

## Physical Properties of Uranium Dinitride UN<sub>2</sub> by Using Density Functional Theory (DFT and DFT+U)

**Authors :** T. Zergoug, S. E. H. Abaidia, A. Nedjar, M. Y. Mokeddem

**Abstract :** Physical properties of uranium di-nitride (UN<sub>2</sub>) were investigated in detail using first principles calculations based on density functional theory. To treat the strong correlation effects caused by 5f Uranium valence electrons, on-site Coulomb interaction correction via the Hubbard-like term, U (DFT+U) was employed. The UN<sub>2</sub> structural, mechanical and thermodynamic properties were calculated within DFT and Various U of DFT+U approach. The Perdew-Burke-Ernzerhof (PBE.5.2) version of the generalized gradient approximation (GGA) is used to describe the exchange-correlation with the projector-augmented wave (PAW) pseudo potentials. A comparative study shows that results are improved by using the Hubbard formalism for a certain U value correction like the structural parameter. For some physical properties the variation versus Hubbard U is strong like Young modulus but for others it is weakly noticeable such as the density of state (DOS) or bulk modulus. We noticed also that up from U=7.5 eV, elastic results become not conform to the cubic cell elastic criteria since the C<sub>44</sub> values turn out to be negative.

**Keywords :** uranium diNitride, UN<sub>2</sub>, DFT+U, elastic properties

**Conference Title :** ICMSE 2015 : International Conference on Materials Science and Engineering

**Conference Location :** Paris, France

**Conference Dates :** January 23-24, 2015