# DFT-analysis of 3D & 2D model of $MoS_2$ [2H] Density of states and band structure

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#### Abstract

Analysing  $MoS_2$  using DFT calculations with PE-GGA functional provides an inaccurate approximation to experimental data of the bandgap of bulk and monolayer [1.29eV][1.90eV], respectivly [1] [2]. Where the bandgap for bulk was calculated to 0.93eV, with a relative error  $\epsilon_{rel} = 27.9\%$ , and for monolayer bandgap of 1.70eV with  $\epsilon_{rel} = 10.5\%$ . Looking at both bulk and thin layer  $MoS_2$ , it was discovered that the band structures for these two structures differed quite a lot. Where the bulk  $MoS_2$  gave an indirect bandgap and the thin layer  $MoS_2$ , showed a direct bandgap.

#### I. Introduction

Molybdenum disulphides has a so-called isotypic crystal structure and is classified as transition metal dichalcogenide. In this paper, one will investigate both bulk and monolayer structures of  $MoS_2$  2H. This kind of arrangement of molybdenum sulphide is trigonal prismatic, which means that the sulphides is arranged in a symmetric order above and underneath the molybdenum atom.

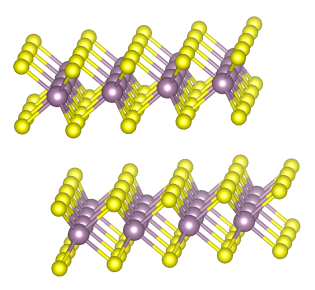


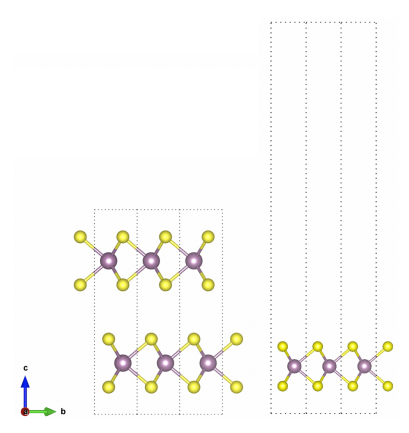
Figure 1: Illustration of the bulk  $MoS_2$  2H. Where the yellow illustrates the sulphides and purple molybdenum atoms

The structrue  $MoS_2$  [2H] has a hexagonal structure indicated by [H].  $MoS_2$  occurs naturally in nature as molybdenite. in its bulk form, as we will investigate, it appears as a dark, shiny solid. The bulk layer sheets is held up by weak Van der Waals forces. These adhesive forces binds the bulk solid togheter. In bulk form it is often used as a lubricant, due to its weak interlayer forces. As for the thin layer  $MoS_2$ , we extrapolate the layer of the bulk  $MoS_2$ . We will investigate which properties that change as we go from bulk to thin layer film. By looking at density of states for both systems and investigating the band structures. Further more when analysing  $MoS_2$ , we will use standardize self-consistent calculation method in which the Hartree Fock method is applied.

The transition metal dichalcogenide  $MoS_2$  is known for its electrical properties as well as its optoelectronic properties. According to experimental values the bulk  $MoS_2$  has a indirect bandgap of 1.29eV [1], while the monolayer  $MoS_2$  has a direct bandgap of 1.90eV [2].

# II. COMPREHENSIVE DFT-ANALYSIS

In this section there will be a brief but comprehensive description on how the results where manifested. The DFT-analysis where done on two materials. Bulk and monolayer  $MoS_2$ , which consist of 6 and 3 atoms in the desired supercell. The fact that the supercell consisted of a small number of atoms, our DFT-calculations where minimized with respect to computational power.



**Figure 2:** Visualization of  $MoS_2$  2H - bulk (left) and the slab (right). Supercells is defined within the stippled lines.

As we can see the by figure 2, Slab contains quite different length in the c-direction. A vacuum layer of of 12Å where inserted to reduce interlayer interactions due to Van der Waals forces. The POSCAR file<sup>1</sup>, contain structural information on the material  $MoS_2$ . Once we have established which structure to work on, we need to start our DFT calculations. One commonly starts with Cut-off energy and **k**-points density. It is important to have sufficient cut-off energy and **k**-points densities before starting the calculations. Before diving in to this, we first need to take a closer look at how to create the slab.

