)
Atom(21241643, S:C, H:3, 2D:[(243.37358553521833, 144.35028348173765)],
AtomType(21241643, FC:O, NC:O, EV:0, Isotope(21241643, Element(21241643, S:C,
ID:a13))))

js> jcp.addPhenyl(newCarbon)
js> |
```