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Structural, mechanical, electronic and optical properties of N-decorated single-walled silicon carbide nanotube photocatalyst for hydrogen evolution via water splitting: a DFT study

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Abstract

This work investigated the fundamental photocatalytic properties of nitrogen-doped single-walled silicon carbide nanotubes (N-doped SWSiCNTs) for hydrogen evolution for the first time. Investigations of the structural, mechanical, electronic, and optical properties of the studied systems were carried out using popular density functional theory implemented in quantum ESPRESSO and Yambo codes. Analysis of the structural properties revealed high mechanical stability with the 3.6% and 7.4% N-doped SWSiCNT. The calculated band gap of the N-doped SWSiCNT with 3.6% demonstrated a value of 2.56 eV which is within the photocatalytic range of 2.3 eV – 2.8 eV. The hydrogen evolution reaction (HER) and oxygen evolution reaction

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(OER) potentials of the 3.6% N-doped SWSiCNT also showed good agreement with previous theoretical data. The studied material showed the best photocatalytic performance in both parallel and perpendicular directions by absorbing photons in the visible region. Therefore, the observed structural, mechanical, electronic and optical behaviors demonstrated by the 3.6% N-doped SWSiCNT exposed it as a better photocatalyst for hydrogen production under visible light.

Keywords: Photocatalyst, silicon carbide nanotubes, water splitting, hydrogen energy.

1. Introduction

There have been reports of excessive usage of non-renewable energy in recent decades, which has raised anticipation of a rapid end to this source of energy [1]. Global warming and climate change were also promoted due to the discharge of greenhouse gases through the use of fossil fuels [2]. Moreover, our ecosystems, and water bodies including surface waters were polluted and contaminated through this process [3]. These issues, along with others including the potential for depleting non-renewable energy sources, have led to the development of new ideas as fossil fuel alternatives, one of which is the production of hydrogen fuel and CO₂ reduction. It has been reported that CO₂ can be successfully reduced by tailoring surfaces and interfaces of copper-based catalysts by electrochemical methods [4]. Zhang et al also reported efficient CO₂ reduction using a Cu-Zn –based alloy/oxide interface which yields C⁺ products [5]. Apart from CO₂ reduction, electrocatalysts were also found to take part in hydrogen evolution reactions in alkaline water [6]

The sun is the largest source of renewable energy, producing up to 1.23×10^{35} Joules of energy in one year, and many technologies have been harvested based on it to meet the global energy demand [7]. More specifically, to solve the ever-growing energy crisis and environmental problems, solar energy has been exploited through various means and many methods. Photocatalysis (i.e., light energy is used to drive chemical reactions) is one of the successful techniques that has evolved to make hydrogen energy as a possible replacement for fossil fuels [8].

Studies in the literature have demonstrated that H₂ and carbon-free fuel are possible perfect energy carriers because, when combined with O₂ in a fuel cell (FC), they only result in the production of water due to the chemical energy stored in the H-H bond [9]. However, several

issues need to be addressed such as the problem of an efficient industrial-scale hydrogen development process [10], the problem of efficient large-scale storage facilities and the potential of the hydrogen capability to eliminate any waste products with a particular focus on carbon dioxide and other greenhouse gases [11]. Some of the processes that have been adopted for hydrogen production include biological, thermal-chemical, electrolytic, photocatalytic, and photo-electrolytic processes [12]. In the photocatalytic process, it was found that several materials serve as photocatalysts that successfully generate hydrogen throughout the water-splitting process. Recent findings revealed that nitrogen deficient Boron Nitride Nanotubes (BNNTs) performed better as photocatalysts in the decontamination of pollutants and hydrogen evolution due to minimum recombination of charge carriers [13].

Research conducted by Yuali et al. showed that the porosity of flourey MoS₂ enhances its photocatalytic performance through adsorption and transfer of methylene blue (MB) to the active site [14]. However, this type of photocatalyst did not last long under strong sunlight and therefore needs to be improved. TiO₂ nanotubes have shown excellent performance against solar radiation in the visible range [15] and have been successfully used for water decontamination [16]. This property has been remarkably achieved due to their high surface area-to-volume ratio, high stability and favorable photoelectric property conversion characteristics. In a study conducted by Wang et al. (2019) reported that narrow band gap (2.6–2.8 eV) of (Zn_{1+x}Ge)(N₂O_x) were the reasons for its photocatalytic properties [17] [18]. It has also been reported that doping sites significantly affect the photocatalytic activities of semiconductors [19].

In addition, other photocatalysts such as carbon nanotube composites, metal oxide nanotubes, etc. have been tested for hydrogen evolution by water-splitting technology. The main problem is their ability to absorb large amounts of photons in the visible range, which depends on the electronic interactions between atoms in the valence bands and conduction bands [20]. During the generation of hydrogen by photocatalysts, photons with sufficient energy hit the semiconductor surface, which excites electrons into the conduction band. This leaves holes in the valence band whose interactions, along with those of the conduction band electrons, stimulate the absorption of electromagnetic energy from the Sun. Water splitting is the process in which hydrogen is evolved from the reduction reaction and the oxidation reaction of water molecules. The process is called Hydrogen Evolution Reaction (HER) and Oxygen Evolution Reaction (OER). The equations for total water splitting are given as [21]



Photocatalytic water splitting technology is a good technology that uses solar radiation to generate hydrogen from water and also helps provide clean energy sources. For a material to work well as a photocatalyst for water splitting, the range of the band gap should be between 2.2 eV to 2.8 eV and has to be more than the free energy of water splitting (1.23 eV) [22], therefore, the energy of the photo-generated electrons must be greater than the 1.23 eV. In addition, it should absorb solar energy in the visible range. Also, the band gap should be less than 3.0 eV to prevent poor luminous efficiency and the HER and OER potentials should be 0.02 eV and -0.77 eV respectively [23]. To provide more materials (i.e., explore new materials) that are better suited for the production of hydrogen throughout the water-splitting process, this research investigated the mechanical, electronic, and optical properties of new photocatalysts optimized from nitrogen doping to single-walled (7, 7) silicon carbide nanotubes (SWSiCNT). It is worth mentioning that no other studies have been conducted previously on the photocatalytic properties of N-decorated single-walled silicon carbide nanotube for the purpose of hydrogen evolution via the overall water-splitting technique. Additionally, we have been prompted to continue research in photocatalysts because of the diverse applications of photocatalytic processes in different fields. For instance, the practical applications of the photocatalytic process have already been demonstrated by many researchers worldwide including Professor Santiago Eesplugas, who has reported the significance of various photocatalysts in areas such as wastewater treatment, treatment of metoprolol, photooxidation, etc [3, 12].

2. Research method

The optimizations of the geometric structure of the nitrogen-doped SWSiCNT were performed using DFT with generalized gradient approximation (GGA) parameterized exchange-correlation and the Perdew–Burke–Ernzerhof (PBE) functional [24], [25]. All calculations were performed using quantum ESPRESSO and Yambo codes [26]. Quantum ESPRESSO was chosen for this work because it is the leading high-performance open-source quantum mechanical software package for the nanoscale modeling of materials [27]. Due to some of its limitations such as band gap underestimations, we interface the quantum ESPRESSO with Yambo hybrid soft code to obtain an accurate description of the investigated systems.

