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Nota di bibliografia	Includes bibliographical references at the end of each chapters and index.
Nota di contenuto	The DV-Xa Molecular Orbital Calculation Method and Recent Development Algebraic Molecular Orbital Theory Analytical Expression of Molecular Integrals over Slater-Type Functions for Generating their Polynomial Expressions Atom-Atom Interaction Potential from the Gaussian Quadrature Method and Classical Molecular Dynamics Comparison of Contributions to Interatomic Interactions between Covalent and Ionic Bonds from Total Energy Calculations Total Energy Calculation by DV-Xa Method and Its Accuracy Energy Expression of the Chemical Bond between Atoms in Hydrides and Oxides and Its Application to Materials Design Comparative Study on Optical Properties of YPO4: Mn, Zr Phosphor by Experiment and Calculation Applications of DV-Xa Method for New Material Design in Dye-Sensitized Solar Cell Microscopic Approach to Water by Using the DV-Xa Method, and Some Innovative Applications Electronic Structure and Chemical Bonding of Li1.1Nb0.9O2-y as a Negative

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	Electrode Material for Lithium Secondary Batteries Chemical Bonding, Point Defects and Positron Lifetimes in FeSi2 from First-Principles Calculations Structural Analysis of Al2TiO5 at Room Temperature and at 600 °C by DV-Xa Approach (II) Wavelength of Luminescence and Energy Level Structure of Binuclear Copper(I) Complex.
Sommario/riassunto	This multi-author contributed volume contains chapters featuring the development of the DV-X method and its application to a variety of problems in Materials Science and Spectroscopy written by leaders of the respective fields. The volume contains a Foreword written by the Chairs of Japanese and Korea DV-X Societies. This book is aimed at individuals working in Quantum Chemistry.