1. Record Nr. UNINA9910830592703321 Autore Helgaker Trygve Titolo Molecular electronic-structure theory / / Trygve Helgaker, Poul Jorgensen, Jeppe Olsen Pubbl/distr/stampa Chichester, England:,: John Wiley & Sons Ltd.,, 2012 ©2000 **ISBN** 1-119-01957-5 1-119-01955-9 1-119-01956-7 Descrizione fisica 1 online resource (940 p.) Disciplina 541.28 Soggetti Molecular structure Electronic structure Lingua di pubblicazione Inglese **Formato** Materiale a stampa Livello bibliografico Monografia Note generali Description based upon print version of record. Nota di bibliografia Includes bibliographical references and index. Nota di contenuto Cover; Title Page; Copyright; Contents; Preface; Overview; Programs used in the preparation of this book; 1 Second Quantization; 1.1 The Fock space: 1.2 Creation and annihilation operators: 1.2.1 Creation operators; 1.2.2 Annihilation operators; 1.2.3 Anticommutation relations; 1.3 Number-conserving operators; 1.3.1 Occupation-number operators; 1.3.2 The number operator; 1.3.3 Excitation operators; 1.4 The representation of one- and two-electron operators: 1.4.1 Oneelectron operators; 1.4.2 Two-electron operators; 1.4.3 The molecular electronic Hamiltonian 1.5 Products of operators in second quantization 1.5.1 Operator products; 1.5.2 The canonical commutators; 1.6 First- and secondquantization operators compared; 1.7 Density matrices; 1.7.1 The oneelectron density matrix; 1.7.2 The two-electron density matrix; 1.7.3 Density matrices in spin-orbital and coordinate representations; 1.8 Commutators and anti commutators; 1.9 Nonorthogonal spin orbitals; 1.9.1 Creation and annihilation operators; 1.9.2 One- and two-electron operators; 1.9.3 Biorthogonal operators; References; Further reading; Exercises; Solutions; 2 Spin in Second Quantization

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Sommario/riassunto

Ab initio quantum chemistry has emerged as an important tool in chemical research and is appliced to a wide variety of problems in chemistry and molecular physics. Recent developments of computational methods have enabled previously intractable chemical problems to be solved using rigorous quantum-mechanical methods. This is the first comprehensive, up-to-date and technical work to cover all the important aspects of modern molecular electronic-structure theory. Topics covered in the book include:* Second quantization with spin adaptation* Gaussian basis sets and molecular-integral evaluation* H