# **Special Issue**

# Computer-Aided Molecular Modeling and Simulation in Drug Design

## Message from the Guest Editors

Computer-aided molecular modeling and simulation has become an essential tool in drug design, allowing researchers to study and understand molecular interactions between a drug candidate and its target. This field combines principles from computational chemistry, bioinformatics, and biophysics to simulate and visualize the behavior of molecules at the atomic level. Many computational methods are also used to predict the toxicity and pharmacokinetic properties of drug candidates. For example, QSAR models can be used to predict the activity of a drug candidate based on its chemical structure, while ADME models can predict how the drug candidate will behave in a person's body. Overall, computer-aided molecular modeling and simulation techniques are important tools in modern drug design, allowing researchers to identify potential drug candidates more efficiently and accurately than traditional experimental methods can. In this Special Issue, we aim to draw together research from experts in the field that highlight traditional and new computational methods and strategies to discover and design new drugs for clinical treatments.

## **Guest Editors**

Dr. José Rogério A. Silva

- 1. Laboratory of Computer Modeling of Molecular Biosystems, Federal University of Pará, Belém 66075-110, Pará, Brazil
- 2. Catalysis and Peptide Research Unit, University of KwaZulu-Natal, Durban 4000, South Africa

Prof. Dr. Gert Kruger

School of Chemistry, University of KwaZulu-Natal, Durban 4001, South Africa

### Deadline for manuscript submissions

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

mdpi.com/journal/ pharmaceuticals





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We hope to handle your contribution to *Pharmaceuticals* soon.

### Editor-in-Chief

### Prof. Dr. Amélia Pilar Rauter

Departamento de Química e Bioquímica (DQB) e Centro de Química Estrutural (CQE), Institute of Molecular Sciences, Faculdade de Ciências, Universidade de Lisboa, Lisboa, Portugal

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