Enhancing Efficiency in MoTe2-Based Solar Cells: The Impact of Sb2S3 Hole Transport Layer and Thickness Optimization

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Abstract

In this study, SCAPS-1D simulator was used to investigate the performance of a solar cell structure based on Molybdenum Telluride (MoTe2) with Sb2S3 (Antimony Sulfide) Hole Transport Layer (HTL). The motivation behind choosing MoTe2 as an absorber layer for its higher optical absorption efficiency, cost-effectiveness, reliable and stable operation. The comparative study of this introduced (Al/FTO/CdS/MoTe2/Sb2S3/Pt) and baseline solar cell (Al/FTO/CdS/MoTe2/Pt) has been implemented. Various photovoltaic parameters like open-circuit voltage, short-circuit current, fill factor, and efficiency have been investigated varying absorber and HTL thickness, doping density, rare surface recombination velocity, defect density, series as well as shunt resistance and temperature. The proposed solar cell performance of η , Voc, Jsc, and FF was found to be 40.33%, 1.13 V, 40.78 mA/cm2, 87.63% optimizing absorber thickness value of 0.5 µm and doping concentration value of cm-3. The determined values of performance parameters Voc, Jsc, FF, and η are 0.95 V, 38.15 mA/cm2, 81.09% and 29.35%, respectively for baseline solar cell. The implantation of Sb2S3 layer contributes to improve the performances by diminishing carrier recombination losses. The present research results indicate the feasible way for obtaining a lower-cost, and higher-efficiency MoTe2-based SC with Sb2S3 HTL layer.

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Keywords: TMDC, MoTe₂, Sb₂S₃ HTL, Solar cell, SCAPS 1D, and Efficiency.

1. Introduction

The demand for energy is constantly growing up all across the world. The availability of suitable energy sources is diminishing regularly since there are no available specialized energy storage systems. As a result, a clean energy source that is renewable, long-term, cost-effective, and environment-friendly is required, and solar energy is a candidate here. Solar cells are an excellent choice since they reduce the power generation cost and minimize the need for fossil fuels [1] [2]. Photovoltaic solar cells made of polycrystalline silicon lead the worldwide market and contribute to over 90% of total PV manufacturing [3] [4]. The drawbacks of Si solar structure are massive production charge and low efficiency. The highest power conversion efficiency was recorded at 26.7 % and 24.4 % for crystalline and polycrystalline Si-based PV cells [5]. Nowadays, researchers have been concentrating their attention on manufacturing highly effective thin-film solar cells (TFSCs) using earth-abundant, non-poisonous, and suitable materials. Among the TFSCs materials, Molybdenum Ditelluride (MoTe₂) is a very much effective and promising absorber materials because of its non-poisonous, stability, and cost-effectiveness. It is a transitional dichalcogenide materials (TMDs) with layered architecture of MX_2 , where M = (Mo, W) is the infusion atom belonging to groups 3 to 12 and X = (S, Se, Te) is the chalcogen atom belonging to group 16. The various electronic, optical, catalytic, and magnetic characteristics of TMDs materials have been intensively studied by several research groups [6] [7] [8]. MoTe₂ has a sandwiched structure similar to conventional TMDs, and the weak Van Dar Waals forces bind the atoms together [6] [7]. The textured crystalline of MoTe₂ with the orientation of the c-axis perpendicular to the plane of the grains may contribute to perform as an effective absorber layer [9]. MoTe₂ is a TFSCs material having high carrier flexibility and lower defect density [7] [8]. The advantages of MoTe2 thin films are optical absorption higher than 10⁵ cm⁻¹, efficient, costeffective fabrication process, reliable and stable operation as well as long-term stability. The direct bandgap of MoTe₂ is 1.1 eV with an intense work function value of ~ 4.7 eV which enables an expansion of optoelectronic applications from the visible to the infrared spectrum [9] [10] [11]. While, most of the TMDC materials such as MoSe₂, WS₂ and MoS₂ operate in the visible range [12]. Experimental and theoretical research on TFSCs by utilizing the MoTe₂ absorber layer has been reported by several research groups.