The optimized structure was visualized with VESTA and BURAI. As a representative model of our study, we considered the armchair shape of the SWSiCNT with (7, 7) configurations having 28 atoms. The entire structure consists of alternating Si and C atoms separated by a bond length of 1.7 Å. To investigate the free hydrogen evolution reaction (HER) and the oxygen evolution reaction (OER), we adopted the technique [28] by Nørskov and coworkers, which was calculated using the following equation (3),

$$\Delta G = \Delta E + \Delta ZPE - T\Delta S \quad (3)$$

Where, the terms ΔE , ΔZPE , ΔS and T denote the difference in total energy, zero potential energy, entropy and temperature (300 K), respectively. The graphical representation of the whole photocatalytic process is shown in Figure 1.

The optimized system of N-doped SWSiCNT was built such that the nitrogen impurity concentration in the pristine SWSiCNT was 3.6% and 7.4% under different doping. To achieve a good agreement between computational cost and accuracy, we used a kinetic energy cut-off value of 60 Ry to perform all ground state computations on the basis set containing 1x1x30 sample k-points in the gamma point of the Brillouin zone. To determine the mechanical stability of the pristine and N-doped systems of SWSiCNT, we used the given Murnaghan equation of state (shown in Eq. 4), which describes the relationship between the volume of the pristine and N-doped systems with the tensile pressure [29].

$$S(V) = \frac{K_0}{K'_0} \left[\left(\frac{V}{V_0} \right)^{-K'_0} - 1 \right] \quad (4)$$

Where $\frac{V}{V_0}$ is the reduction in the volume of the nanotubes under stress and $\frac{K_0}{K'_0}$ is the reduction of the force constant under stress. In order to analyze the photocatalytic energy range of the pure and N-doped systems of SWSiCNTs, the electronic band gap and the total density of states were evaluated. Concerning the photo absorption properties of the systems, we determined the optical absorption energies of the studied systems using the imaginary dielectric functions based on the Kramers-Kronig relations, as shown in Eq. (5) [30].

$$\varepsilon_2 = -\frac{2\omega}{\pi} P \int_0^\infty \frac{\varepsilon_1(\omega') - 1}{\omega'^2 - \omega^2} d\omega' \quad (5)$$

Where ω is real and where P denotes the Cauchy principal value.

3. Results and discussion

3.1 Structural and mechanical (elastic) properties

To study the photocatalytic properties of nanosystems, it is crucial to analyze their resistance to mechanical exposure to solar radiation. Photocatalysts with lower Young's moduli, low surface area to volume ratio, and low tensile stress are not efficient for photo absorption [31]. As part of this work, we have investigated the structural and elastic properties of the systems under investigation and analyzed the results accordingly. The optimized structure of the pristine SWSiCNTs, C replaced N-doped SWSiCNTs and Si-replaced N-doped SWSiCNTs are shown in Figures 2(a), 2(b) and 2(c), respectively, while Table 1 presents the distribution of formation energies with different states of the SWSiCNT systems. It can be seen from Table 1 that the diameter of the SWSiCNT decreased from 30.01 Å to 29.51 Å and 29.02 Å when 3.6 wt% and 7.4 wt% N dopants were used, respectively. This is because, with increasing tensile (external) pressure on the nanotube, its volume decreases. After all, Si and C atoms are pressed together and bond lengths are shortened. Conversely, a decrease in the tensile pressure leads to an increase in the volume of the nanotube since the Si and C atoms are now far apart. In addition, the formation energy is highest for Pristine SWSiCNT and lowest for 7.4 wt% N-doped SWSiCNT. The claim to this statement rests on the fact that the release of energy for bond formation is always negative since this process is always exothermic. Also, N dopants favored the N-doped SWSiCNT system via reduction of the formation energy of Si vacancies hence affecting the charge transfer of neighboring atoms [32].

It was found that SWSiCNT was not stable when N atoms replaced the C atoms. However, the system showed good stability when N atoms in different concentrations were used to replace Si atoms which were in good accordance with the obtained experimental results [33]. The doping was done by replacing C atoms with N atoms in the original structure of (7, 7) SWSiCNT, giving the atomic composition of the doped system 50% Si, 46.4% C and 3.6% N atoms. The calculated bulk modulus for pristine SWSiCNT was 913.6 GPa, while that of the N-doped (7, 7) SWSiCNT was 929.4 GPa. The tensile pressure (strain) showed an inverse variation with volume for all nanotube systems, which is consistent with the obtained theoretical insights [34].

In the case of bare SWSiCNT, the most stable volume of the optimized nanotube corresponds to 1777.44 (a.u)³ as shown in Figure 3(a). In this region, any additional increase in volume on the system will lead to severe defects such as dislocations, etc. Figure 3(b) shows that the most

stable volume of the 3.6% N-doped SWSiCNT is blue-shifted. It has happened due to N-doping following the reduced atomic radius of the N atom. Due to the increase in the concentration of the N-dopant from 3.6% to 7.14% (figure 3(c)), the most stable volume further decreased to 1776.36 (a.u.)^3 . However, it should be noted that the calculated volumes of these nanotubes are still within the experimental range and therefore have not changed the mechanical stability of the systems studied. Figure 3(d) shows the effect of volume variation on the total energy of the pristine SWSiCNT system. It can be seen that the total energy varies directly and then inversely with the volume of the nanotube. The optimal volume to achieve the overall optimized energy of the system was found to be 1283.91 (a.u.)^3 , which corresponds to a total energy of -985.35 eV . In this region, the electron-hole interactions outweigh the nucleus-nucleus interactions. The optimized volumes determined for 3.6% and 7.4% N-doped SiC nanotubes were 1283.82 (a.u.)^3 and 1281.91 (a.u.)^3 , respectively. It was found that the obtained values for the optimized volumes of the N-doped systems all agreed well with the standard experimental data for nanotubes [35]. Furthermore, the calculated mechanical strength found from the N-doped SWSiCNT systems demonstrated their good ability to withstand much mechanical pressure during photo absorption.

To determine Young's modulus of the nanotubes under consideration, we have evaluated the total energy of both pristine and N-doped (7, 7) SWSiCNTs under different bond lengths using the pure DFT method. Results were obtained for variation of deformation energy with different Si-C and C-N bond lengths and the corresponding Young's modulus was obtained. Figure 4(a) presents the obtained results by varying bond lengths. From the results obtained, the closest distance between the two nearby Si-C atoms was 1.71 \AA which corresponds to a bond energy of -985.35 eV . The previously obtained optimized volume of 1283.91 (a.u.)^3 (shown in Figure 3a) is in good agreement with the obtained experimental values of 1.7 \AA [36]. Figures 4(b) and 4(c) showed that the calculated bond lengths of the 3.6% and 7.4% N-doped SWSiCNTs were 1.69 \AA and 1.68 \AA , respectively. The lower value of the bond lengths of these nanotubes compared to pristine SWSiCNT makes them stronger and therefore able to withstand unfavorable changes in mechanical states. To confirm this statement, we found that the calculated bond lengths of all three systems were inversely proportional to the bond order. Strong bond orders are therefore accompanied by strong attractive forces that hold the interacting atoms together. Furthermore, for all three systems, the inverse variations in bond energy with bond length were due to repulsive forces, while the obtained direct variations in bond energies were due to attractive forces. On this basis, it was found that the stability of the N-doped systems shows a significant

increase, thus requiring higher enthalpy to break the bond. Figures 4(d), 4(e), and 4(f) show the analysis of the calculated Young's moduli of the investigated systems with different lengths of the nanotubes. The 7.4% N-doped SWSiCNT was found to be stiffer due to the reduced bond length and higher dopant concentration, while pure SWSiCNT exhibited lower stiffness due to longer bond length. A summary of the calculated mechanical properties of the original SWSiCNT, 3.6% N-doped, and 7.4% N-doped SWSiCNT structures is presented in Table 2. It can be seen from the Table that this doping does not cause any significant change in the structural properties of the systems studied. For example, the volume of the original SWSiCNT changed by only 0.09 (a.u)³ when 4.6% N atom was doped and by 1 (a.u)³ when 7.4% N atom was doped. Therefore, it can be concluded that our doped SWSiCNT system is structurally and mechanically stable for further investigation of its photocatalytic properties.

