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Electronic and magnetic properties of Ba₂CoWO 6: First principal investigation

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Abstract : We report an investigation of the structural; electronic and magnetic properties of a new double perovskite Ba₂CoWO₆ by means of density functional calculations (DFT); within the generalized gradient approximation (GGA). The lattice constants of Ba₂CoWO₆ was obtained and found toagree very well with published experimental reports. The total and partial density of state are calculated and discussed. The results reveal that the Ba₂CoWO₆ has stable antiferromagnetic character. Our results predicts that Ba₂CoWO₆ have metallic nature.

Keywords: ab initio, FP-LAPW, Electronic structure, Double perovskite