Record Nr.	UNINA9910298580803321
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Titolo	Computational Materials Science : From Ab Initio to Monte Carlo Methods / / by Kaoru Ohno, Keivan Esfarjani, Yoshiyuki Kawazoe
Pubbl/distr/stampa	Berlin, Heidelberg : , : Springer Berlin Heidelberg : , : Imprint : Springer, , 2018
ISBN	3-662-56542-0
Edizione	[2nd ed. 2018.]
Descrizione fisica	1 online resource (XII, 427 p.)
Disciplina	620.11011
Soggetti	Optical materials Electronic materials Physics Chemistry, Physical and theoretical Nanotechnology Solid state physics Optical and Electronic Materials Numerical and Computational Physics, Simulation Theoretical and Computational Chemistry Solid State Physics
Lingua di pubblicazione	Inglese
Formato	Materiale a stampa
Livello bibliografico	Monografia
Nota di contenuto	Ab-Initio Methods Tight-Binding Methods Empirical Methods and Coarse-Graining Monte Carlo Methods Quantum Monte Carlo (QMC) Methods.
Sommario/riassunto	This textbook introduces modern techniques based on computer simulation to study materials science. It starts from first principles calculations enabling to calculate the physical and chemical properties by solving a many-body Schroedinger equation with Coulomb forces. For the exchange-correlation term, the local density approximation is usually applied. After the introduction of the first principles treatment, tight-binding and classical potential methods are briefly introduced to indicate how one can increase the number of atoms in the system. In the second half of the book, Monte Carlo simulation is discussed in detail. Problems and solutions are provided to facilitate understanding.

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Readers will gain sufficient knowledge to begin theoretical studies in modern materials research. This second edition includes a lot of recent theoretical techniques in materials research. With the computers power now available, it is possible to use these numerical techniques to study various physical and chemical properties of complex materials from first principles. The new edition also covers empirical methods, such as tight-binding and molecular dynamics.