3.2 Electronic properties

To obtain relevant data on the electronic properties of the N-doped (7, 7) SWSiCNT photocatalyst, it is crucial to understand its pristine electronic behaviors. For this reason, we first analyzed the electronic properties of the pure (7, 7) SWSiCNT system. The results are shown in Figure 5(a) and the calculations were performed with both Kohn-Sham DFT and GW approximations (GWA). Results from KS-DFT using the GGA-PBE exchange function revealed a bandgap of 1.6 eV. In addition, it can be seen that both the valence band and the conduction band have the same momentum. For this reason, the obtained band gap of the pristine SWSiCNT is a direct band gap and this agreed well with the obtained experimental values [37]. Therefore, this type of material generates light during the recombination of electrons and holes (high charge carrier concentration). Due to the reported bandgap problem by the GGA functional, we have repeated our calculations using the hybrid GW within the G_0W_0 approximation and obtained accurate results in which a band gap of 1.7 eV was obtained. Also, the VBM was found to be -0.3 eV which is less than the normal value of -0.77 eV for OER. Similarly, the CBM was found to be 1.2 eV which is also less than 1.23 eV for HER. Therefore, pristine SWSiCNT does not satisfy the condition to serve as a photocatalyst for water splitting because the band gap value of 1.7 eV is outside the range of 2.3 eV to 2.8 eV for efficient absorption of visible light. To tune the pure (7, 7) SWSiCNT, we doped the system with 3.6% nitrogen and observed changes in its electronic behavior after doping. Later on, we increased the N-dopant concentration to 7.4% and

different behavior was observed. Figure 5(b) shows the result of the calculated electronic band gap of the 3.6% N-doped SWSiCNT system. The effect of the N-doping can be seen through the shift of the bands from 1.7 eV to 2.54 eV (with KS-DFT) and 2.56 eV (with G_0W_0) which is blue-shifted. It can be seen that both KS-DFT and GW methods report a band gap within the specified range for photocatalysts. The obtained band gap (2.56 eV) of the 3.6% N-doped SWSiCNT was found to be greater than the reported 1.23 eV for hydrogen evolution [38] by water splitting, and this presents overpotential to overcome the energy variation of the SWSiCNT photocatalyst. It also falls within the range of 2.2 eV to 2.8 eV. Additionally, N-doping promoted a decrease in charge carrier concentration which also reduces electron-hole recombination. As a result of this, the electronic motion becomes limited and trapped by N-impurities. Therefore, hydrogen reduction can take place by the photo-generated electrons of the 3.6% N-doped SWSiCNT. Moreover, the potential of the VBM was found to be -0.4 eV which is lower than -0.77 eV for normal OER [39]. The CBM was also analyzed and found to be 2.1 eV which is also greater than HER [40] potential of 0.02 eV [23], thus indicating that hydrogen can be liberated well on the active sites of N-doped SWSiCNT (3.6%). Based on this, the 3.6% N-doped SWSiCNT demonstrated itself as a better photocatalyst for hydrogen evolution via the water-splitting process.

The widening of the band gap of the N-doped SWSiCNT was due to the atomic orbitals of hydrogen atoms transforming into a series of energy levels as a result of quantum mechanical occupations of the equilibrium band by many electrons. It is also due to interactions between lattice phonons of the pure SWSiCNT and free holes of N atoms as a result of increased charge carrier concentration [41]. Furthermore, the carrier concentration and electrical conductivity of SiCNT decrease with pentavalent nitrogen atoms due to intra-atomic cluster [42]. As presented in Figure 5(c), an increase in the concentration of N atoms further broadened the band gap to the tune of 4.3 eV which is outside the standard range for photocatalysts to make perfect interactions with photons in the visible region. Therefore, it can be reported that the N-doped SWSiCNT photocatalyst performs better when 3.5% of N-dopant is used to dope one unit cell of the nanotube. Based on this, we can report that N-doped SWSiCNT can be a better water-splitting photocatalyst when the minimum energy state in the conduction band occurs at the conduction band edge of 2.1 eV and the maximum energy state in the balance band occurs at the band edge of 0.4 eV.

Analysis of the variation of the dopant concentration with the calculated band gaps of the SWSiCNT system was performed appropriately and obtained results are presented in Figures 6(a) and (b). As shown in Figure 6(a), both GGA and G_0W_0 results revealed that the band gap of pristine SWSiCNT broadens as the N-concentration increases. Compared with Figure 6(b), it can be seen that one atom of Nitrogen corresponds to 3.6% while two atoms of nitrogen correspond to 7.4%. Therefore, to get optimum performance by SWSiCNT photocatalyst, the sample shall be prepared such that one N atom per unit cell of SWSiCNT is obtained.

In order to know the degree of occupation and the density of electrons in a given state, it is necessary to obtain a detailed description of its electronic density of states (DOS), which is possible by analyzing the number of different energy states at a given energy level. Figures 7(a), 7(b) and 7(c) show the overall density of states for both pristine and N-doped SWSiCNT structures under different concentrations. Although both systems registered empty states at the Fermi level, the probability of finding electrons at the Fermi level is higher for the pristine SWSiCNT than for the N-doped SWSiCNT. Therefore, the band gaps in the doped systems are wider. In addition, a higher intensity of the DOS in Figure 7(b) indicated that the energy populations in the doped system are higher than in the pristine system. Also, the shapes of the DOS patterns are almost similar except in the area corresponding to 5 eV in Fig. 7(b), which has a lower intensity than the same 5 eV area in Figure 7(a). Because at this point in the conduction band, N atoms took up space for themselves. Therefore, the band gap opening in the doped system was due to the effects of N atoms in the conduction band. To obtain different orbital contributions to the photocatalytic potentials of the N-doped SWSiCNT, PDOS calculations were performed and the results are shown in Figure 8. Although N orbitals showed the highest valence band occupation, it has no contribution between valence band maxima (VBM) and conduction band minima (CBM). C orbital behavior was observed very close to the Fermi level near the VBM and showed no contribution in the conduction band. Regarding Si orbitals, clear features were observed in both VBM and CBM. Therefore, the widening of the band gap of the N-doped SWSiCNT is due to the blue shift by Si orbitals as a result of the occupation of spaces in the conduction band by N orbitals.

