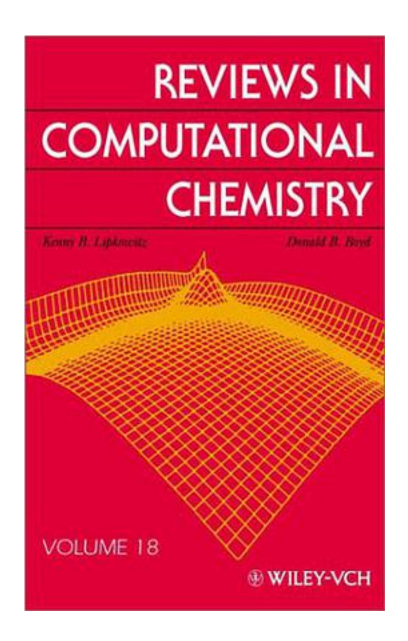
Reviews in Computational Chemistry, Reviews in Computational Chemistry, Volume 18



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This volume, like those prior to it, features chapters by experts in various fields of computational chemistry. Topics covered in Volume 18 include molecular modeling, computer-assisted molecular design (camd), quantum chemistry, molecular mechanics and dynamics, and quantitative structure-activity relationships (qsar).
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