1. Record Nr. UNISA996466708703316 Computer simulations in condensed matter systems : from materials to **Titolo** chemical biology / / edited by Mauro Ferrario, Giovanni Ciccotti, Kurt Binder Berlin, Germany; New York, New York:,: Springer-Verlag,, [2006] Pubbl/distr/stampa ©2006 **ISBN** 1-280-80517-X 9786610805174 3-540-35273-2 Edizione [1st ed. 2006.] Descrizione fisica 1 online resource (715 p.) Lecture Notes in Physics, , 0075-8450 ; ; 703 Collana Disciplina 530.410113 Soggetti Condensed matter - Computer simulation Lingua di pubblicazione Inglese **Formato** Materiale a stampa Livello bibliografico Monografia Proceedings of a school held at the Ettore Majorana Foundation and Note generali Center for Scientific Culture, Erice, Sicily in July 2005. Nota di bibliografia Includes bibliographical references and indexes. Nota di contenuto Introduction: Condensed Matter Theory by Computer Simulation --Transition Path Sampling Methods -- Sampling Kinetic Protein Folding Pathways using All-Atom Models -- Calculation of Classical Trajectories with Boundary Value Formulation -- Transition Path Theory -- Multiscale Modelling in Molecular Dynamics: Biomolecular Conformations as Metastable States -- Transport Coefficients of Quantum-Classical Systems -- Linearized Path Integral Methods for Quantum Time Correlation Functions -- Ensemble Optimization Techniques for Classical and Quantum Systems -- The Coupled Electron-Ion Monte Carlo Method -- Path Resummations and the Fermion Sign Problem -- to Cluster Monte Carlo Algorithms -- Generic Sampling Strategies for Monte Carlo Simulation of Phase Behaviour --Simulation Techniques for Calculating Free Energies -- Waste-Recycling Monte Carlo -- Equilibrium Statistical Mechanics, Non-Hamiltonian Molecular Dynamics, and Novel Applications from Resonance-Free Timesteps to Adiabatic Free Energy Dynamics --Simulating Charged Systems with ESPResSo -- Density Functional Theory Based Ab Initio Molecular Dynamics Using the Car-Parrinello

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