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Special Issue on Density Functional Theory Research

Call for Papers

Density functional theory is computational technique used to predict the properties of molecules and bulk materials. It is a method for investigating the electronic structure of many-body systems and is based on a determination of a given system's electron density rather than its wavefunction. The goal of this special issue is to provide a platform for scientists and academicians all over the world to promote, share, and discuss various new issues and developments in the area of density functional theory research.

In this special issue, we invite front-line researchers and authors to submit original research and review articles that explore **density functional theory research**. In this special issue, potential topics include, but are not limited to:

- Density functional theory calculations
- Application of density functional theory
- Thomas–Fermi model
- Hohenberg–Kohn theorems

Authors should read over the journal's <u>Authors' Guidelines</u> carefully before submission. Prospective authors should submit an electronic copy of their complete manuscript through the journal's <u>Paper Submission System</u>.

Please kindly notice that the "**Special Issue**" under your manuscript title is supposed to be specified and the research field "**Special Issue** – *Density Functional Theory Research*" should be chosen during your submission.

Special Issue timetable:

Submission Deadline	March 10th, 2016
Publication Date	May 2016

Guest Editor:

For further questions or inquiries Please contact Editorial Assistant at jmp@scirp.org