To further analyze the electronic behaviors of the investigated systems, we report the effects of tensile pressure (stress) and Young's modulus on the band gaps of the SiC nanotube systems. Results of the analysis are presented in Figures 9(a) and 9(b). It has been observed that the band

gap of the SWSiCNT photocatalyst was found to decrease from 4.3 eV (3.35 eV with GGA) to 1.25 eV (1.1 with GGA) with external pressure from 10.51 GPa to 11.12 GPa because when external pressure is increased, gamma states are produced into the energy band gap [43]. Figure 9(b) also revealed that Young's modulus increases with decreasing band gap, which agrees well with the reported mechanical behaviors of carbon nanotubes photocatalysts [44]. Therefore, the nanotube structure doped with 7.4% N atoms demonstrated higher mechanical strength while pristine SWSiCNT showed the lowest mechanical strength.

3.3 Optical properties

Analysis of an optoelectronic system cannot be completed until there is an adequate description of its behavior related to the interactions with electromagnetic waves. For an electronic system, the optical response to the incident electromagnetic waves is determined from the real and imaginary dielectric functions given by [45], as expressed by equation (6),

$$\varepsilon(\omega) = \varepsilon_1(\omega) + \varepsilon_2(\omega) \quad (6)$$

Where $\varepsilon_1(\omega)$ is the real part and $\varepsilon_2(\omega)$ is the imaginary part of the optical dielectric constant. The real part of the dielectric function describes the photon dispersion, while the imaginary part of the dielectric function describes the photon absorption. In the present study, we consider the imaginary part of the dielectric function to analyze the amount of solar energy absorbed by the pristine and doped SWSiCNT structures under different nitrogen dopant concentrations. For these studies, the optical absorption spectra were examined in both parallel and perpendicular directions. Furthermore, the studies were conducted using the random phase approximation method which takes care of the electron-electron interactions only. Figure 10(a) shows the optical absorption spectra of the pristine SWSiCNT with respect to the imaginary dielectric function. As can be seen, pristine SWSiCNT showed higher energy absorption in directions parallel than perpendicular to the nanotube axis. Few absorption peaks also appeared at 1.3 eV and 2.1 eV in the perpendicular direction. Based on the different behaviors observed by nanotubes in different directions, it can be called anisotropic. High intensities were recorded at 5.1 eV in both directions, corresponding to the ultraviolet range. 5.1 eV is also higher than the experimental optical gap of efficient photocatalysts, hence the pure (7, 7) SWSiCNT is considered an unsuitable photocatalyst. The N-doped SWSiCNT photocatalyst shown in Fig. 10(b) showed good performance, consistent with other perfect photocatalysts [46] [47]. It absorbs a high amount of energy at a band edge of 2.7 eV, which is in the visible range. In

addition, its absorptions are both parallel and perpendicularly symmetric, with a high absorption being recorded in parallel. Absorption at 2.7 eV means that the doped SWSiCNT interacts with visible electromagnetic energy from 2.2 eV to 2.8 eV range for photocatalysis. Therefore, the optimized system of N-doped (7, 7) SWSiCNT can be considered a better photocatalyst for the whole water-splitting technology. Figure 10(c) revealed different behavior of the N-doped SWSiCNT system when the percentage doping was increased to 7.4%. A new and smaller peak can be seen at 3.89 eV as a result of increased doping. The material also absorbs a higher amount of photons in the ultraviolet region which accounts for only 3% of solar radiation. Therefore, the 7.4% N-doped SWSiCNT is regarded as an unsuitable photocatalyst but can be used for other applications such as medical treatments using UV light. The energy loss function analysis was performed on all systems examined. Figure 10(d) shows that pristine SWSiCNT neither emits nor absorbs photons below 5 eV. More energy losses were also recorded in the UV range. Relative to 3.6% N-doped SWSiCNT, there were significantly lower electron energy losses in the range corresponding to 2.7 eV, which corresponds to the obtained optical band gap of this photocatalyst. This behavior indicated that higher photon energy was absorbed in this region, which agreed well with the absorption spectra previously reported in Figure 10(b). Figure 10(f) showed higher energy loss than 3.6% SWSiCNT, which means it showed poor absorption in the visible range.

A summary of the obtained characteristics of the investigated systems can be found in Table 3. As can be seen, the N-doped SWSiCNT photocatalyst performs efficiently when 3.6% of N dopant is used per unit cell of the pristine SWSiCNT because the N atoms have strong columbic interactions with Si and C atoms [48]. In this concentration, more photon energies were absorbed, interactions took place in the visible range and both electronic and optical band gaps correlated very well with the obtained theoretical data. Analysis of absorption energy presented in Table 4 also revealed that 3.6% N-doped SWSiCNT absorbs more energy than the pristine and 7.4% N-doped SWSiCNT systems. This Table also shows that atoms in pristine SWSiCNT are more bound together than in the doped systems. Table 5 presents various reports from some available literature regarding materials' photoactivity in the water-splitting process. It has been found that C-doped TiO₂ photocatalyst [49] and the results from current work presented a very good photocatalytic performance. Additionally, the optical and electronic band gaps of these materials were in good agreement with each other. Therefore the ability of the Si-doped SWZnONT can be enhanced by the photo-generated holes in the Si-doped ZnO nanotube

surface. Based on this, hydrogen can be produced when protons produced by water splitting penetrate the Si atom and react with the photo-generated electrons on the ZnO nanotube.

In terms of potential application areas, Table 6 presents the data obtained from the previous work in comparison with the current work. As shown, several works were performed on either the pristine SiCNT or the doped systems of SiCNTs. However, its potential to serve as a photocatalyst to produce hydrogen under solar irradiation has not been explored except in the current work. Therefore, the novelty of this work offers more possibilities to use the versatile properties of these nanotubes for a wide range of applications.

4. Conclusions

This study reported the photocatalytic properties of the nitrogen-doped single-walled silicon carbide nanotube based on the analysis of structural, elastic, electronic and optical features of this semiconductor. All calculations for the aforementioned properties of the investigated systems were done using the popular density functional theory as implemented in quantum ESPRESSO and Yambo codes. Doping of the pristine SWSiCNT system was performed using different concentrations of nitrogen atoms. It has been observed that the diameter of the SWSiCNT decreased from 30.01 Å to 29.51 Å and 29.02 Å when 3.6 wt% and 7.4 wt% N dopants were used. Additionally, it was found that the electronic band gap varies directly with N concentration and inversely with tensile pressure and Young's modulus. Better photocatalytic performance (energy range of 2.2 eV – 2.8 eV) was found with the 3.6% N-doped SWSiCNT which demonstrated electronic and optical band gaps of 2.56 eV and 2.70 eV, respectively. Finally, the observed structural, mechanical, electronic and optical behaviors demonstrated by the N-doped SWSiCNT exposed it as a better photocatalyst for hydrogen production under the visible light/energy spectrum. On the final note, it is recommended that these materials should be synthesised and analysed in order to identify any potential differences between experimental and theoretical data.

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Ethics

Procedure for data collection and study protocols were permitted by Bauchi State University centre for excellence research and innovation, and was conducted according to the guidelines declared. Also no human parts were used in the conduct of this research.

