Special Issue

Modeling Adsorption Properties of Molecular and Nanostructured Systems for Environmental Applications

Message from the Guest Editors

Understanding how adsorption properties towards common pollutants can be improved offers the possibility of solving emerging ecological problems. In terms of aspects of renewable energy, it is also imperative to find materials with superior adsorption properties towards molecular hydrogen. Various molecular and nanostructured systems have been proposed for the efficient adsorption of pollutants and hydrogen, but the quest for perfect adsorbing materials remains open. Ab initio and density functional theory (DFT) calculations, molecular dynamics (MD) simulations and other approaches are indispensable computational tools in the area of adsorption research. A perfect adsorbing material should allow both adsorption and desorption under mild conditions to allow further technical processing of adsorbed molecules, while the adsorber can be recycled several times. This imposes several research challenges, such as appropriate interval of binding energies, specific adsorption mechanisms, high adsorbing capacity, etc.

Guest Editors

Dr. Sania J. Armakovic

Department of Chemistry, Biochemistry and Environmental Protection, Faculty of Sciences, University of Novi Sad, Trg Dositeja Obradovića 3, 21000 Novi Sad, Serbia

Dr. Stevan Armaković

- 1. Department of Physics, Faculty of Sciences, University of Novi Sad, Trg Dositeja Obradovića 4, 21000 Novi Sad, Serbia
- 2. President of the Association for the International Development of Academic and Scientific Collaboration—AIDASCO, 21000 Novi Sad, Serbia

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Molecules
Editorial Office
MDPI, Grosspeteranlage 5
4052 Basel, Switzerland
Tel: +41 61 683 77 34
molecules@mdpi.com

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Editor-in-Chief

Prof. Dr. Thomas J. Schmidt

Institute of Pharmaceutical Biology and Phytochemistry, University of Münster, Corrensstrasse 48, D-48149 Münster, Germany

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