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Monday–Friday, March 13–17, 2017; New Orleans, Louisiana

Session E50: Magnetic Nanoparticles and Nanostructures

8:00 AM–11:00 AM, Tuesday, March 14, 2017

Room: 397

Sponsoring Units: GMAG DMP

Chair: Dario Arena, University of South Florida

Abstract: E50.00003 : Anisotropy of Zr-Co and Hf-Co nanoclusters using the evolutionary algorithm

8:48 AM–9:00 AM

[Preview Abstract](#)

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Nanostructures of Hf-Co and Zr-Co rare earth free magnetic material that exhibit a high room-temperature energy product. In our study, the evolutionary algorithm coupled with density functional (DFT) method is used to identify the global energy minimum atomic structure of Zr-Co and Hf-Co clusters. Using evolutionary crystal structure optimization algorithm, as implemented in USPEX, we studied the atomic structure, binding energies, magnetic properties, and anisotropy of Zr_xCo_y and Hf_xCo_y ($x=1,2$ and $y=5,7,11$) clusters. A set of metastable and global minimum atomic structures are identified. Several new lower energy configurations were identified for Zr_2Co_{11} , Zr_1Co_5 , Zr_1Co_7 , Hf_2Co_{11} , Hf_1Co_5 and Hf_1Co_7 clusters by our calculations. We discussed the magnetic interaction between the atoms of the clusters which is critical in finding the lowest energy structure. Our calculation show that Zr-Co and Hf-Co have ferromagnetic coupling and large magnetization. We will also discuss the magnetocrystalline anisotropy (MAE) variation in these clusters.