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First Principles Study of Copper Sulfides

(for applications as photoconductors)

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Abstract. The Tetrahedrite's family constitutes a complete solid-solution series, and is among the most frequent complex sulfides in Nature. This kind of structure can be generically expressed by the composition, $Cu_{12}Sh_{4}S_{13}$. We have calculated the electronic band structure of $Cu_{12}Sh_{4}S_{13}$ and $Ag_{6}Cu_{6}Sh_{4}S_{13}$ (with band gaps of 1.24 and 1.20 eV, respectively) to demonstrate that different elements occupying certain sites of the crystal structure will make a difference in what concerns the conduction process in Tetrahedrites. We will use this effect and *ab initio* calculations to show that the electronic properties of these compounds make them promising candidates as solar cells photovoltaic materials since not only they possess a direct band gap but their energy falls within the range of energies of photovoltaics. Moreover, we can optimize these properties by doping and will be compared.

Introduction

Tetrahedrite is a copper antimony sulfosalt mineral, $Cu_{12}Sb_4S_{13}$, with I -43m crystal symmetry, seeming to be a promising candidate for solar cells photovoltaics. The Copper-Antimony-Sulfur compound can occur as a stable mineral [1] (Fig. 1) (at ambient conditions), or being synthesized from the constituent elements in stoichiometric proportion [2,3].



Fig1: A specimen of natural Tetrahedrite, from Huaron Mine, Peru.

The crystal structure of the Tetrahedrite can be interpreted as a derivation of the crystal structure of the Sphalerite, which presents a cubic structure F -43m, with composition ZnS. This compound can be transformed into a Tetrahedrite, 1-43m. This transformation can be explained and supported by the principles assisting the geometrical crystallography. It only implies a transformation from F to 1 cell type, keeping the point group (-43m). The latter transformation requires doubling the unit cell size from a_0 = 5.25 Å to a = 10.50 Å. The transformation is described as following: the cubic crystal structure of the Tetrahedrite has two formula units per cell, being derived from the Sphalerite structure by replacing a tetrahedral cluster of sulfur atoms by four packing vacancies (4Vac) in each sixteen packing sulfur positions. Then the vacancies, 4Vac, are replaced by one single S-atom according to the sequence (1), where S^t corresponds to the generic tetrahedral position ansumed by one in sixteen atoms of sulfur (Fig. 2).

 $[S_{16}^{t}] \rightarrow [S_{12}^{t} \operatorname{Vac}_{4}] \rightarrow \{[S_{12}^{t}][S_{0}^{0}]\}$

(1)

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