The η , FF, V_{oc}, J_{sc} of the SnO₂/Zn₂SnO₄/CdS/MoTe₂/Mo Solar structure had been recorded to be 17.06 %, 73%, 0.730 V and 23.74 mA/cm² respectively. By inserting an As₂Te₃ HTL layer within the left metal contact and absorber in this same structure, the performances of η , FF, V_{oc} and J_{sc} enhanced to 25.06 %, 84.7%, 1.8 V, and 27.60 mA/cm², respectively [10]. The same research groups investigated the structure SnO₂/Zn₂SnO₄/CdS/MoTe₂/Mo with different BSF layers to increase the behaviour of this solar cell. The η determined was 25.11 %, 25.16 %, and 25.17% for CZT, ZnTe, and In₂Te₃ HTL, respectively [13] [11] [14]. The conversion efficiency of approximately 15.45% had been observed for FTO/CdS/MoTe₂/Ag solar cell and the highest PCE of 29.13% had been attained for optimized FTO/ZnO/MoTe₂/Cu₂Te₃/Ni cell at 1500 nm thickness [1]. But higher efficiency can achieve from the bilayer structure such as for bilayer Graphene/HgCdTe dual and single junction photodetector efficiency is 85.8 and for bilayer, Graphane/ZnO/Silicon heterojunction efficiency is 71% [15].

Since the determined behaviour of the parameters of this introduced MoTe₂ solar cell cannot compete with other heterojunction TFSCs. The conventional MoTe₂ solar cells are needed to be remodelled to achieve excellent performance after inserting a useful hole transport layer within the absorber and the left metal electrode. Band alignment and band gap tuning between the HTL and absorber layer are also crucial. Therefore, it's important to continue more research work for achieving better performance of the MoTe₂ absorber layer. Herein, both V_{oc} and J_{sc} are the essential criteria for enhancing the η of MoTe₂ SC. The obtained values of V_{oc} and J_{sc} of the MoTe₂ solar are still lower than the expected results because of the recombination of the carrier at the intersection. Besides, the inefficient carrier transportation and collection in the electrodes instead of the recombination of the carrier in the interface causes the reduction of the V_{oc} and J_{sc} which results in lower photo conversion efficiency [16][17]. Therefore, redesigning the MoTe₂ solar cell with useful HTL is also essential.

An HTL layer with commendable bodily properties, optical behaviour, and chemical stability may simplify the exact transportation and collection of photo-generated carriers between the absorber and metal electrode. The HTL layers of NiO₂, CuI, Sb₂S₃, CuSCN or CuO_x have been employed in various types of solar cells [18][19][20][21]. Among them, Sb₂S₃ is a very much useful and cost-effective material which can be used as an HTL layer. The Sb₂S₃ is suitable as a transportation layer with a larger band gap, earth-abundant, non-poisonous, and useful for band matching. A

suitable band alignment is anticipated by introducing Sb_2S_3 between the absorber and the left contact which will be benefited for the transportation of the hole and restrict electrons from the absorber layer [22]. Therefore, the photo-generated carriers (holes) can able to transport to the left metal contact through Sb_2S_3 and minority carrier electrons are inhibited by the potential barrier which was developed at $Sb_2S_3/MoTe_2$ interface.

2. Simulation Methodology and Device Modeling

2.1. Simulation Methodology

So many tools are used by the researchers for analysis of the characteristics of the TMSCs like COMSOL, SILVACO ATLAS, AMPS, SCAPS etc. [23]. In this study, we conducted our research using SCAPS software (Version 3.3.05) which is a one-dimensional simulator (SCAPS-1D) software. It has been employed for modelling and simulation works that were developed at Electronics & Information System (EIS) department at Gent University, Belgium. The exploration of the different physical parameters has been done by utilizing carrier transport equations [24]. This simulation software has also been applied for elementary semiconductor equations like electrons and holes transportation, Continuity, and Poisson equations. It may extract the J_{sc} vs V_{oc} (J-V) curve and also the ac characteristic curves like C-F, C-V, and η curve of a PV SC. The performances of this presented structure have been analyzed to exhibit the outcomes of defect density, doping concentration, thickness, temperature, rare contact work function, R_s and R_{sh} using the SCAPS-1D program. The simulations were carried out below the enlightenment of AM1.5G at a working temperature of 300 K.