# 2.1. Creation of $MoS_2$ Slab

In able to get a good approximation of a 2D-material one need to insert a vacuum layer in the material, to distance the interlayer in the bulk. A self-consistent DFT calculation where done here, by inserting a vacuum layer in the c-direction. Gradually increasing the vacuum layer with 2Å per calculation. In the INCAR-file the tag IVDW = 1 where inserted. This tag takes care of the Van

der Waals forces in between the layers. The main object here, is to see when the total energy is converged.

Listing 1: Initial POSCAR-file for bulk

```
MoS2
 1.000
      3.190
                0.000
                            0.000
                            0.000
                2.763
      -1.595
      0.000
                0.000
                           14.879
Direct
         0.667
                 0.250
  0.333
          0.333
                  0.750
  0.333
                  0.855
          0.667
  0.667
          0.333
                  0.145
  0.667
          0.333
                  0.355
  0.333
         0.667
                  0.645
```

Listing 1 the POSCAR-file for bulk  $MoS_2$  2H is shown, where for creation of slab one increases the c-direction with a vacuum layer.

Table 1: Inserted vacuum layer inserted in c-direction in POSCAR file, with corresponding energies

Vacuum layer [Å]	$E_{tot}$ [eV]
2	-22.356
4	-22.355
6	-22.354
8	-22.355
10	-22.355
12	-22.355

Table 1, gives a good convergence of energies as vacuum layer is increased. In this analysis a vacuum layer of 12Å where used for further calculations to ensure no interlayer interactions. Once the slab has been created, the convergence test of cut-off energy and **k**-points density may be established.

# 2.2. Convergence tests

# 2.2.1. Cut-off energy $[E_{cutoff}]$

This subsection will contain information on cut-off energy and **k**-points density convergence tests. The convergence test is done by checking the relative total energy of bulk and slab. Energy cut-off is an important parameter in DFT-calculations. One need a large energy cut-off in order to reproduce the wavefunction to some desired accuracy. The cut-off energy contains many **k**-points, which all represent a plane wave [3].

$$\psi_{\mathbf{k}}(\mathbf{r}) = u_{\mathbf{k}}(\mathbf{r})e^{i\mathbf{k}\cdot\mathbf{r}} \tag{1}$$

Where  $u_{\mathbf{k}}(\mathbf{r})$  is a bloch function which describes the periodic potential in the crystal. The periodicity of the bloch function means that it can be represented as a set of plane waves that spans the reciprocal space:

$$u_{\mathbf{k}}(\mathbf{r}) = \sum_{\mathbf{G}} c_{\mathbf{G}} e^{i\mathbf{G} \cdot \mathbf{r}} \tag{2}$$

$$\psi_{\mathbf{k}}(\mathbf{r}) = \sum_{\mathbf{G}} c_{\mathbf{k}+\mathbf{G}} e^{i(\mathbf{k}+\mathbf{G})\cdot\mathbf{r}}$$
(3)

Where **G** is the vector defined by  $\mathbf{G} = m_1 \mathbf{b_1} + m_2 \mathbf{b_2} + m_3 \mathbf{b_3}$  These waves is being combined in the calculations to represent the wavefunction. In order to represent the wavefunction one need a sufficient amount of plane-waves, however, an unlimited  $E_{cutoff}$  is unrealistic since it needs an unlimited amount of computing power.

$$E = \frac{\hbar^2}{2m} |\mathbf{k} + \mathbf{G}| \to E_{cut} = \frac{\hbar^2}{2m} G_{cut}^2$$
 (4)

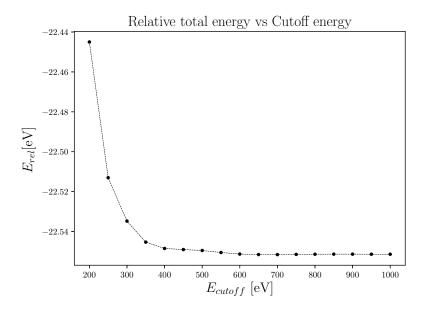
Which further yields a constraint to equation 3:

$$\psi_{\mathbf{k}}(\mathbf{r}) = \sum_{|\mathbf{k} + \mathbf{G}| < G_{cut}} c_{\mathbf{k} + \mathbf{G}} e^{i(\mathbf{k} + \mathbf{G}) \cdot \mathbf{r}}$$
(5)

Therefore we do a convergence test on cut-off energies to see which energy cut-off is sufficient. According to table 2 and figure 3, a sufficient energy cut-off is found at 450eV.

