
Subject: C=C double bonds/search by extended stereochemistry

Posted by [nbehrnd](#) on Tue, 30 Nov 2021 16:11:02 GMT

[View Forum Message](#) <> [Reply to Message](#)

Hello Thomas,

my understanding of DataWarrior's manual is that the program internally stores the molecules in a format capable to discern and (if selecting .sdf v3000) export of double bonds including those without explicit assignment of (E)/(Z) configuration. As anticipated, the SMILES strings

C/C=C/C

C/C=C\C

CC=CC

yield three different representations of but-2-ene.

For a small test set (cf. `double_bond.dwar` and the screen photos in the .zip archive attached), I aimed to use the features of extended stereochemistry as a criterion to retain a sub set (Chemistry -> From Chemical Structure -> Add Substructure Count), including the unassigned double bond visualized as a crossed double bond. So far, the attempts failed.

As anticipated, the search pattern of (cis)-butene, retained both (cis)-butene and cyclohexene and withdrew the oxime. However -- contrasting to anticipation -- (trans)-butene and the unassigned butene equally were retained. Does the search for, e.g. crossed double bonds, require a particular instruction?

Version info: DataWarrior 5.5.0 for Linux in Debian 12/bookworm (branch testing), including the update as «development version» as fetched on Sunday, 2021-11-28.

Norwid

File Attachments

1) [double_bonds.dwar](#), downloaded 156 times

2) [double_bond_documentation.zip](#), downloaded 157 times
