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Theoretical Investigation on Non-covalent Interactions

Guest Editors:

Dr. Alexander S. Novikov

1. Institute of Chemistry, Saint Petersburg State University, Universitetsky pr., 26, 198504 Stary Petergof, Russia 2. Infochemistry Scientific Center, ITMO University, Lomonosova st., 9, 191002 Saint Petersburg, Russia

Prof. Dr. Longjiu Cheng

Computational chemistry, Anhui University, Hefei 230093, China

Deadline for manuscript submissions:

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Message from the Guest Editors

Dear Colleagues,

The problem of non-covalent interactions in crystals is one of the paradigms for computer modelling and theoretical studies in chemistry and related fields of knowledge (crystallography, biology, physics, mathematics, and computer science). Modern methods of data science, artificial intelligence, and quantum and computational chemistry are widely used for the investigation of nature properties of different non-covalent and various interactions (hydrogen, halogen, chalcogen, pnictogen, tetrel, and semi-coordination bonds; agosic and anagosic interactions; stacking, anion/cation-π interactions; metallophilic interactions, etc.).

Our Special Issue welcomes contributions from researchers focused on this subject to highlight and overview modern trends and attract the attention of the scientific community to the problem of theoretical investigation on non-covalent interactions

All types of papers (reviews, full papers, communications, technical notes, highlights, etc.) are welcome for consideration







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Editor-in-Chief

Prof. Dr. Alessandra Toncelli Department of Physics, University of Pisa, 56126 Pisa, Pl, Italy

Message from the Editor-in-Chief

Welcome to *Crystals*, the journal dedicated to the fascinating world of crystallographic research! Crystals are more than mere decorative elements; they hold the key to understanding the fundamental structure of matter. Our mission is to explore the crucial significance of this research across various fields. From medicine to technology, chemistry to geology, crystals play a vital role. Their structure provides insights into new advanced materials, innovative drugs, and groundbreaking technologies. Through *Crystals*, we delve into the microscopic world to discover solutions that will shape the future. Join us on a journey through the *Crystals*, where science merges with beauty and innovation.

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