The SCAPS analysis is based on Poisson's equation, electron continuity and hole continuity, mentioned below [24]:

 $\frac{1}{q}\frac{dJp}{dx} = G_{op}(x) - R(x) \quad \dots \quad (3)$

Where, Ψ , N_D , N_A , ρ_p , ρ_n , q, Jp, Jn, are the electrostatic potential, donor density, acceptor density, hole distribution, electron distribution, electron charge, current densities of hole and current densities of electrons respectively stated as a function of the position coordinate x.

2.2. Device Modeling

Figures 1(a)-(b) display the structural design of the proposed Al/FTO/CdS/MoTe₂/Sb₂S₃/Pt SC and the energy band diagram, respectively. In this model, a p-type MoTe₂ has chosen as the absorber layer and CdS is the n-type buffer layer. The p^+ -type Sb₂S₃ HTL layer is implanted in between the absorber and rare contact layer. The FTO has been devoted to a layer of translucent directing oxide (TCO).

The Pt and Al have been used as the rare and right contact respectively. The values of the physical parameters of several layers have been listed in Table 1. We carefully accumulated these values from several experimental and theoretical works. Table 2 includes the values of the interface defect parameters. The hole and electron thermal velocity are considered 10^7 cm/s at room temperature. The absorption coefficient values of Sb₂S₃, MoTe₂, CdS, and FTO has received from the SCAPS-1D program model [25]. The exterior recombination rate of electrons and holes has assumed 10^7 cm/s for both the back and front sides. For the simulation, the capture cross-section of both hole and electron has been chosen as 10^{-19} cm⁻².



Fig: 1. (a) The structural design and (b) illuminated energy alignment of the Al/FTO/CdS/MoTe $_2$ /Sb $_2$ S $_3$ /Pt Solar Structure.

Table 1. Thysical input parameters utilized of More ₂ -based subcture required in the simulation [11] [22].						
Parameters (unit)	FTO	CdS	MoTe ₂	$\mathbf{Sb}_2\mathbf{S}_3$		
Thickness (nm)	50	50	100 - 1500	10 - 120		
Band Gap, Eg(eV)	3.5	2.4	1.1	1.7		
Electron Affinity, χ (eV)	4	4.3	4.2	3.8		
Dielectric Permittivity, ε_r	9	10	13	7		
CB effective density, N _c (cm ⁻³)	2.2×10^{18}	$1.8 imes 10^7$	107	10 ¹⁹		
VB effective density, N_v (cm ⁻³)	10 ¹⁹	10 ¹⁷	2.4×10^{19}	$1.8 imes 10^{19}$		
Electron Mobility, µe (cm²/V-s)	20	350	110	0.07		
Hole Mobility, µh(cm²/V-s)	10	25	426	0.02		
Donor Density, N _D (cm ⁻³)	10 ¹⁹	10 ¹⁷	0	0		
Acceptor Density, N _A (cm ⁻³)	0	0	5×10^{13} -	$10^{11} - 10^{22}$		
			5×10^{22}			
Defect Type		SA		SD		
Defect Density, N _t (cm ⁻³)	10 ¹⁴	1014	$10^{12} - 10^{21}$	1014		
Energetic Distribution		Gaussian	Gaussian	Gaussian		

Table 1: Physical input parameters utilized of MoTe₂-based structure required in the simulation [11] [22].

Parameters (unit)	MoTe ₂ /Sb ₂ S ₃ interface	CdS/MoTe ₂ interface
Category of Defect	Neutral	Neutral
Capture cross-section of electrons (cm ⁻²)	10 ⁻¹⁹	10 ⁻¹⁹
Capturecross-sectionof holes (cm-2)Positionfor	10 ⁻¹⁹	10 ⁻¹⁹
imperfection energy level, E _t	Above the highest E _v	Above the highest E _v
Energy regarding reference (eV)	0.6	0.6
Total density (cm ⁻²)	$10^{10} - 10^{20}$	$10^{10} - 10^{20}$

Table 2: Boundary constraints cast-off in Al/FTO/CdS/MoTe₂/Sb₂S₃/Pt heterojunction cell.

