Symmetry in Quantum and Computational Chemistry

Message from the Guest Editor

Dear Colleagues,

The problem of symmetry in quantum and computational chemistry is a paradigm of development of this field of knowledge. Modern ab initio and semi-empirical methods as well as density functional theory widely use the group theory formalism for investigation of nature and various properties of different periodic chemical systems (crystalline solids, polymers, surfaces and films, nanotubes) and molecules. Researchers in various fields of theoretical chemistry and related disciplines (physics, crystallography, mathematics, computer software development) are welcome to submit their works on this topic in our Special Issue “Symmetry in Quantum and Computational Chemistry”.

The aim of this Special Issue is to highlight and overview modern trends and attract the attention of the scientific community to the problem of symmetry in quantum and computational chemistry.

All types of papers (reviews, mini-reviews, full papers, short communications, and technical notes, highlights) are welcome for consideration.
Message from the Editor-in-Chief

Symmetry is ultimately the most important concept in natural sciences. It is not surprising then that very basic and fundamental research achievements are related to symmetry. For instance, the Nobel Prize in Physics 1979 (Glashow, Salam, Weinberg) was received for a unified symmetry description of electromagnetic and weak interactions, while the Nobel Prize in Physics 2008 (NambuKobayashi-Maskawa) was received for the discovery of the mechanism of spontaneous breaking of symmetry, including CP symmetry. Our journal is named Symmetry and it manifests its fundamental role in nature.

Author Benefits

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Rapid publication: manuscripts are peer-reviewed and a first decision provided to authors approximately 14.3 days after submission; acceptance to publication is undertaken in 3.9 days (median values for papers published in this journal in the second half of 2019).

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