Data Availability Statement: All data that we used in this manuscript regarding quasi-particle energies, bands structure, DOS and optical spectra analysis were analysed and obtained with the quantum ESPRESSO codes which are freely available at www.quantum-espresso.org. Correction to quasi-particle interactions were done via Yambo interacting codes at www.yambo.org. We have optimized the geometric system of gallium nitride nanotube by using ‘tubegen’ database which is freely available at <https://turin.nss.udel.edu/research/tubegenonline.html>

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Figures

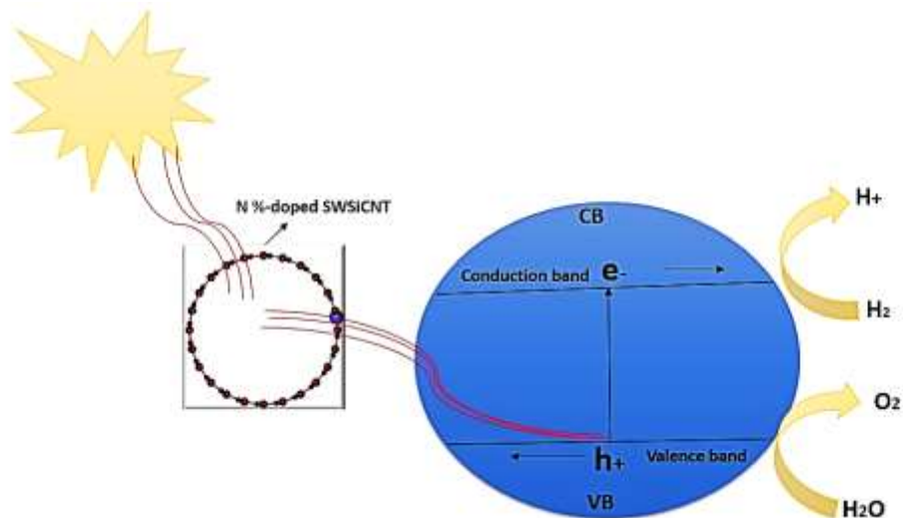


Figure 1. Graphical representation of the photocatalytic process of N-doped SWSiCNT

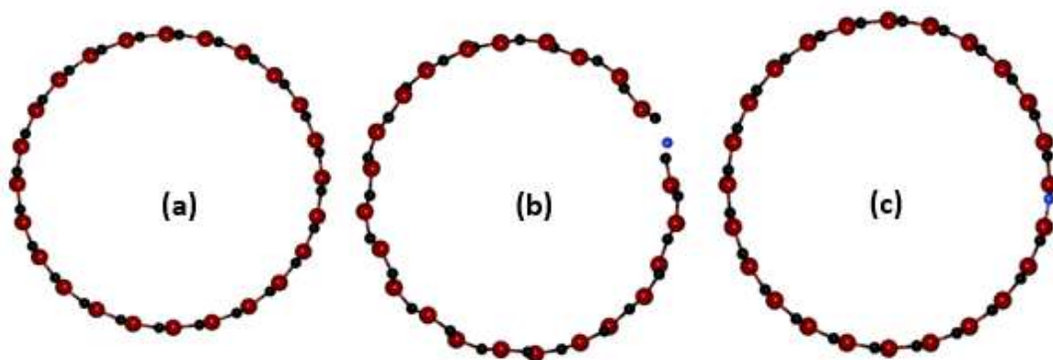
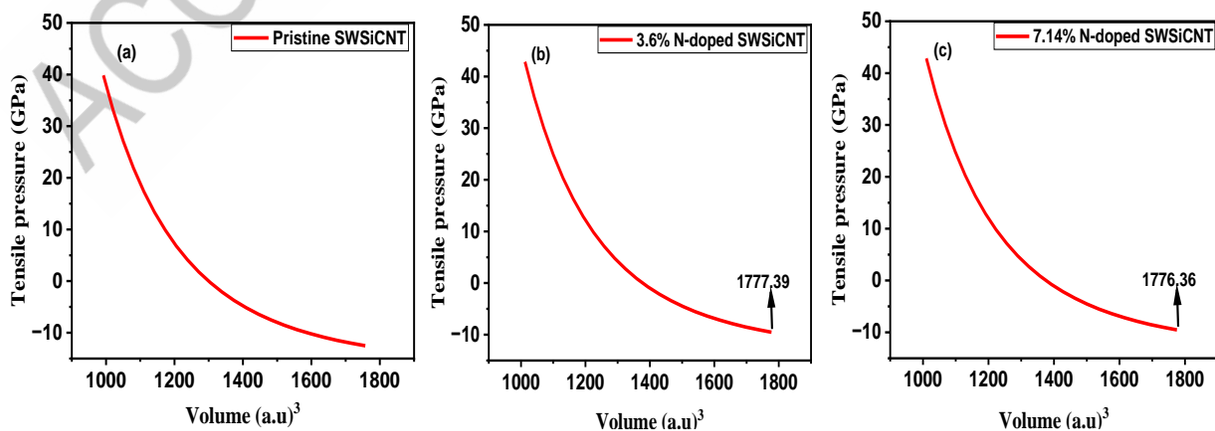


Figure 2. The optimized structures of (a) pristine SWSiCNT, (b) C replaced N-doped SWSiCNT and (c) Si replaced N-doped SWSiCNT



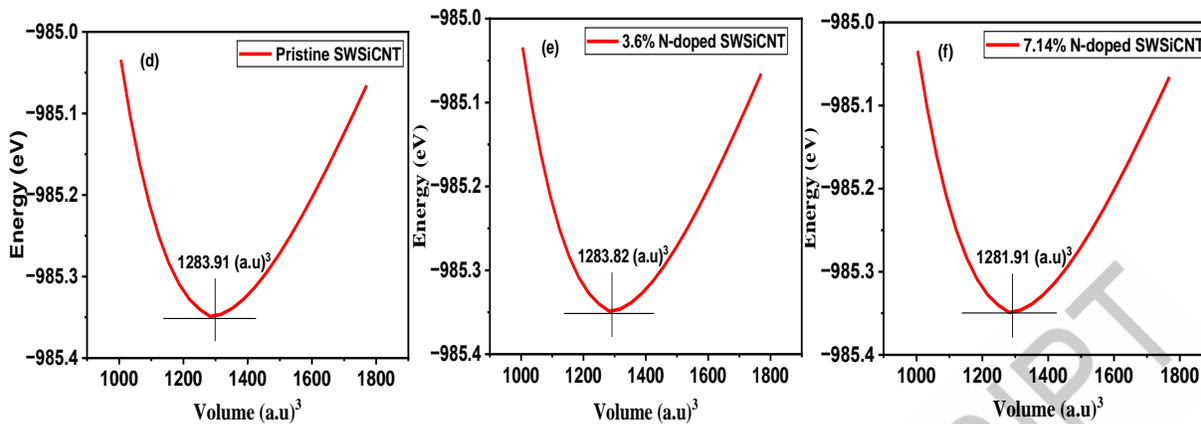


Figure 3. Effects of tensile pressure on the volume of (a) pristine SWSiCNT (b) 3.6%N-doped SWSiCNT (c) 7.4% N-doped SWSiCNT, and effects of Total energy on the volume of (d) pristine SWSiCNT (e) 3.6% N-doped SWSiCNT and (f) 7.4% N-doped SWSiCNT