3. Results and Discussion

3.1. Investigating how the HTL layer's properties are impacted by changes in thickness and doping density on PV solar cell

The proper transportation and collection of photo-generated holes can be obtained by the suitable HTL layer and it can also restrict the flow of electrons (minority carrier). As a result, it reduces carrier recombination. Hence, it's highly essential to demonstrate the consequences of the HTL layer on SC performance parameters. The effect of thickness and doping density of Sb₂S₃ on the performance parameters of the proposed SC has been represented in Fig. 2. The values of thickness have been changed within 10 to 120 nm and the value of doping concentration within 10^{11} to 10^{22} cm⁻³, respectively, while other parameters remain constant. The values of V_{oc} and J_{sc} act with a little variation with rising the thickness of the HTL layer. However, the FF and η demonstrate an almost insignificant amount of reduction. The parameter values of V_{oc}, FF, and η initially show a constant tendency with increasing the doping concentration of the HTL layer upto 10^{21} cm⁻³. After that, it rises slightly when the doping density is more than 10^{21} cm⁻³ and maintains a constant value. The increase in doping concentration and collection of carriers and as a result the rise in the performances of V_{oc}, FF, and η . The decrease of FF with reducing doping concentration in Sb₂S₃ HTL indicates

the rise of series resistance in the introduced solar cell [26][27]. The optimal thickness may be considered as 100 nm and the optimal doping density may be considered to be and 10^{18} cm⁻³ for obtaining the maximum η of 40.33%.



Figure 2. HTL thickness and doping density has varied for presenting the parameters of (a) V_{oc} (b) J_{sc} (c) FF and (d) η .

3.2. Examining the effects of varying thickness and doping density on the MoTe₂ absorber layer

The simultaneous influence of the $MoTe_2$ layer on the performances with the variation of thickness and carrier concentration has been observed and narrated in Fig. 3. The percentage of incident photons entered into the MoTe₂ contributes for enhancing J_{sc} and V_{oc} changes because of absorber concentration. Thus, it's highly needed to optimize the thickness as well as carrier density. These values have been switched within 100 to 1500 nm and 10¹³ to 10²² cm⁻³, respectively. It has been demonstrated that Voc enhances due to the rise of the doping concentration while Jsc reduces. On the contrary, the increase in thickness reduces V_{oc} below the doping concentration of 5×10^{17} cm⁻³ and then it rises marginally. The FF and n rise because of the enhancement of the two varied parameters (thickness and doping concentration). The higher thickness of the MoTe₂ contributes to enhance the absorption of the incident photons which displays higher values of η , FF and Jsc. The rise of doping density may also attribute to enhance the carrier recombination rate and hence, the value of J_{sc} and V_{oc} reduces [28] [29] [30]. At a thickness value of 500 nm and doping concentration of 5×10^{16} cm⁻³, the numerically simulated value of η , FF, V_{oc}, and J_{sc}, is 1.13 V, 40.78 mA/cm⁻², 87.63%, and 40.33%, respectively. For achieving a better performance of Al/FTO/CdS/MoTe₂/Sb₂S₃/Pt SC, we consider the optimum points of doping density and thickness to be 5×10^{16} cm⁻³ and 500 nm, respectively





Figure 3. Absorber (MoTe₂) thickness and doping density has varied for presenting the parameters of (a) V_{oc} (b) J_{sc} (c) FF and (d) η .

