

Resource Summary Report

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Chemtool

RRID:SCR_023992

Type: Tool

Proper Citation

Chemtool (RRID:SCR_023992)

Resource Information

URL: <http://ruby.chemie.uni-freiburg.de/~martin/chemtool/>

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Description: Software tool for drawing organic molecules. Runs under the X Window System using the GTK widget set. Used for drawing chemical structures on Linux and Unix systems using the GTK toolkit under X11.

Synonyms: chemtool

Resource Type: software application

Defining Citation: [DOI:10.1002/nadc.20010491112](https://doi.org/10.1002/nadc.20010491112)

Keywords: drawing organic molecules, drawing chemical structures,

Availability: Free, Available for download, Freely available

Resource Name: Chemtool

Resource ID: SCR_023992

Alternate URLs: <https://sources.debian.org/src/chemtool/>

Record Creation Time: 20230824T050211+0000

Record Last Update: 20260725T191035+0000

Ratings and Alerts

No rating or validation information has been found for Chemtool.

No alerts have been found for Chemtool.

Data and Source Information

Source: [SciCrunch Registry](#)

Usage and Citation Metrics

We have not found any literature mentions for this resource.