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Titolo	Introduction to modern methods of quantum many-body theory and their applications [[electronic resource] /] / editors, Adelchi Fabrocini, Stefano Fantoni, Eckhard Krotscheck
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Descrizione fisica	1 online resource (428 p.)
Collana	Series on advances in quantum many-body theory ; ; v. 7
Altri autori (Persone)	FabrociniA FantoniS (Stefano) KrotscheckEckhard
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Note generali	Lecture notes of the second European Summer School on Microscopic Quantum Many-Body Theories and their Applications, held in Miramare, Trieste, Italy, on September 3-14, 2001.
Nota di bibliografia	Includes bibliographical references and indexes.
Nota di contenuto	CONTENTS ; PREFACE ; Chapter 1 DENSITY FUNCTIONAL THEORY ; 1. Introduction ; 1.1. Units and notation ; 1.2. Hartree-Fock theory ; 1.3. Homogeneous electron gas ; 1.3.1. Free electrons ; 1.3.2. Exchange energy ; 2. What is density functional theory? ; 2.1. Hohenberg-Kohn theorem ; 2.2. A simple example: the Thomas-Fermi theory ; 2.2.1. Variational equation of Thomas-Fermi theory ; 2.2.2. Thomas-Fermi atom ; 2.2.3. An example ; 3. Kohn-Sham theory ; 3.1. Local density approximation ; 3.2. Spin and the local spin density approximation ; 3.3. The generalized gradient approximation ; 4. Numerical methods for the Kohn-Sham equation ; 4.0.1. Exact exchange ; 4.0.2. O(N) methods ; 5. Some applications and limitations of DFT ; 5.1. Two examples of condensed matter ;

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 polarizability ; 7.4. Dielectric function
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 ; 8.3. Sternheimer method ; 8.4. Real time method
 ; 9. Applications of TDDFT ; 9.1. Simple metal
 clusters ; 9.2. Carbon structures
 ; 9.3. Diamond ; 9.4. Other applications
 9.5. Limitations

Sommario/riassunto

This invaluable book contains pedagogical articles on the dominant nonstochastic methods of microscopic many-body theories - the methods of density functional theory, coupled cluster theory, and correlated basis functions - in their widest sense. Other articles introduce students to applications of these methods in front-line research, such as Bose-Einstein condensates, the nuclear many-body problem, and the dynamics of quantum liquids. These keynote articles are supplemented by experimental reviews on intimately connected topics that are of current relevance. The book addresses the striking