3.3. The role of MoTe₂ thickness and defect density in shaping the properties of PV Structure

The presence of defects in the SC system has an opposing impact on PV performances. For that, it's highly beneficial to supervise the impact of defect density on the MoTe₂ layer. The simultaneous influence of defect density and thickness of MoTe₂ on device performances FF and has defined in Fig. 4. It is noticeable that the parameters values of V_{oc} , J_{sc} , FF and η have been decreased from 1.24 to 1.00 V, 40.77 to 17.32 mA/cm⁻², 83.19 to 44.18%, and 42.05 to 5.62% respectively for changing the defect density within 1.0×10^{12} to 1.0×10^{21} cm⁻³ at a constant thickness of 500 nm. The PV parameters enhance with the rising thickness up to the defect density 10^{18} cm⁻³ and reduce after that limit. Whereas values of V_{oc} reduce noticeably with rising thickness. The simulated results demonstrate a clear correlation between defect density and the reduction in parameter values for the MoTe₂ SC. Shockley Red Hall recombination losses are influenced by defects located within the band gap of the absorber layer. A consequence of the SRH mechanism is the reduction in photo-generated carriers, leading to diminished SC performances [31] [32] [33]. At the defect density of 10¹⁴ cm⁻³ and thickness of 500 nm, the numerically simulated determined performance parameters of Voc, Jsc, FF and n are 1.33 V, 40.78 mA/cm⁻², 87.68% and 40.33%, respectively. We have chosen the value of defect density and thickness 10¹⁴ cm⁻² and 500 nm. respectively for determining the optimum performance of the displayed solar cell. By setting the defect density at 10^{14} cm⁻² and thickness at 500 nm, we aimed to determine the best performance for the SC presented.



Figure 4. Absorber (MoTe₂) thickness and defect density has varied for displaying the parameters of (a) V_{oc} (b) J_{sc} (c) FF and (d) η .

3.4. Analysing the effects of altering the temperature and rare surface recombination velocity on PV Structure

The simulation work, displayed in Fig. 5(a), investigated the influence of working temperature (ranging from 274 K to 525 K) on the performance of the MoTe₂ SC. Increasing the working temperature shows a consistent reduction in the PV performances except for J_{sc} . It has been noticed that the value of all the PV parameters (V_{oc} , FF, and η) decreases monotonically except J_{sc} with increasing the operating temperature. The numerically simulated η for the SC without the HTL layer is 29.63% and 22.28% at a temperature of 275K and 525K, respectively. While the maximum η of the newly designed SC with Sb₂S₃ HTL is 40.76% and 32.97% at the temperature of 275K and 525K, respectively. Increased working temperature promotes greater collision rates between photogenerated electron-hole pairs and vibrational atoms, resulting in J_{sc} fluctuations in the proposed SC. The diminishing energy bandgap in the semiconductors, caused by the increasing temperature, is responsible for the drop in V_{oc} [34] [35]. These outcomes are consistent with the findings of multiple research groups investigating temperature-related effects on PV performances [35,36].

. The examination of the back surface recombination velocity (BSRV) on the PV performances has been carried out for the SC, with and without HTL and is visually represented in Fig. 5(b). It can demonstrate a reduction in the values of V_{oc} , J_{sc} , FF, and η has been reduced from 1.13 to 0.91V, 40.70 to 36.00 mA/cm⁻², 87.4 to 77.8% and 40.33 to 25.20%, respectively for changing BSRV from 1×10^{1} to 1×10^{5} cm/s for the baseline Solar structure. No significant changes are observed in the PV performances as BSRV varies, indicating consistent behaviour. The insertion of the HTL layer creates a high electric field which contributes to decreasing the recombination loss. Therefore, the PV parameters remain constant. On the contrary, the absence of HTL accelerates the recombination and the PV parameters reduce [37] [38].



Figure 5. Variation of Photovoltaic parameters of MoTe₂ SC with (a) temperature and (b) BSRV.

3.5. Exploring how the PV parameters of the solar cell are affected by its interface defect density

We have investigated the effect of defect density at $MoTe_2/Sb_2S_3$ and $MoTe_2/CdS$ interfaces of the $MoTe_2$ heterojunction SC under the enlightenment of AM1.5G and 300 K temperature as represented in Fig. 6 (a) and (b). The variation of defect density within 10^{10} to 10^{20} cm⁻² is applied at the interface of $MoTe_2/Sb_2S_3$ keeping others parameters fixed. The results demonstrate a constant behaviour in all PV parameters up to 10^{14} cm⁻², followed by a decrease in response to rising defect density.

