

03. - 07.09.2023 (Frankfurt am Main)

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FEMS EUROMAT is the most important international congress in materials science and technology in Europe. It continues a successful congress series promoting the transfer of knowledge and the exchange of experience between academia and industry. **Submission deadline:28 February 2023**

Area D: Characterization and Modeling

D07: Atomic Scale Modeling of Advanced Materials- Ab Initio, Molecular Dynamics and Monte-Carlo Simulations

The aim of the symposium is to assess the state of the art in applications of theoretical tools that allow for simulations of materials properties at the atomic scale for the knowledge-based design of advanced materials. We will discuss advances and challenges in applications of ab initio calculations, molecular dynamics, and Monte-Carlo techniques focusing on the need to carry out simulations at the most realistic conditions in which materials operate in tools and devices.

The dominant approach in searching for new materials is an experiment. However, it is recognized that the traditional way is too slow, and the goal is to reduce development time and to deploy advanced materials in a more expeditious and economical way. Atomistic simulations allow one to gain far greater insight into physical mechanisms, synthesis, and the properties of materials. Moreover, significant improvement in the predictive power of theoretical modeling has led to an expectation of a shift from a fully empirical paradigm in materials design to the knowledge-based materials design concept, in which atomistic simulations play an important role. During the symposium, we will discuss theoretical methodologies, starting from the basic concepts of quantum simulations and proceeding along the entire atomistic simulations chain, covering molecular dynamics and Monte-Carlo techniques. Furthermore, the use of new computational data-driven techniques to parametrize interatomic potentials or to extend the time and length scales of atomistic simulations shall be discussed.

Targeted Topics include:

- Novel theoretical approaches allowing for improvement of reliability of theoretical simulations, e.g. improved description of many-electron effects in ab initio simulations
- Ab initio simulations at finite temperature
- Treatment of magnetic effects in atomistic simulations
- Improved coarse-graining techniques for multiscale modeling
- Increasing time and length scales of atomistic simulation
- Data-driven approaches in atomistic simulations, including ML/AI

Symposium Organizer



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