**Table 2:** Convergence of total energy of both bulk and slab as a function of energy cut-off, also relative energy of bulk and slab

$E_{cutoff}$ [eV]	$E_{tot}$ [bulk] [eV]	E <sub>tot</sub> [slab] [eV]	$\rm E_{rel}~[eV]$
250	-44.865	-22.352	-22.513
300	-44.908	-22.373	-22.535
350	-44.929	-22.384	-22.545
400	-44.936	-22.387	-22.549
450	-44.937	-22.388	-22.549
500	-44.938	-22.388	-22.549



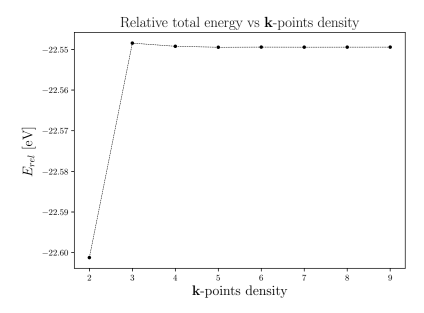
**Figure 3:** Convergence test of  $E_{rel}$  vs  $E_{cutoff}$ , with step-size of 50eV for energy cut-off. Relative taken as  $E_{rel} = E_{tot}[bulk] - E_{tot}[bulk]$ 

## 2.2.2. **k**-points density

Choosing the **k**-point density is also an important aspect for getting accurate DFT-calculations. **k**-mesh is the integration points in our calculations. If our unitcell is large in one direction we need a small **k**-point in that direction. for small unitcells we need a high **k**-mesh in order to produce the results with high enough accuracy. Giving an accurate groundstate density  $n(\mathbf{r})$  and total energy  $E_{tot}$  of the system. By this the number of integration points is altered as we insert a vakuum layer. For the bulk structure the python-file makekpoints -d #density<sup>2</sup> where used as well as for the monolayer structure. Choosing the appropriate **k**-mesh is vital for accurate calculations.

**Table 3:** Convergence of **k**-points density as a function of total energy, and relative energy of bulk and slab.

k-points	E <sub>tot</sub> [bulk] [eV]	$E_{tot}$ [slab] [eV]	$E_{rel}$ [eV]
2	-45.043	-22.442	-22.601
3	-44.935	-22.386	-22.549
4	-44.937	-22.387	-22.549
5	-44.937	-22.388	-22.550
6	-44.937	-22.388	-22.550
7	-44.937	-22.388	-22.550



**Figure 4:** Convergence test of  $E_{rel}$  vs **k**-points density. With step-size of 1 **k**-points density. Relative taken as  $E_{rel} = E_{tot}[bulk] - E_{tot}[bulk]$ 

As we can see by table 3 and figure 4 the convergency rapidly occurs as the **k**-points density is increased. For further calculations a **k**-point density of 5 was submitted. This gave sufficient convergence when the relative total energy where analysed. When it comes to the mesh-points for a thinlayer structure it is sufficient to set **k**-mesh to one in the direction where the vakuum layer is being implemented.

<sup>&</sup>lt;sup>2</sup>In the directory: ~olem/fm4111/bin/makekpoints

## 2.3. Electronic Density of states & Band structure

In this section we will undergo a brief study of relaxation of the structure before going forwards with density of states and band structure. To relax the bulk structure was demanding. Problems occurred during the iterations, towards relaxing the structure. The fact that the energies diverged at some point was troubling. Managed to overcome this problem, by readjusting the lattice vector in the c-axis. Small change gave an easy relaxation of the structure.

The relaxation was done on both the ions and the unitcell. Where the IBRION = 1 tag is used to relax the ions into their instantaneous groundstate. However, the problem may have been that the initial position guess was not suitable for IBRION = 1. By setting IBRION = 2 we got a satisfactory answer, where the structure was relaxed without further implications. Once the structure was relaxed, one can calculate the density of state and band structure.

Density of states was calculated using NSW = 0, a static self-consistent calculation. Used DOSCAR to plot the density of state as a function of E [eV]. In order to get an accurate representation of density of state (DOS), one needed to use a large number of **k**-points. The reason is that when calculating DOS one analyses the integrals in **k**-space, a more compact **k**-mesh gives a more accurate description of the integrals.

$$\bar{g} = \frac{V_{cell}}{(2\pi)^3} \int_{BZ} g(\mathbf{k}) d\mathbf{k} \tag{6}$$

Previously in our calculation a **k**-mesh of  $10 \times 10 \times 3$  for bulk and  $10 \times 10 \times 1$  for monolayer. With **k**-density of 5. By increasing the number of **k**-points to  $24 \times 24 \times 6$  and for monolayer  $24 \times 24 \times 1$ , with **k**-density of 12. We where able to get a more accurate description of DOS.

