Minnesota Solvation (MNSOL) Database

An extensive and vetted database of free energies of solvation.



Applications

- Chemical engineering
- Drug development
- Developing solubility and bioavailability predictive models

Overview

The Minnesota Solvation Database consists of a collection of 3037 experimental free energies of solvation or transfer free energies for 790 unique solutes in 92 solvents (including water) and gas-phase M06-2X/MG3S optimized molecular geometries in Cartesian coordinates for the corresponding solutes. All of the 790 solutes in this database (541 neutrals and 249 singly-charged ions) contain at most the following elements: H, C, N, O, F, Si, P, S, Cl, Br, and I. User manual with detailed description of the dataset and example calculations is included.

Phase of Development

Database -- version 2012 (MNSol-v2012) -- available for download.

External Link (chem.umn.edu)

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Publications

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Category

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Software & IT/Databases
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<u>Universal Solvation Model Based on Solute Electron Density and on a Continuum Model of</u> <u>the Solvent Defined by the Bulk Dielectric Constant and Atomic Surface Tensions</u>

Journal of Physical Chemistry B, 113, 6378–6396 (2009)

Generalized Born Solvation Model SM12

Journal of Chemical Theory and Computation, 9, 609-620 (2013)

External Links

AMSOL 7.1: Software to Calculate Free Energies of Solvation

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- Non-profit organizations must submit a request here.
- Commercial organizations must purchase the online license using the link provided on the right panel.