TINKER - Software Tools for Molecular Design

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TINKER is a modular program package for molecular mechanics-based potential energy calculations, geometry optimization, molecular dynamics simulation, distance geometry and structural analysis.

Selected References for the TINKER Package:

- (1) Y. Shi, et al., J. Chem. Theory Comput., **9**, 4046-4063 (2013)
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Prof. Jay W. Ponder

Department of Chemistry, Box 1134 Phone: 1-314-9354275 Washington University in Saint Louis Fax: 1-314-9354481

One Brookings Drive

Saint Louis, MO 63130 U.S.A.

Email: ponder@dasher.wustl.edu