Journal of Computer-Aided Molecular Design

Incorporating Perspectives in Drug Discovery and Design

Editors-in-Chief: F. Gago; T.R. Stouch; R. Lewis

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The Journal of Computer-Aided Molecular Design provides a form for disseminating information on both the theory and the application of computer-based methods in the analysis and design of molecules. The scope of the journal encompasses papers which report new and original research and applications in the following areas: theoretical chemistry;

- computational chemistry;
- computer and molecular graphics;
- molecular modeling;
- protein engineering;
- drug design;
- expert systems;
- general structure-property relationships;
- molecular dynamics;
- chemical database development and usage.

Contributions on computer-aided molecular modeling studies in pharmaceutical, polymer, materials and surface sciences, as well as other molecular-based disciplines, are particularly welcome.

As of Volume 15, a number of issues per volume will be dedicated to Perspectives in Drug Discovery and Design and from time to time overviews will be included in `regular' JCAMD issues.

Impact Factor: 2.356 (2017), Journal Citation Reports®

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