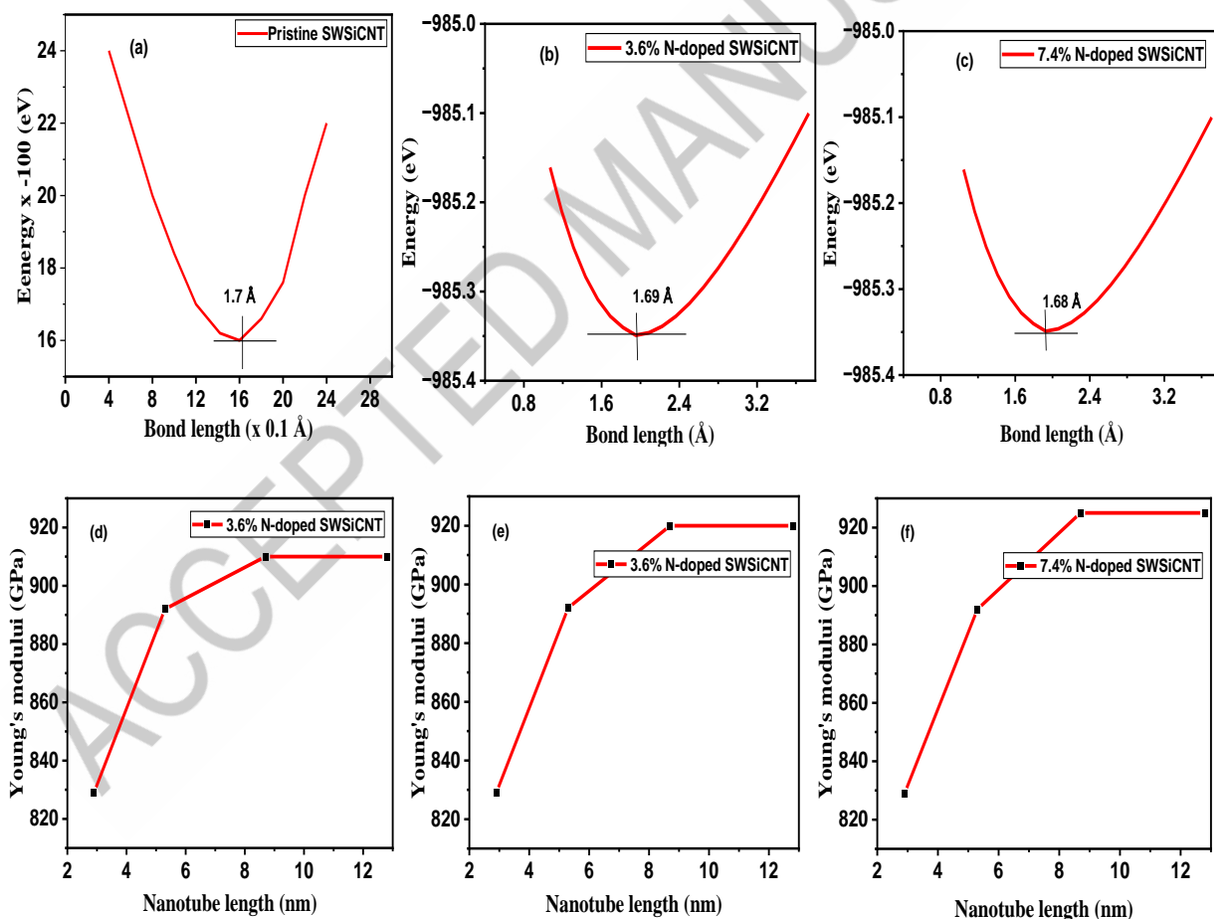


Figure 4. Effects of bond energies on bond lengths of (a) pristine SWSiCNT (b) 3.6% N-doped SWSiCNT, (c) 7.4% N-doped SWSiCNT. The calculated Young's moduli for (d) pristine SWSiCNT, (e) 3.6% N-doped SWSiCNT and (f) 7.4% N-doped SWSiCNT

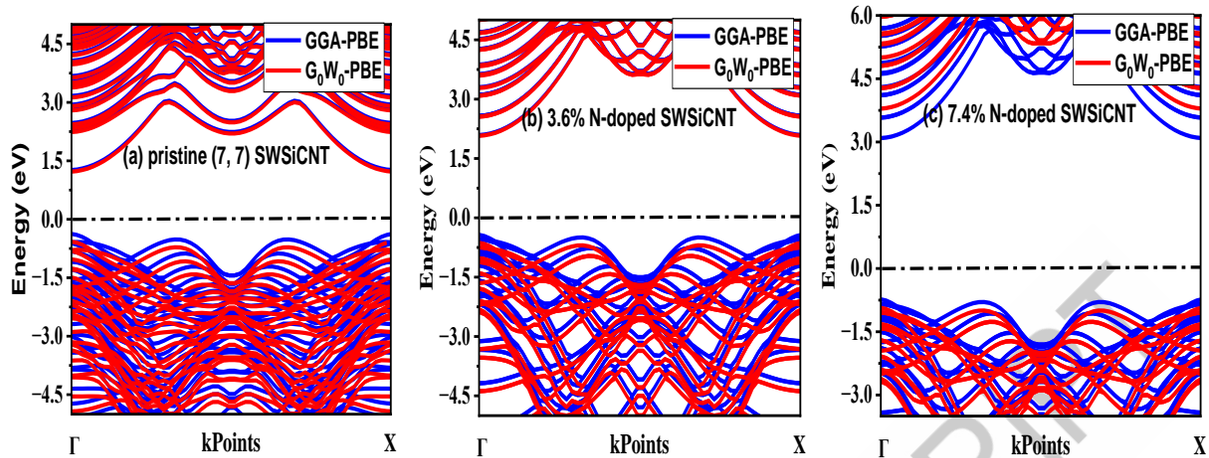


Figure 5. Electronic band gaps of (a) pristine SWSiCNT (b) 3.6% N-doped SWSiCNT and (c) 7.4% N-doped SWSiCNT.

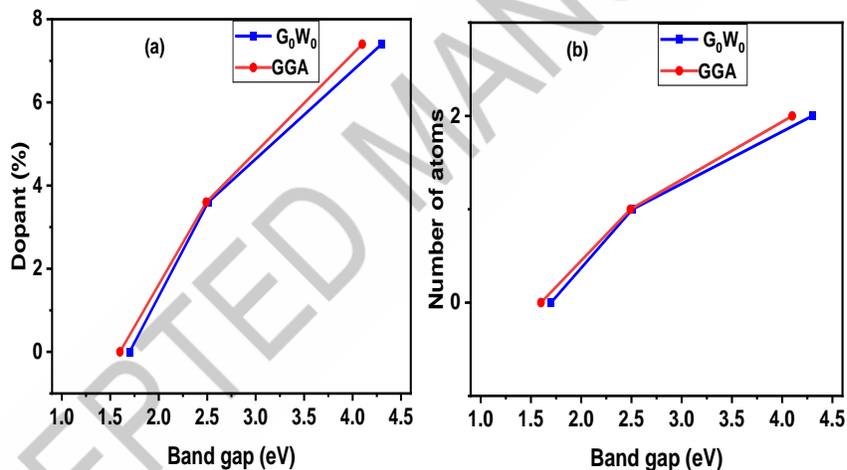


Figure 6. (a) Variation of N concentration with a band gap of SWSiCNT (b) variation of number of N atoms with a band gap of SWSiCNT

