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Autore	Raabe Dierk
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Nota di contenuto	Front Matter Introduction and Fundamentals. Introduction Modeling and Simulation in Materials Science Fundamentals and Solution of Differential Equations Simulation Techniques at the NanoscopicMicroscopic Scale. Fundamentals Statistical Mechanics in Atomic-Scale Simulations Monte Carlo Simulation and Integration Molecular Dynamics Simulation Techniques at the MicroscopicMesoscopic Scale. Introduction Discrete Dislocation Statics and Dynamics: Sections 9.19.3 Discrete Dislocation Statics and Dynamics: Sections 9.49.8 GinzburgLandau-Type Phase Field Kinetic Models Cellular Automata Mesoscale Kinetic Monte Carlo and Potts Models Geometrical and Component Models Topological Network and Vertex Models Simulation Techniques at the MesoscopicMacroscopic Scale. Introduction Finite Element and Difference Methods at the Meso Macroscale Polycrystal Elasticity and Plasticity Models Integrated Modeling and Simulation. Fundamentals Space and Time Scales in Microstructure Simulation Appendices. Appendix A: General Reading Appendix B: Computer Classification Appendix C: Advanced Empirical Methods Appendix D: Percolation Theory References Index.

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Modeling and simulation play an ever increasing role in the development and optimization of materials. Computational Materials Science presents the most important approaches in this new interdisciplinary field of materials science and engineering. The reader will learn to assess which numerical method is appropriate for performing simulations at the various microstructural levels and how they can be coupled. This book addresses graduate students and professionals in materials science and engineering as well as materialsoriented physicists and mechanical engineers.