Theoretical study of structural and thermalproperties of Fe2 (Zr, Nb) Laves alloys

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Abstract : The aim of this is to use ab initio calculations (with the (PP) Pseudopotential method) to give newinsights on structural and thermal properties ofbinary and ternary C15-Laves Fe2(Zr,Nb) alloys. The temperature effects have been obtained using the quasi-harmonic Debye model exploiting the total energy calculations of the PP method. The main results show that: The structural properties obey to the Vegard's law, meaning that $Fe_2(Zr, Nb)$ are quite ordered alloys independent of the Zr/Nb ratio. In addition, The use of the quasi-harmonic Debye model was successfully applied to determine the thermal properties of the $Fe_2(Zr, Nb)$ alloys in the 0-1000 K temperature range. The effect of temperature on bulk modulii and volume expansions is important in Fe_2Nb while lesser in Fe_2Zr .

Keywords: Laves Phases, First-Principles calculations, DFT, Quasi-Harmonic Approximation, Debye's Model, Thermal Properties