When calculating the band structure, the first step contained the same  $\mathbf{k}$ -mesh. The first calculations done where self-consistent calculations using PAW PBE potential with Perdew-Burke-Ernzerhof-generalized gradient approximation functional (PE-GGA), the produced CHGCAR file was then used to calculate the band structure. The next step was to alter the KPOINTS-file to contain the symmetry path of  $\mathbf{k}$ -points. By this we needed to understand how the Brillouin zone for a

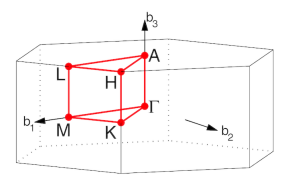


Figure 5: Illustration of Brillouin zone in hexagonal lattice with irreducible Brillouin zone marked in red line

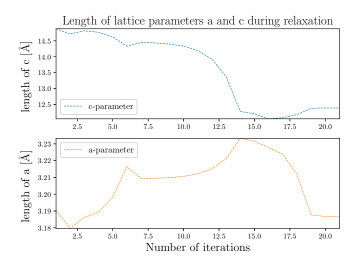
hexagonal structure was structured.<sup>3</sup> Figure 5 show the Brillouin zone as well as the high-symmetry path. These **k**-points paths where written into the KPOINTS-file. The full high-symmetry path is as follows:  $\Gamma - M - K - \Gamma - A - L - H - A|L - M|K - H$ , however, the path was reduced to  $\Gamma - M - K - \Gamma$  in this text. This is because the area of interest lies within this region. We also need to add the tag ICHARG = 11 in order to read the CHGCAR file.

<sup>&</sup>lt;sup>3</sup>Figure 5 is taken from the article High-throughput electronic band structure calculations: Challenges and tools [4]

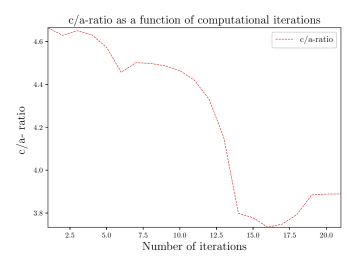
## III. NUMERICAL RESULTS

This section will present results regarding the bulk  $MoS_2$  as well as the monolayer  $MoS_2$ . The electronic density of state for both cases will be presented, the resulting band structures and relaxation of the bulk structure. The plots where possible due to the scripts [plotbands.py, dosplot.py], which where available through the directory:  $\sim$ olem/fm4111/bin. Alternation of the py-files has been done, in order to match up with the given structure at hands. The scripts can be found at GitHub.

# 3.1. Relaxation of bulk structure

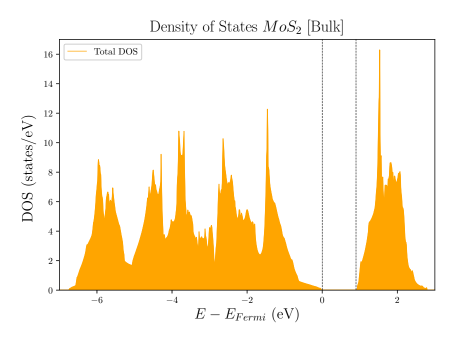


**Figure 6:** Plot of lattice parameters a and c as a function of relaxation processes. By using IBRION=2, the bulk structure where relaxed at 22 iterations.



**Figure 7:** Plot of c/a-ratio as a function of iteration of calculations done through the relaxation process.

# 3.2. Bulk structure of $MoS_2$ [2H]



**Figure 8:** Electronic density of states (DOS) of  $MoS_2$  [bulk], as calculated by VASP using PAW-PBE potential with PE-GGA functional. Where the dashed lines indicates the bandgap.

