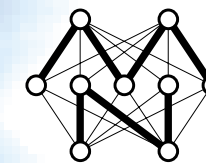


# Accessing the Three-Dimensional World of Molecules

## 3D Structure Generator CORINA™

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Molecular Networks

### Overview

#### CORINA™ - Fast and Efficient Generation of High-Quality Three-Dimensional Molecular Models

The three-dimensional structure of a molecule is closely related to a large variety of chemical, physical and biological properties. The need for computer generated 3D molecular structures has clearly been recognized in drug design and related disciplines. Since the number of experimentally determined molecular geometries is limited - about 250.000 X-ray structures in the Cambridge Crystallographic Database compared to approximately 22 millions of known compounds - one needs a method to predict 3D atomic coordinates of molecules fast and efficiently.

The rule- and databased program system CORINA™ (COoRdINates)<sup>[1]</sup> automatically generates three-dimensional atomic coordinates from the constitution of a molecule as expressed by a connection table or linear string. It can handle massive volumes of structures such as those in company databases and combinatorial chemistry experiments. The scheme on the right side shows the general principles of CORINA™.

#### Features

- Applicable to the entire range of classical organic chemistry, including stable molecules, radicals, reactive intermediates, multi-fragment records (e.g. salts), fused rings, spiro, polycyclic, macrocyclic and polymacrocyclic systems
- Generation of low-energy conformations
- Consideration of stereochemical information
- Processing of structures with atoms having up to six neighbors
- Conversion of organo-metallic complexes
- No upper limits to the size of the molecules or size of ring systems<sup>[2]</sup>
- Generation of multiple ring conformations for rings consisting of less than ten atoms
- Processing of a large variety of structural data file formats (e.g. MDL SDFfile, MDL RDFfile, SYBYL MOLfile and MOL2, SMILES linear code and PDB format)
- Interface to the flexible ligand docking program FlexX™ for the generation of multiple ring conformations during the docking process
- Several options to influence the generation process, e.g. addition of implicitly given hydrogen atoms in the input file, neutralization of charged molecules, or orientation of the 3D structures according to their moments of inertia
- Available for common UNIX platforms (SGI, Sun, DEC, and Linux) as well as for MS Windows systems

#### Performance

- High-quality molecular models: Lowest RMS deviations from published X-ray structures among all commercially available model builders
- Fast: Conversion in less than one second for medium-sized organic molecules on a common UNIX workstation
- Robust: Conversion of the open part of the NCI Database with a rate of more than 99% without intervention and program crash
- General: Processing of a database with over six million compounds and a conversion rate of more than 99%

#### References

- [1] J. Sadowski, J. Gasteiger *Chem. Reviews* **1993**, *93*, 2567.
- [2] Schönberger, H.; Schwab, C.H.; Hirsch, A.; Gasteiger, J. *J. Mol. Model.* **2000**, *6*, 379-395.
- [3] J. Sadowski, J. Gasteiger, G. Klebe *J. Chem. Inf. Comput. Sci.* **1994**, *34*, 1000.