At the CdS/MoTe₂ interface, higher defect density results in a significant drop in the values of FF, η and V_{oc}. The values of Jsc remains unaltered at 40.87 mA/cm² until the defect density reaches 10^{16} cm⁻². The degradation of η and V_{oc} from 40.33 to 27.30% and 1.1 to 0.3 V for altering the defect density in between 10^{10} cm⁻² to 10^{20} cm⁻², respectively. The defect density at CdS/MoTe₂ interface has a dominant impact over that of the MoTe₂/Sb₂S₃ interface of the demonstrated structure. The interface defects contribute to increasing the rate of photo-generated carrier recombination which degrades the PV performances [29] [39] [40]. The optimum performance was achieved at the position of 10^{14} cm⁻² of the defect density for both interfaces. The trap state exists at the interfaces behave like a recombination centre and contributes to the degradation of the photo-generated carriers. In this way, the interface defects demand a reduction in the performance of the SC [41].



Figure 6. Influence of interface defect at (a) MoTe₂/Sb₂S₃ and (b) CdS/MoTe₂ interfaces on the PV parameters.

3.6. Understanding the effects of series and shunt resistance on the PV performances

Several factors are liable for the non-ideality behaviour of a solar structure. One of them is parasitic resistance ($R_s \& R_{sh}$). Both these resistances diminish the output performances of the solar structure. The R_s and R_{sh} resistance significantly affects SC performance parameters. The R_s originated from the connection among various terminals between front and back contact in the heterojunction SC structure. In contrast, R_{sh} is generated from defects in the SC that cause to increase in reverse saturation current. Therefore, these resistances have a significant role in SC PV performance.



Figure 7. Comprehensive analysis of (a) series and (b) shunt resistances to enhance the overall performance of MoTe₂ SC.

Fig. 7 (a) and (b) present a comparative analysis of how R_s and R_{sh} affect the PV parameters of the MoTe₂ structure, both in the absence and presence of an Sb₂S₃ HTL. The values of FF, and η reduces with the variation of R_s from 0 to 6 Ω -cm² at a fixed R_{sh} of 10³ Ω -cm². Otherwise, values of V_{oc} and J_{sc} remain fixed over the entire range of R_s . The improper connection at the front contact, back contact and combination among various terminals causes power loss which in turn outcomes in a fall down in the conversion efficiency [28][42]. With the enhancement of R_s , the behaviour of FF decreases dramatically because of solder bond degradation, but the Jsc drop for the optical transmission loss resulted from the encapsulate discolouration [43].

Furthermore, we have investigated the effect of R_{sh} on PV parameters by altering values within 10^1 to $10^{11} \Omega$ -cm² at constant R_s of 1Ω -cm² and displayed in Fig. 7(b). From the simulated results, it is highly noticeable that the performance parameters enhance with increasing the R_{sh} except J_{sc} which is constant. The parameters values increased up to $10^3 \Omega$ -cm², and then got saturated above the $10^3 \Omega$ -cm². The calculated η has been determined 4.16% and 40.33% at the R_{sh} value of 10^3 to $10^6 \Omega$ -cm², respectively. The presences of bulk and interface defects in the proposed Al/FTO/CdS/MoTe₂/Sb₂S₃/Pt SC are liable for decreases in the values of R_{sh} which in turn reduces the PV performance parameters [28] [44].

3.7. The CV characteristics of the optimized device

To characterize the semiconductor devices like solar cells, the C-V profile is an integral part and is used to enhance the device's performance. This study reveals additional basic features of a device structure, where doping density and build-in potential can be derived with the help of two following relations [24]:

Where, the mentioned C, N_a, ε_0 , ε_s , A, V, q are the values of measured capacitance, doping density (1/cm³), the permittivity of the free space (8.85×10⁻¹⁴ F/cm, dielectric constant as per table 1, area

of the cell (cm²), applied potential, the charge of an electron (1.6×10⁻¹⁹ C), respectively. From these two relations, we can find the build-in potential (V_{bi}) and doping density (N_a) from the intercept and the slope of the line in C⁻² vs V plot.

