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Density Functional Theory (DFT) and Semi-empirical Quantum Mechanical (SQM) Methods in Organometallic Chemistry

Guest Editor:

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Message from the Guest Editor

Dear Colleagues,

In recent decades, Kohn–Sham density functional theory (DFT) has become the workhorse of computational chemistry, and numerous methods with different focus have been developed. Moreover, modern semi-empirical quantum mechanical (SQM) methods are also seeing more and more applications in various fields of organometallic chemistry due to their outstanding efficiency and their improved reliability in describing organometallic systems. Both represent valuable tools for detailed studies of a wide variety of chemical problems, and their predictive power enables rapid and targeted scientific progress. Accordingly, these methods allow a deep understanding of the very nature of organometallic molecules and reactions far beyond the limits of the experiment.

This Special Issue aims to highlight cutting-edge applications and methodological developments of DFT and SQM methods regarding chemical challenges in the field of molecular organometallic chemistry.











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Editor-in-Chief

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Message from the Editor-in-Chief

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