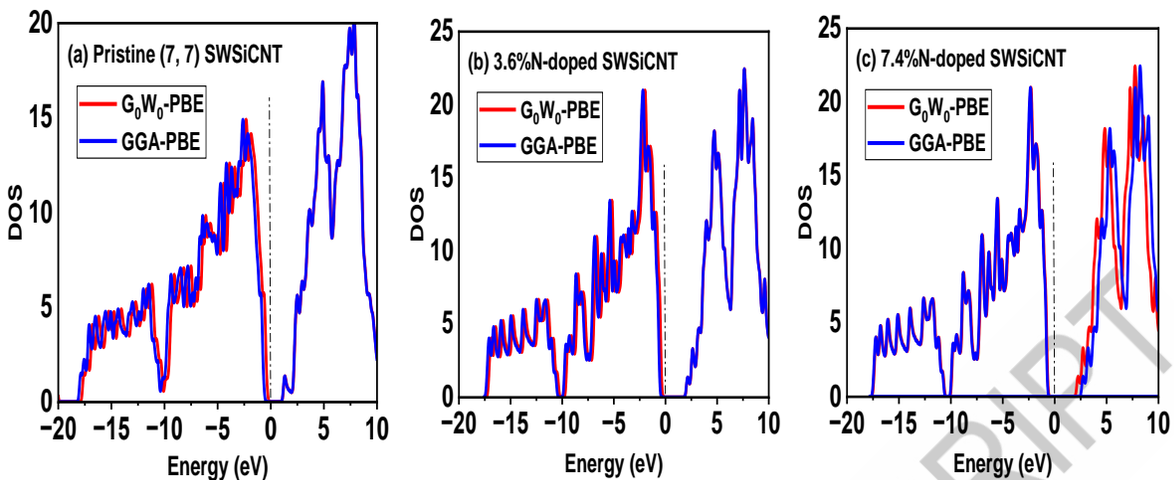


Figure 7. DOS of (a) Pristine SWSiCNT (b) SWSiCNT doped with 3.6% wt N (c) SWSiCNT doped with 7.4% wt N

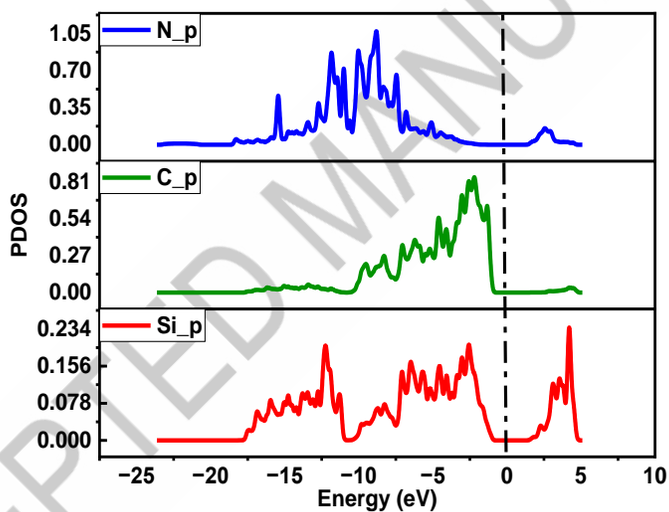


Figure 8. PDOS of the 3.6% N-doped SWSiCNT photocatalyst

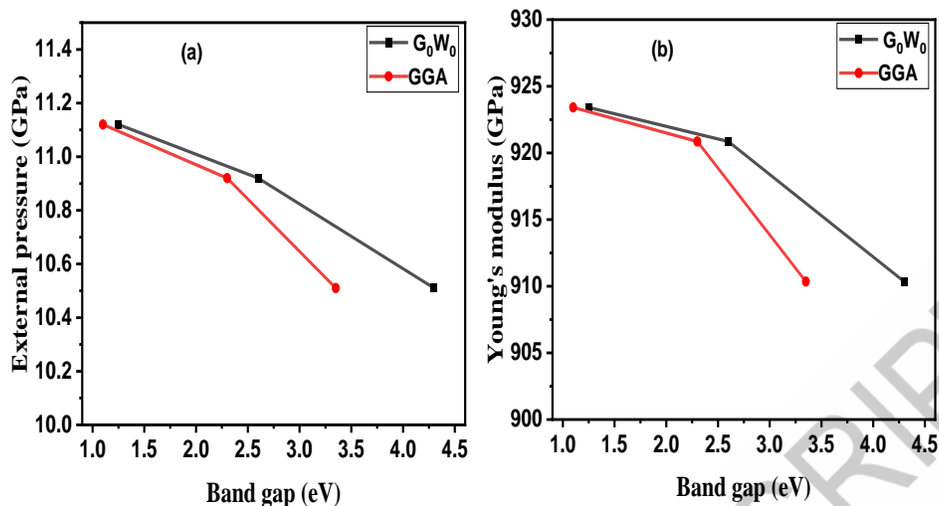


Figure 9. Band gap variation with (a) tensile pressure and (b) Young's modulus of SWSiCNT photocatalyst under N doping