To evaluate the consistency of the studies, the C-V measurements may examine at diverse frequencies ranging from 0.1 kHz to 1 MHz. The total p-n junction may be the summation of the diffusion and depletion capacitance. At forward bias, the depletion capacitance is smaller than diffusion capacitance, while the relationship is opposite for reverse biased condition. At zero bias condition, the capacitance is found 48 nF/cm² as shown in Figure 10 (a). The mathematical expression for the depletion capacitance is as follows [45]:

Their physical meanings are stated above. At a constant frequency, the capacitance exhibits a substantial and rapid rise with the increasing polarization potential. The Mott-Schottky plot of the proposed structure is shown in Figure 10 (b). The intersection of the $1/C^2$ with the voltage axis originates the flat band potential. Here, the negative slope reveals that holes are the majority carrier and electrons are the minority carrier and the space charge area broadly occupies the p-type MoTe₂ absorber layer. Localized deep states in the MoTe₂ layer lead to a noticeable marginal deviation in the $1/C^2$ curve. The C-V characteristics with varying absorber layer thickness (0.1 µm to 0.8 µm) are shown in Fig. 10 (c), keeping frequency constant at 1 MHz. The capacitance value is almost constant up to a certain level and changes dramatically after that level. The same kind of relation has been discussed in the earlier report [46].

The C-V characteristics with varying absorber doping density at a constant frequency of 1 MHz has studied in Fig. 10 (d). With the increases in bias voltage, the capacitance value also changes proportionally. The charge development at the interface increases with the increases in absorber doping density which further contributes to enhancing the capacitance value [41].



Fig: The impact of C-V parameters on SC (a) varying frequency (b) $1/C^2$ vs V curve (c) varying absorber thickness and (d) varying absorber doping concentration.

The depletion width (W_d) and the length of carrier diffusion can be calculated by these two obtained values (V_{bi}, V) , by the following two relations [24]:

Where D_n is the diffusivity of the electrons, τ is the minority career lifetime, μ_n is the electron mobility, $\frac{kT}{q}$ is the thermal voltage at room temperature. Before recombination, electrons may travel by this average distance towards the boundary of the space charge region. The active thickness required for the photocurrent in a MoTe₂ Solar structure can now be easily determined through the equation $L_p=L_n+W_d$ [24].

3.8. The final output performance MoTe₂ solar structure

There is two MoTe₂-based heterostructure solar cell. One is the baseline (without HTL) MoTe₂ SC structure and the other is the displayed (with HTL) heterojunction SC design of Al/FTO/CdS/MoTe₂/Sb₂S₃/Pt, where Sb₂S₃ has been used as an HTL layer and placed within left contact and MoTe₂ absorber layer. Figure 9(a) represents a comparison of the J-V curve of baseline and proposed SC structures. The thickness of FTO, Sb₂S₃, MoTe₂, CdS and has been chosen as 0.05 μ m, 0.1 μ m, 0.5 μ m, 0.05 μ m, respectively. It has been demonstrated that the PV performance of this displayed SC design is greater than the reference MoTe₂ solar structure. The PV parameters of the baseline MoTe₂ SC are 0.95 V, 38.15 mA/cm², 81.09%, and 29.35%, respectively. While the enhanced η of the presented Al/FTO/CdS/MoTe₂/Sb₂S₃/Pt SC is 40.33% containing V_{oc} of 1.13 V, J_{sc} of 40.78 mA/cm², and FF of 87.63%. The absorber layer with Sb₂S₃ HTL creates a strong electric field via the creation at the p⁺-p junction which contributes to a degradation at the carrier recombination rate at the MoTe₂/Sb₂S₃ interface [21] [26] [27] [46] [47]. This reduction of carrier recombination contributes to enhancing the performance of our presented solar structure.



Figure 9. (a) Simulated current density vs voltage (J-V) features and (b) EQE versus wavelength graph of the MoTe₂ SC in the presence and absence of HTL.

The wavelength-dependent external quantum efficiency (EQE) curves of MoTe₂ SC for both reference and presented structures have displayed in Fig. 9(b). It has also been cleared that the EQE of MoTe₂ SC with the HTL layer of solar cells is higher than without HTL and the wavelength has altered within (300~1200 nm) for detecting this effect. The enhancement of the EQE in the longer wavelength may be because of the insertion of the Sb₂S₃ HTL layer which diminishes carrier recombination by creating an electric field at the interface of MoTe₂/Sb₂S₃ [21] [40].

