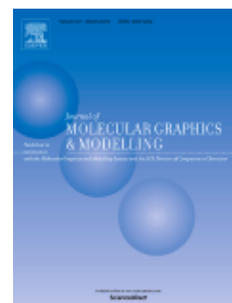




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The *Journal of Molecular Graphics and Modelling* is devoted to the publication of papers on the uses of **computers** in theoretical investigations of molecular structure, function, interaction, and design. The scope of the journal includes all aspects of **molecular modeling** and **computational chemistry**, including, for instance, the study of molecular shape and properties, molecular simulations, protein and polymer engineering, drug design, materials design, structure-activity and structure-property relationships, database mining, and compound library design.

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The *Journal of Molecular Graphics and Modelling* is devoted to the publication of papers on the uses of computers in theoretical investigations of molecular structure, function, interaction, and design. The scope of the journal includes all aspects of molecular modeling and computational chemistry, including, for instance, the study of molecular shape and properties, molecular simulations, protein and polymer engineering, drug design, materials design, structure-activity and structure-property relationships, database mining, and compound library design.

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