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Drug Discovery: New Concepts Based on Machine Learning

Guest Editors:

Dr. Miroslava Nedyalkova

Department of Chemistry, University of Fribourg, Chemin du Musée 9, 1700 Fribourg, Switzerland

Dr. Andrew S. Paluch

Department of Chemical, Paper, and Biomedical Engineering, Miami University, 64 Engineering Building 650 E High Street, Oxford, OH 45056, USA

Deadline for manuscript submissions: closed (1 April 2023)



mdpi.com/si/141342

Message from the Guest Editors

Dear Colleagues,

The ML/AI-based methodology era enables and opens new avenues that can boost the growth of new methods and their increasing importance. The computational-aided drug design exposes the impacts on drug discovery (new targets, the targeting of small molecules, targeted protein– protein interactions, SAR generation using data-driven experimental databases and integrated platforms, drug delivery pathways, etc.).

The Special Issue will cover the following topics:

targeting small molecules; protein–protein interactions; protein dynamics; docking studies; logP and pKa computational methods; solvation-free energy; QSPR/QSAR studies; fragment-based drug discovery (FBDD).

We also welcome papers dedicated to computational and machine learning for drug discovery. The new ML approach for drug design and CADD was developed and designed for de novo drug design methods to generate a space for novel chemical compounds with desirable properties in a costefficient manner. We are happy to welcome papers dedicated to fragment-based drug discovery (FBDD) as a powerful tool to recognize and classify a new compound as the initial point for drug development.

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Editor-in-Chief

Prof. Dr. Patrick J. Sinko

Department of Pharmaceutics, Ernest Mario School of Pharmacy, Rutgers University, Piscataway, NJ 08854, USA

Message from the Editor-in-Chief

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Pharmaceutics Editorial Office MDPI, St. Alban-Anlage 66 4052 Basel, Switzerland Tel: +41 61 683 77 34 www.mdpi.com mdpi.com/journal/pharmaceutics pharmaceutics@mdpi.com X@MDPIpharma