Table 4 reveals the comparison of the performance features of MoTe₂ SC studied by various research groups and proposed structures. This study shows that the η varied from 22.43% to 29.13% in the previously investigated MoTe₂ SC structures. The value of PV performance parameters is lower because of the mismatch of band arrangement at the Sb₂S₃/ MoTe₂ interface that inhibits the photo-generated carrier's conduction. Moreover, the presence of shallow and deep-level defects facilitates SRH recombination [47]. Thus the photovoltaic performances of MoTe₂ solar cell structure need to be enhanced. Our proposed structure has achieved better values by restricting carrier recombination at Sb₂S₃/MoTe₂ and MoTe₂/CdS layers [47]. The present study

of Al/FTO/CdS/MoTe₂/Sb₂S₃/Pt SC demonstrates insight into the development of high-efficiency MoTe₂ SC by suppressing carrier recombination.

No	Structure	Depth (µm)	V _{oc} (V)	Jsc (mA/cm ²)	FF (%)	PCE (%)	References
1	FTO/ZnO/MoTe ₂ /Cu ₂ Te	0.5	0.88	38.14	85.26	29.13	[1]
2	$ZnO/Zn2SO4/CdS/CdTe/MoTe_2$	0.08	0.7	28		28.00	[8]
3	CdS/MoTe ₂ /As ₂ Te ₃	0.1-10	1.07	27.79	84.20	25.06	[10]
4	CdS/MoTe ₂ /In ₂ Te ₃	0.1-5.0	1.07	27.90	84.20	25.17	[14]
5	FTO/CdS/CdTe/Sb ₂ S ₃	1.0	1.15	28.74	86.03	28.41	[19]
6	SnO ₂ /Zn ₂ SnO ₄ /CdS/CdTe/ZnTe	0.1-2.9	1.05	24.34	87.20	22.43	[48]
7	SnO ₂ /Zn ₂ SnO ₄ /CdS/CdTe/SnTe	0.1-3.0	1.05	24.40	87.20	22.48	[49]
8	Zn2SnO4/CdS/MoTe2/CZT	0.1-10	1.07	27.85	84.10	25.11	[13]
9	Al/FTO/CdS/MoTe ₂ /Pt (without HTL)	0.5	0.95	38.15	81.09	29.35	Proposed SC
10	Al/FTO/CdS/MoTe ₂ /Sb ₂ S ₃ /Pt (with HTL)	0.5	1.13	40.78	87.63	40.33	Proposed SC

Table 4: Comparison values of PV parameters with various research works.

4. Conclusions

In this study, we have constructed Al/FTO/CdS/MoTe₂/Sb₂S₃/Pt heterojunction PV solar structure with Sb₂S₃ HTL layer. Utilizing the SCAPS-1D simulator, the simulation successfully examined and compared the PV performances in the presence and absence of Sb₂S₃ HTL. We have extensively investigated the PV parameters due to the changes in thickness, doping density, temperature, series as well as shunt resistances. The impact of Sb₂S₃ HTL insertion has been explored in this research for MoTe₂-based PV cells to acquire the highest efficiency through lower absorber thickness. The constructed Al/FTO/CdS/MoTe₂/Sb₂S₃/Pt solar structure with Sb₂S₃ HTL has performed the η of 40.33%, FF of 87.63%, V_{oc} of 1.13 V, J_{sc} of 40.78 mA/cm² optimizing thickness of 0.5 µm of MoTe₂ absorber layer. While the baseline Al/FTO/CdS/MoTe₂/Pt solar cell has η of 29.35%, V_{oc} of 0.95 V, J_{sc} of 38.15 mA/cm², and FF of 81.09%. The presence of the Sb₂S₃ HTL generates an electric force at the P⁺-P interface, leading to a decrease in carrier recombination. The decrease of carrier recombination contributes to enhancing the performance

parameters e.g. V_{oc} , J_{sc} , FF, and η . These findings indicate that MoTe₂-based heterojunction may be implemented in the solar semiconductor community in a cost-effective as well as feasible way.

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