Subject: Re: ChemAxon calculated properties

Posted by chemist on Tue, 17 Jan 2017 09:45:31 GMT

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Thanks for creating really great software!

I have been struggling with Chemaxon calculated properties. I also have this free academic lisence but enabling these chemaxon properties seems quite difficult.

What I have tried:

- 1. Go to chemaxon directory
- 2. Searh .jar
- 3. copy all of them to new folder
- 4. extract all .jars by ticking [x]"Extract archives to subfolders"
- 5. zip all to capka.zip
- 6. rename -> capka.jar
- 7. move to datawarrior.exe location

I have also lisence file in correct location.

I just started to think if in the academic version there is some limitation to batch calculation of pKa values.

I have lisence for following products:

Marvin Applets

Marvin Beans

Instant JChem

JChem Base

Standardizer

Screen

Reactor

JKlustor

Metabolizer

Markush Search

Protonation Plugin Group

Partitioning Plugin Group

Isomers Plugin Group

Markush Enumeration Plugin

Structure to Name Plugin

Name to Structure

JChem for Excel

Structure Search

IUPAC naming plugin

Web Services Server

Structure Checker

Predictor Plugin

MCES

3D Screen
Molecular Descriptors
Instant JChem VIZ
ECFP/FCFP
Document to Structure
NMR Predictor
JChem for Office
Markush Editor
Solubility Plugin
Structural Calculations