First principles investigation of the optoelectronic properties of Molybdenum dinitride for optical sensing applications †

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Abstract: The electronic and optical properties of the newly synthesized Molybdenum dinitride (MoN₂) in the hypothetical 2H structure analogous to MoS₂ is investigated using the Density Functional Theory (DFT) full potential linearized augmented plane wave (FP-LAPW) method and the Modified Becke-Johnson (mBJ) approximation. The aim is to investigate the optoelectronic properties of this compound for potential optical sensing applications and compare with the capabilities of MoS₂ in this field. As compared to MoS₂, which is a semiconductor, MoN₂ is found to be a semi metal from the band structure plots. The dielectric function, optical conductivity and the optical constants, namely, the refractive index, the reflectivity, the extinction and absorption coefficients are evaluated and compared with those of MoS₂ and discussed with reference to the sensing performance.

Keywords: Layered materials; Electronic structure; Dielectric function; Optical conductivity; optical constants; optical sensing

1. Introduction

The high potential of transition metal dichalcogenides (TMD) for electronic, sensing, photonic and thermoelectric device applications has been exploited this past decade and especially MoS₂: a prototype TMD material has shown a lot of promise [1-3]. It has been studied and characterized extensively for structural, electronic, optical and transport properties both in bulk and in the 2D limit [4-6]. Interest in TM nitrides has been rekindled because they exhibit a number of unique and advanced catalytic properties for photo and electrochemical catalysis [7, 8]. There were no layered structures in any of these studies. Layered structures provide more flexibility in doping, ease of going down to lower dimensions and materials design.

The search for layered nitrogen rich TM nitrides, particularly those of MoS₂-type, led to the recent synthesis and discovery of 3R-MoN₂, which has the rhombohedral MoS₂ structure [9]. It was synthesized through a high P–T route of solid-state ion-exchange and has shown great potential for applications in catalysis and hydrogenation. In addition, the very recent first principles study of MoN₂ monolayer by Zhang et al [10] showed the 1H configuration to be the most stable among the structures considered in their study. Their study revealed the importance of 2D MoN₂ as a high capacity electrode material for metal ion batteries. Further, the first principles study of Ramanathan and Khalifeh [11] has shown the 2H MoN₂ to be a promising thermoelectric material.

All the above interesting results for MoN₂ provide a strong motivation to study this compound. Considering that, to date no optical characterization of MoN₂ has been performed; the present study is devoted to the determination of the electronic and optical properties from first principles and to look at the various possibilities for optical sensing.
applications of MoN\textsubscript{2}. Since, an optical sensor measures a physical property of light and depending upon the sensor usage converts it to a readable output, it is highly essential to characterize the optical properties of the new layered material MoN\textsubscript{2}. The hypothetical 2H structure analogous to MoS\textsubscript{2} of MoN\textsubscript{2} is investigated using the DFT full potential linearized augmented plane wave (FP-LAPW) method and the mBJ approximation. In addition, the 2H MoS\textsubscript{2} optoelectronic properties are determined by the same method for the sake of completeness and comparison.

2. Calculation details

The geometry of MoN\textsubscript{2} is optimized using the ABINIT software program [12, 13] with the generalized gradient approximation (GGA) of Perdew, Burke and Ernzerhof (PBE) PAW (projector augmented wave) pseudopotentials [14]. All the structural calculations are performed with convergence criteria of less than $1 \times 10^{-6}$ Ha for the Self Consistent Field (SCF) iterations and a threshold of less than 1 mRy/a.u. for the optimization of the geometries [15, 16]. The fully relaxed MoS\textsubscript{2} lattice constant values are taken from our previous work [6].

The optimized structures and lattice constant values are then used with the WIEN2k [17] code to perform full-potential linearized-augmented plane wave (FP-LAPW) calculation employing GGA\_PBE to obtain the ground state energy and electronic properties at a 20×20×4 k-point grid. The optical properties are evaluated using denser grids of 40×40×5 with the more accurate mBJ exchange correlation of Trans Blaha (TB-mBJ).

3. Results and discussion

3.1. Structural and electronic

The 2H-MoN\textsubscript{2}/MoS\textsubscript{2} unit cells have hexagonal symmetry and consist of two stacks of 3 atomic layers; each stack consists of a Mo atomic plane sandwiched between two N/S atomic planes respectively. The atoms are bonded covalently in plane and the stacks are held together by weak Van der Waals force.

The non-magnetic state is the ground state for both the layered compounds and the structural relaxation of the system with a complete relaxation of all the atoms simultaneously gives us the equilibrium geometry. The lattice parameter a and c values of MoN\textsubscript{2} are 3.094 and 11.975Å respectively. The lattice parameter values of MoN\textsubscript{2} are much smaller than that of MoS\textsubscript{2} due to the shorter bond lengths of Mo-N as compared to Mo-S. The lattice parameters for MoS\textsubscript{2}; a=3.193 and c=12.359Å taken from previous work [6] using the LDA (local density approximation) are in good agreement with the experimental values $a_{\exp} = 3.16Å$ and $c_{\exp} = 12.29Å$ and within 2.3 and 0.6% respectively.

The equilibrium lattice constant values for MoN\textsubscript{2} and MoS\textsubscript{2} are used with the GGA-PBE Wien2k code to extract the electronic band structures with a 20×20×4 k-point grid.

