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First principles study of structural, elastic, electronic and optical properties of triclinic CsAlS₂ chalcogenide.

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B Phale¹, RR Maphanga^{2, 3} and PS Ntoahae¹

¹Department of Physics, University of Limpopo, Private bag x 1106, Sovenga, 0727

²Next Generation Enterprises and Institutions, Council for Scientific and Industrial Research, P.O. Box 395, Pretoria, 0001

³National Institute for Theoretical and Computational Sciences, NITheCS, Gauteng, 2000

Abstract

The structural, elastic, electronic and optical properties of triclinic model CsAlS₂ are investigated using the plane wave ultrasoft pseudopotentials approximation in the framework of density functional theory as implemented in CASTEP code of Material Studio package. The exchange correlation potential is treated with the generalized gradient approximation within the scheme of Perdew-Burke-Ernzerhof. The ground state properties are determined and the calculated elastic constant show that the CsAlS₂ structure obeys the triclinic criterion. The obtained band structure and density of states predict the material to be an insulator with a direct band gap of 3.246 eV at 0 GPa. The optical properties are obtained and discussed, including reflectivity and absorption coefficient, which provide useful information for the future application of CsAlS₂ in photovoltaics.

Apply to be considered for a student ; award (Yes / No)?

Yes

Level for award;(Hons, MSc, PhD, N/A)?

MSc

Primary author: Mr PHALE, Boitemogelo (University of Limpopo)

Co-authors: MAPHANGA, Rapela (CSIR); Dr NTOAHAE, Petros (University of limpopo)

Presenter: Mr PHALE, Boitemogelo (University of Limpopo)

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