

Special Issue

Design, Synthesis, and Evaluation of Multi-Target Ligands

Message from the Guest Editor

Designing entirely new multi-target drug, although it seems very easy in concept, is in fact a complex task. The development of various strategies based on ligands or structures, experimental techniques, in silico, and artificial intelligence methods, increase the chances of success in this field. Computer-aided drug design based on modern techniques such as molecular docking, which accelerates screening, and molecular dynamics simulations, which add a "degree of flexibility" to structures, provide insight into the mechanism of ligand-target binding. In silico and artificial intelligence methods predict the physicochemical profiles of ligands and reveal structure-activity relationships. Both of these classes of methods significantly facilitate ligand modelling, analogue design and optimisation of leading compounds. In this Special Issue, we welcome submissions on recent advances in the design, synthesis, evaluation and applications of multi-target drugs.

Guest Editor

Prof. Dr. Jolanta Natalia Latosińska

Faculty of Physics, Adam Mickiewicz University, Uniwersytetu
Poznańskiego 2, 61-614 Poznań, Poland

Deadline for manuscript submissions

closed (15 September 2024)



Processes

an Open Access Journal
by MDPI

Impact Factor 2.8
CiteScore 5.5



mdpi.com/si/182657

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

mdpi.com/journal/

[processes](https://processes.mdpi.com)





Processes

an Open Access Journal
by MDPI

Impact Factor 2.8
CiteScore 5.5



[mdpi.com/journal/
processes](https://mdpi.com/journal/processes)



About the Journal

Message from the Editor-in-Chief

You are invited to contribute either a research article or a comprehensive review for consideration and publication in *Processes* (ISSN 2227-9717). *Processes* is published in open access format – research articles, reviews, and other content are released on the internet immediately after acceptance. The scientific community and the general public have unlimited, free access to the content. As an open access journal, *Processes* is supported by the authors and their institutes through the payment of article processing charges (APCs) for accepted papers. We would be pleased to welcome you as one of our authors.

Editor-in-Chief

Prof. Dr. Giancarlo Cravotto

Department of Drug Science and Technology, University of Turin, Via P. Giuria 9, 10125 Turin, Italy

Author Benefits

Open Access:

free for readers, with article processing charges (APC) paid by authors or their institutions.

High Visibility:

indexed within Scopus, SCIE (Web of Science), Ei Compendex, Inspec, AGRIS, and other databases.

Journal Rank:

CiteScore - Q2 (Chemical Engineering (miscellaneous))