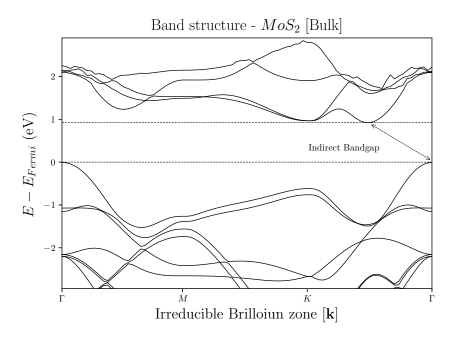


Figure 9: Band structure of bulk  $MoS_2$ , showing an indirect bandgap at high symmetry point  $\Gamma$ , with 30 points from one symmetry point to another. With bandgap calculated to  $E_g = 0.93 eV$ 

# 3.3. $Monolayer\ MoS_2\ [2H]$

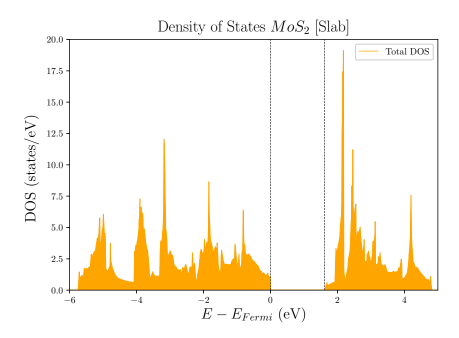


Figure 10: Electronic density of states (DOS) of  $MoS_2$  [Slab], as calculated by VASP using PAW-PBE potential with PE-GGA functional. Where the dashed lines indicates the bandgap.

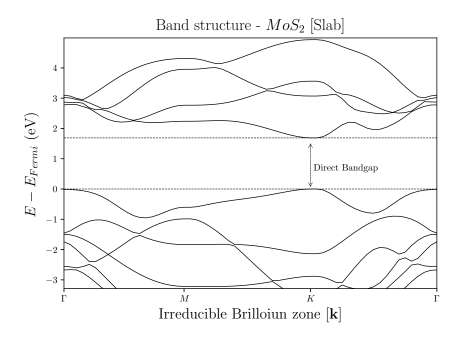


Figure 11: Band structure of monolayer  $MoS_2$ , showing a direct bandgap at high symmetry point K, with 20 points from one symmetry point to another. With bandgap calculated to  $E_g = 1.70 eV$ 

## 3.4. Bandgap of bulk and monolayer

**Table 4:** Bandgap of both structures shown for DFT-calculations using PE-GGA functional and experimental values [1] [2]

Structure	Bandgap [eV]	
	$DFT ext{-}calulations$	$Experimental\ values$
Bulk	0.93	1.23
Slab	1.70	1.90

#### IV. DISCUSSION

## 4.1. Relaxation of bulk

In the process of relaxing the structure, which is a crucial part of DFT-calculation, the monolayer structure where relaxed through few iterations, giving a satisfactory result. Relaxing the bulk structure on the other hand showed to be a slight problem when performing the calculations with IBRION = 1, this problem was due to the poorly initial guess on lattice vectors. By listing 1, which contains the original POSCAR file, the structure had a c/a ratio of:

$$\frac{c}{a} \approx 4.66 \tag{7}$$

which is fairly high for a bulk structure, relaxing this structure gave inconclusive results using IBRION = 1, however by changing from IBRION =  $1 \rightarrow 2$ , we where able to better handle the initially bad guess. As we can see by figure 6, the lattice parameters converges as number of iterations reaches 22. The initial positions stated from listing 1, where a poor initial guess. By 7 we can see the overall c/a-ratio as the relaxation takes place. After the relaxations the c/a ratio was altered, mostly due to change in c-direction. The calculations gave new values a = 3.188Å and c = 12.403Å, which gives a c/a-ratio of:

$$\frac{c}{a} \approx 3.89 \tag{8}$$

The the c/a-ratio has decreased with approximately 16.5% after the relaxation, which is quite a lot. Experimental values of stable lattice parameters for hexagonal  $MoS_2$  of a and c is shown to be 3.15Å and 12.3Å respectively [5], these values yields a c/a-ratio of 3.90, which is in good agreement with our DFT-calculations.

#### 4.2. Band structure

The main goal of this article was to investigate the alternation of band structure from bulk to monolayer  $MoS_2$ , as well as look at the given electronic density of states in both cases. As we can see by figures 9,11. The bulk exhibits an indirect bandgap, at the  $\Gamma$  centre point, while the monolayer exhibits a direct bandgap at the symmetry point K. The shift of bandgap location is interesting, as one can see there is clearly a shift going from a bulk to monolayer structure. Some explanation may be that when the structure is in bulk, the monolayers is interacting with each other through Van der Waals forces, the symmetry and geometry of the crystal is different from the single monolayer structure. The most important reason for a change is what happens when we look at the monolayer, it experience a so-called quantum confinement effect, this is because the particles size in the c-direction is to small to be comparable with the electron wavelength. As the dimension is decreased we get discrete energy levels  $\epsilon_{n,z}$ , where n is the quantum number. So this discretization of energy levels in the z-direction(equivalent to c-direction), gives a shift from  $\Gamma$ -centered valence

band to K-centered valence band. For the indirect bandgap, there is needed a change in momentum in order to transition the electron from valence band to conduction band. This change in momentum mostly comes from phonon contribution. For the direct bandgap, there is no need for change in momentum, as long as the photon has energy near the bandgap energy  $E_g$ , it can excite the electron up to the conduction band, and an emission of photon may take place.

