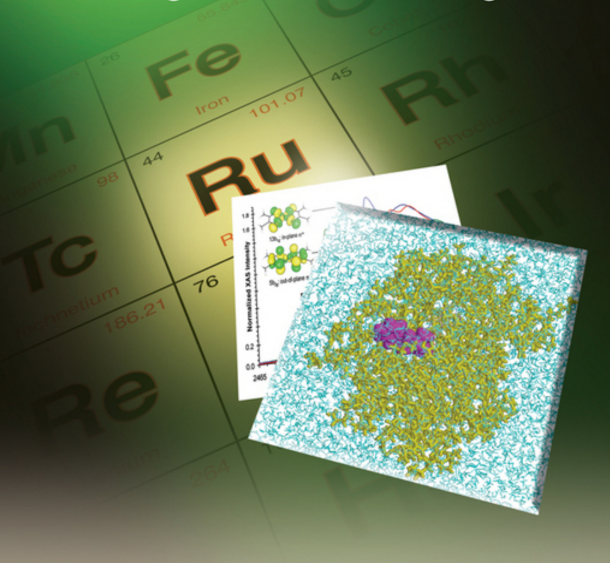
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# COMPUTATIONAL Inorganic and Bioinorganic Chemistry

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Frenking • Calculation of Bonding Properties

Richard A. Columbia University, New York, NY, USA

• Modeling Metalloenzymes with Density Functional and Mixed

Quantum Mechanical/ Molecular Mechanical (QM/MM)

Calculations: Progress and Challenges

Filipp Furche University of California, Irvine, CA, USA

• Approximate Density Functionals: Which Should I Choose?