Conference on Single Molecule Magnets and Hybrid Magnetic Nanostructures | (smr 1664)

Contribution ID: 29 Type: not specified

Density-Functional-Based Simulation of Molecular Magnets: Anisotropy and Exchange Hamiltonians

Friday, 1 July 2005 09:00 (0:45)

Content

Summary

Primary author(s): MARK PEDERSON (Naval Research Laboratory, Washington, U.S.A.)

Presenter(s): MARK PEDERSON (Naval Research Laboratory, Washington, U.S.A.)

Session Classification : Density-Functional-Based Simulation of Molecular Magnets: Anisotropy

and Exchange Hamiltonians