We notice that there is a change in the electronic distribution of MoN\textsubscript{2} as compared to MoS\textsubscript{2} which is reflected in the band structure of MoN\textsubscript{2} as shown in Figure1. The MoS\textsubscript{2} band structure is also shown on the right panel of Figure1. The band structures illustrate the change in behavior of MoN\textsubscript{2} to a semi metal one from the semiconducting one of MoS\textsubscript{2}. There is an overlap between the bottom of the conduction band and the top of the valence band in MoN\textsubscript{2}. This semi metal feature implies that there is a range of energies for which electrons and holes co-exist. The Mo 4d, S 3p and N 2p atomic orbitals play a decisive role in the band structure properties.
3.2. Optical Properties

The TB-mBJ proves to be an excellent choice with a $40\times40\times5$ grid for calculating the optical properties with a high degree of accuracy for MoN$_2$ and MoS$_2$. This section is devoted to the presentation and discussion of the results for the dielectric function and optical conductivity. In addition, the optical constants namely the refractive index, the reflectivity, the extinction and absorption coefficients are obtained and interpreted.

The complex dielectric function ($\varepsilon = \varepsilon_1 + i\varepsilon_2$) is a function of the amount of light absorbed by the material. The imaginary part of dielectric function, $\varepsilon_2(\omega)$, which represents absorption behavior, can be calculated from the electronic band structure of solids. The real part of dielectric function, $\varepsilon_1(\omega)$, which represents the electronic polarization under incident light can be calculated according to Kramers-Kroing relation [20, 21]. Figure 2 shows the real and imaginary plots for the dielectric function for MoN$_2$ and MoS$_2$ in the photon energy range of 0-14eV. We see from the plots the anisotropy of the dielectric function. The general trend is the in-plane values are almost double that of the out of plane direction and the peaks are shifted more towards the right with higher energies.

Figure 2. The dielectric function left panel MoN$_2$ and right panel MoS$_2$: (a) the real $\varepsilon_1(\omega)$ and (b) imaginary part $\varepsilon_2(\omega)$ for the in-plane (xx) and out of plane (zz) directions.

The complex index of refraction of the medium $N$ is defined as $N = \sqrt{\varepsilon} = n + ik$

Where $n$ is the refractive index and $k$ the extinction coefficient. These are depicted in Figure 3.
Figure 3. The refractive index top panel and extinction coefficient bottom panel for MoN$_2$ and MoS$_2$ in the xx in-plane and zz out of plane directions.

Once again we see the anisotropy in the two directions for these optical constants. The amplitudes in the xx direction is larger and closer to the visible range for both $n(\omega)$ and $k(\omega)$ for both compounds. We notice that MoN$_2$ has large static ($\omega=0$) refractive index values of $\approx 11$ and $4$ in the xx and zz directions respectively. In contrast the corresponding values for MoS$_2$ are $4$ and $3$. The larger values of $n(\omega)$ imply higher electron density. In contrast to MoS$_2$, MoN$_2$ has peak extinction coefficient values at $\omega = 0$ of $4.4$ and $0.9$ in the xx and zz directions respectively. Both MoN$_2$ and MoS$_2$ show low $k(\omega)$ values in the infra red and MoS$_2$ continues to have almost zero values upto $1.5$ and $2.5$ eV for xx and zz directions respectively. The first maxims of MoS$_2$ are at $3$ eV of the spectra and the magnitude in the xx direction almost six times larger than in the zz direction. The second k peak for zz direction is in the UV region and slightly larger than the first xx peak.

The conductivity and absorption coefficient graphs for MoN$_2$ and MoS$_2$ are shown in Figure 4.
The graphs show for the xx in-plane direction maximum conductivity is in the UV region for MoN₂ whereas for MoS₂ it is in the visible part of the photon energy. The conductivity in the zz direction has peak positions in the UV region around 8 and 5eV for MoN₂ and MoS₂ respectively. The absorption on the other hand shows the first peak in the visible and second broader peak with a much higher magnitude in the UV region for MoN₂; and a very broad peak of almost constant magnitude for MoS₂ covering the visible and the UV region in the in plane xx direction. Beyond 10eV both MoN₂ and MoS₂ show a rise in the absorption coefficient. With respect to the zz direction there are a set of small peaks beyond the visible and a sharp maximum value peak at around 9eV of the UV region for MoN₂; whereas for MoS₂ the peaks are around 6, 10 and 12eV in the UV region. These characteristics confirm the suitability of MoN₂ and MoS₂ for visible and UV sensing applications. The reflectance is depicted in Figure 5 and we observe large static reflectance value greater than 0.7 for MoN₂ that is double of MoS₂ in the xx direction. In the zz direction the values are 0.35 and 0.25 for MoN₂ and MoS₂ respectively. The graphs show that MoN₂ is a good infra-red reflector, whereas MoS₂ reflects best just beyond the visible range.

4. Conclusions

In conclusion as opposed to MoS₂, MoN₂ is a semi metal. Both layered materials show anisotropy for all the optical properties with different magnitudes and peak positions, although the shapes of the graphs for the same property are similar in the two directions.

The large values of refractive index and good conductivity, absorption and reflectance results obtained reinstate the suitability of these materials for sensing applications in the visible and UV region.

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