# 4.3. Electronic Density of States

Figures 8 shows the electronic density of states for bulk. The DOS gives an calculation of number of electron states within a interval E, E+dE, where it is plotted as a function of  $E-E_{fermi}(eV)$ , At E  $=E_{fermi}$ , and the region where  $E < E_{fermi}$  we see the valence band. The region where the density of states is zero theres a bandgap. The stippled line at  $E - E_{fermi} = 0.93 eV$ , and above shows the conduction band. As we can see, there is a small amount of electron states at the valence band maxima and conduction band minima for the bulk. At zero kelvin T = 0K, there is no electrons in the conduction band due to fermi-dirac statistics. At zero kelvin the fermi-function acts as a step-function, where all electron is populated in the valence band, where as no electron has sufficient energy to be in the conduction band. 10, shows the electronic density of states for the monolayer. A clear distinction between figure 8 and 10 is that there is a higher number of states occupied near the valence band maxima for the monolayer. Tracing back to the discussion about direct bandgap, we can see that there is a chance that more electrons will occupy near the valence band maxima, yielding a higher order of photon emission. Another interesting property of monolayer  $MoS_2$  is that as a photon excites an electron to the conduction band a hole is created in the valence band, yielding an existing exciton (electron-hole par). As the electron and hole is seperated, they create an electrical current in the crystal, which is one of the fundamental properties of a semiconductor. Finally by looking figures 8 and 10, observe some sharp peaks, which a is discontinuously change in the slope of DOS. These discontinuities is characterized as Hove singularities [3].

# 4.4. Calculated bandgap $[E_q]$ vs experimental value

The Bandgap estimated by VASP, was calculated to to 0.93eV for bulk  $MoS_2$ , which is dramatically underestimated compared to the experimental value of 1.29eV [1]. For the monolayer it was calculated to 1.70eV compared to the experimental value of 1.90eV [2]. The calculation done by VASP is as we see roughly underestimates the bandgap for both cases. As standard DFT-calculations using a PE-GGA functional fails to correctly estimate the electronic states near the fermi energy. Relative error is calculated below for bulk case, with  $E_g$  being the calculated bandgap from VASP and  $E_{ex,g}$  being the experimental bandgap.

$$\epsilon_{rel} = \frac{|E_g - E_{ex,g}|}{E_{ex,g}} \times 100\% = \frac{|0.93eV - 1.23eV|}{1.23eV} \times 100\% = 27.9\%$$
(9)

Relative error of bandgap for monolayer  $MoS_2$ :

$$\epsilon_{rel} = \frac{|E_g - E_{ex,g}|}{E_{ex,g}} \times 100\% = \frac{|1.70eV - 1.90eV|}{1.90eV} \times 100\% = 10.5\%$$
(10)

A better way of calculating the bandgap is to use hybrid functionals, one example is HSE06, which provides a better estimate of the bandgap for small bandgap semiconductors. As we see the bandgap of bulk is much narrower than that for monolayer. This can be explained by the decrease of dimension in the c-direction. As the dimension is decreases we reach reach a discretization of energy levels as mentioned above, this increases the the size of the bandgap.

#### V. CONCLUSION

The bulk and monolayer  $MoS_2$  band structures and electronic density of states where calculated, and showed a clear transition from indirect to direct bandgap. The monolayer structure is therefore more suitable for photonic devices, due to no momentum change in  $\mathbf{k}$ . Monolayer  $MoS_2$  is beneficial in application such as photodetectors, photovoltaics and photocatalysis. The bandgap calculated was grossly underestimated compared to the experimental value. This underestimation is due to the choice of functional, if the calculations had been done using a hybrid functional such as HSE06 functional the results would probably coincide with the experimental values. The PE-GGA functional managed described the shape of the band structure and electronic density of states. The bandgap where align along a high symmetry point, which is fairly normal. The transition of bandgap symmetry point alignment could be described though the minimisation of the dimension along c-direction.

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