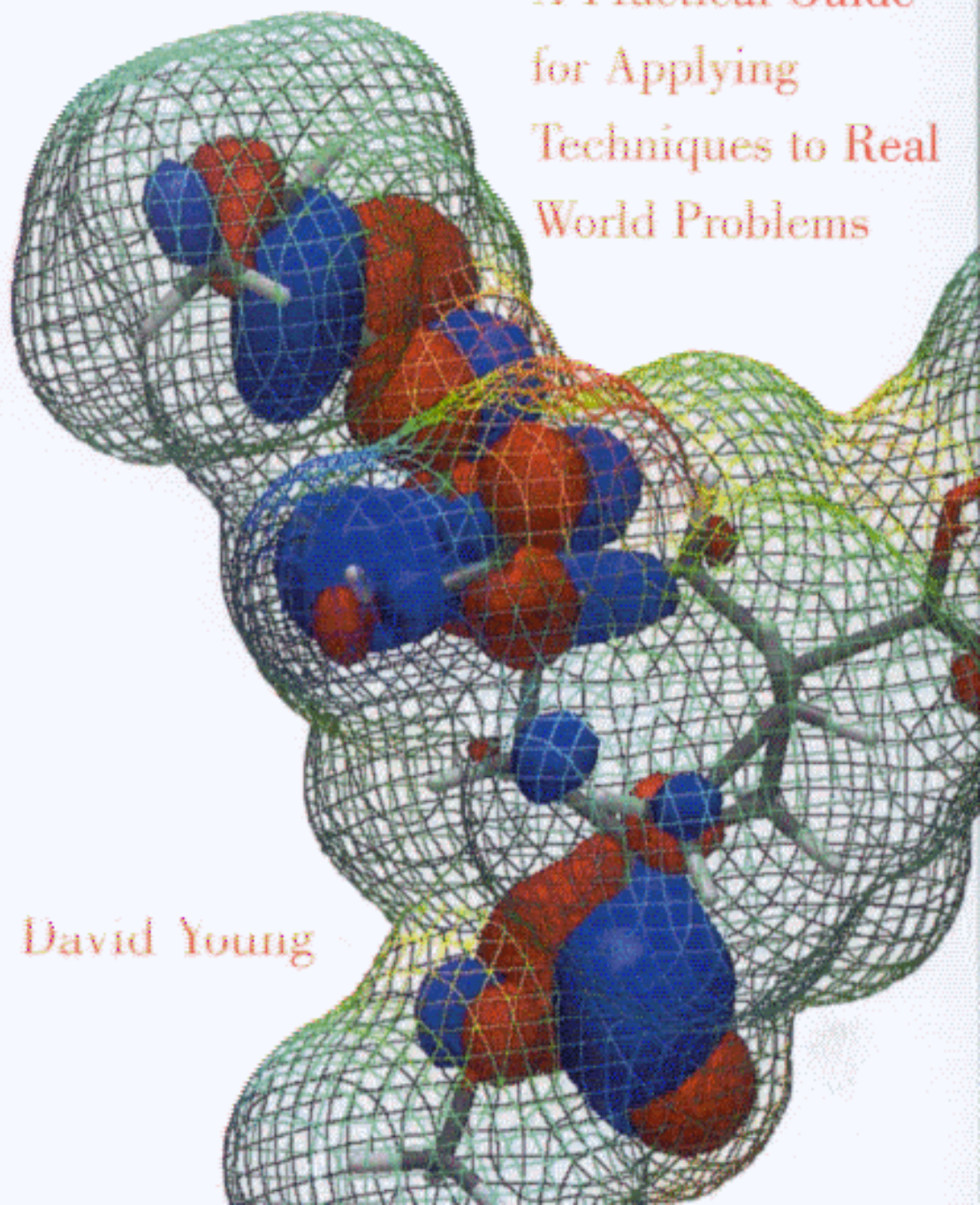


Computational Chemistry

A Practical Guide
for Applying
Techniques to Real
World Problems

David Young



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