

Contribution ID : 58

Type : not specified

## An investigation of magnetic structure and spin reorientation in Cr and Mn doped rare earth ferrites using neutron powder diffraction

Thursday, 4 February 2016 12:15 (15)

Rare earth orthoferrite RFeO3 is a family of perovskite with fantastic property, such as ultra-fast spin switching[1], photomagnetic excitation[2]and multiferrocity[3]. These properties usually determined by their magnetic structure and unique spin reorientation(SR) effect. The antisymmetric interaction(DM interaction)[4] induce a weak ferromagnetism at room temperature, while the large anisotropic interaction of R3+ ion induce a rotation of Fe3+ spin in the ac or ab plane, viz. spin reorientation. Usually there are 3 types magnetic structure for orthoferrite, in terms of Bertaut's notation[5],  $\Gamma_4(G_xA_yFz)$ ,  $\Gamma_2(F_xC_yG_z)$  and  $\Gamma_1(A_xG_yC_z)$ . For most of magnetic  $R^{3+}$ , there is  $\Gamma_4(G_xA_yF_z) \rightarrow \Gamma_2(F_xC_yG_z)$  transition except  $R^{3+} = Dy^{3+}$  upon cooling[6], which show a  $\Gamma_4 \rightarrow \Gamma_1$ . We investigated the magnetic structure and SR transition of Cr doped  $HoFeO_3$  and Mn-doped  $TbFeO_3$  using neutron powder diffraction. We found Cr substitution for Fe leads to an increasing SR transition temperature of  $\Gamma_4 \rightarrow \Gamma_2$  dramatically. On the other side, the Mn substitution of Fe in  $TbFeO_3$  vanishes the  $\Gamma_4 \rightarrow \Gamma_2$  transition while induces a novel  $\Gamma_4 \rightarrow \Gamma_1 \rightarrow \Gamma_4$  transition. This is unusual because it is usually think it is the the anisotropic rare earth ion determines the SR property. Our observation demonstrate a delicate balance of magnetic interaction in system. This will provide us new interesting physics and potential functional materials.

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Session Classification : Contributed talk