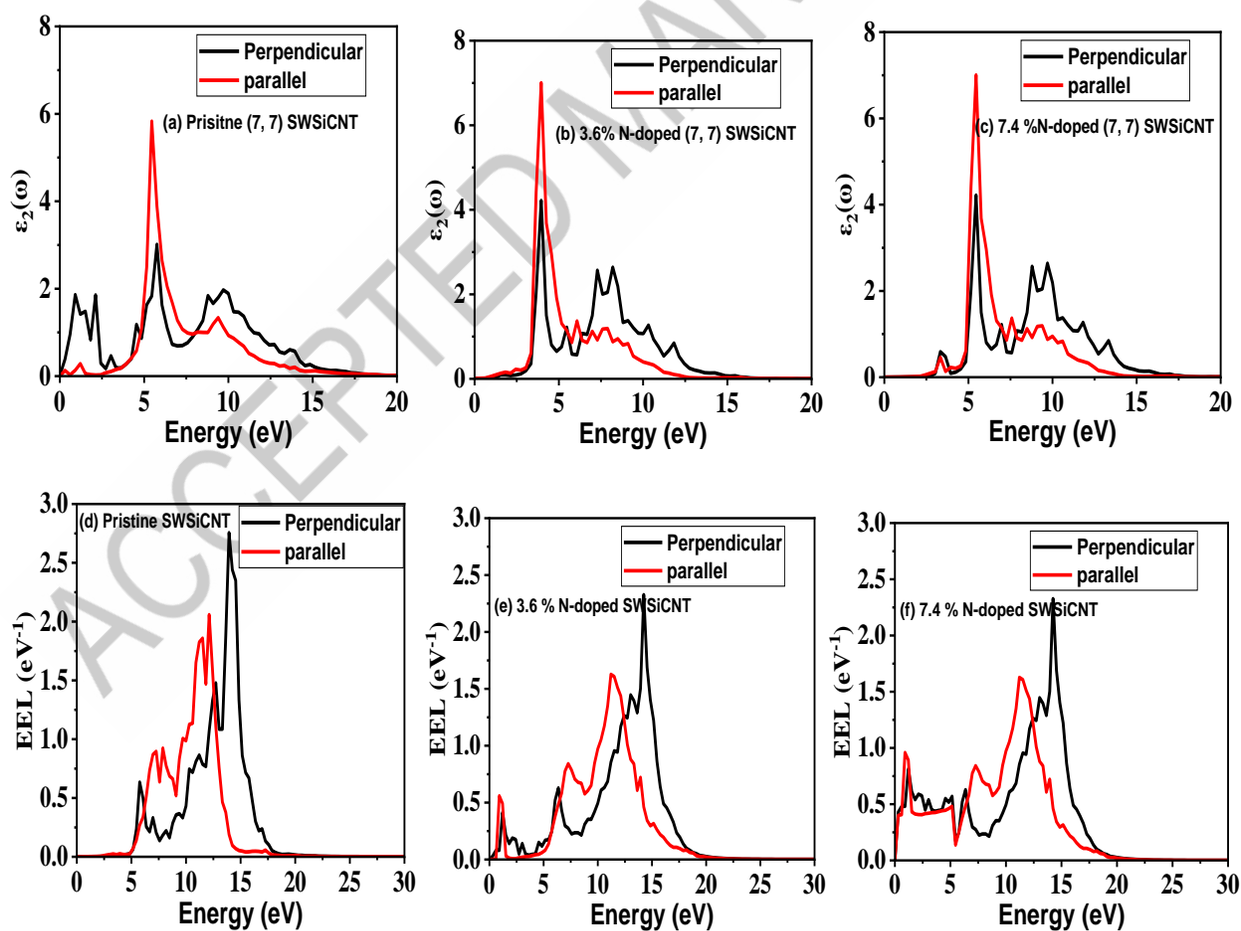


Figure 10. Photon absorption by (a) pristine SWSiCNT, (b) 3.6% N-doped SWSiCNT and (c) 7.4% N-doped SWSiCNT photo-systems. Electron energy loss spectra of (d) Pristine SWSiCNT (e) 3.6% N-doped SWSiCNT and (f) 7.4% N-doped SWSiCNT

Tables

Table 1. Formation energy distribution of the investigated systems

Material	Substitution	Type of defect	Formation energy (eV)	Diameter (Å)
Pristine SWSiCNT	No substitution	No defect	12.34	30.01
3.6% SWSiCNT	N-doped N atoms replaced	Si N_{Si}	10.36	29.51
7.4% SWSiCNT	N-doped N atoms replaced	Si N_C	9.42	29.02

Table 2. Calculated elastic parameters of the investigated systems

Material	Bond length (Å)	Bond energy (eV)	Volume (a.u) ³	Stress (GPa)	Young's Modulus (GPa)
Pristine (7, 7) SWSiCNT	1.71	-985.35	1283.91	11.12	910.35
3.6% N-doped SWSiCNT	1.69	-985.34	1283.82	10.92	920.86
7.4% N-doped SWSiCNT	1.68	-985.30	1282.91	10.51	923.41

Table 3. Summary of the obtained characteristics of the investigated systems.

Material	The electronic band gap (eV)	The optical band gap (eV)	Dielectric absorption (a.u)	Remark
Pristine SWSiCNT	1.7	1.6	6.10	Unsuitable
3.6% N-doped SWSiCNT	2.56	2.70	7.42	Suitable
7.4% SWSiCNT	4.30	4.81	7.32	Unsuitable

Table 4. Effects of absorption and binding energies

Absorbing surface	Bond angle (degrees)	Bond length (nm)	Absorption energy (eV)	Binding energy (Ve)	Reference
Pristine SWSiCNT	102.02	1.71	-1.81	-11.55	[50]
3.6% N-doped SWSiCNT	102.21	1.69	-1.97	-11.48	[51]
7.4% N-doped SWSiCNT	101.91	1.68	-1.91	-11.53	[52]

Table 6. Previous work Vs current work

Material	Method	ϵ_2 (a.u)	The band gap (eV)
N-doped TiO ₂ nanotubes [53]	Experimental hydro-thermal process	19.11	2.7
C-doped TiO ₂	Experimental synthesis method	18.52	2.4
BN-CNT heterostructure [54]	DFT- first principles	Not reported	2.51
Co-doped CNTs [55]	DFT-first principles	23.20	2.55
3.6% N-doped SWSiCNT- current work	DFT with absorption optical spectra analysis	21.53	2.56

Table 6. A critical comparison between some selected past works and the current work

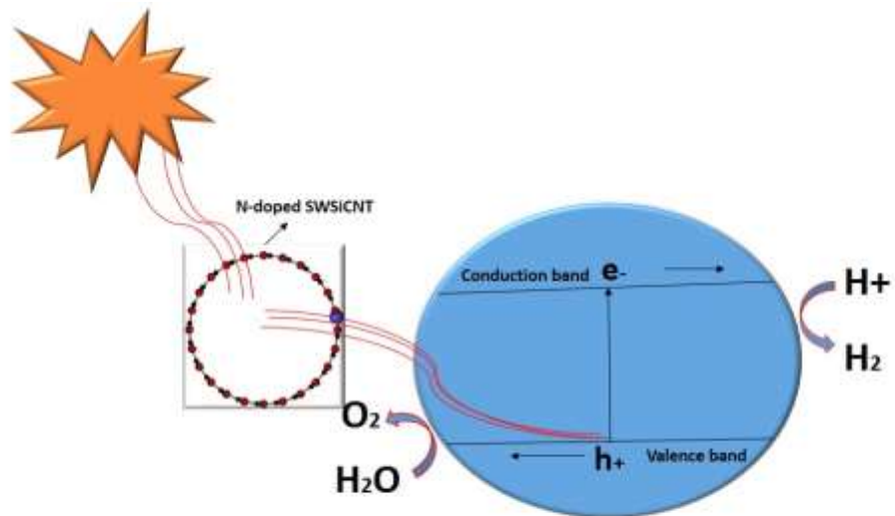
Materials	Analysis	Application	Reference
SWSiCNT	Zheng et al. only report the field emission potentials of the SWSiCNT without considering chirality. While the present work investigated the N-doped system of SWSiCNT with (7, 7) chirality for the production of hydrogen.	Does not specifically state the most suitable area of application of the investigated nanotubes while current work found the material worthy of use as a photocatalyst.	[56]
N-doped (6, 0) and (4, 4) SWSiCNT	Results demonstrated that N impurities do not alter the structure of SWSiCNT which agreed well with the current work. Our findings also revealed that the characteristic surface reactivity of the (7, 7) SWSiCNT was generally influenced by doping of 3.6% nitrogen. However, our work is different because we have extended the studies to the mechanical and optical characteristics of the (7, 7) SWSiCNT.	Mahdaviani et al. investigated the surface reactivity of the N-doped SWSiCNT while the current work studied the photocatalytic potentials of N-doped SWSiCNT	[57]
SWSiCNT doped with B, N and O at once	Maghnaoui and Boufelfel studied the effect of magnetic moment on the doped SWSiCNT without a complete description of the electronic and optical properties. While the current work investigated the effects of different	Maghnaoui and Boufelfel did not specify possible areas of applications of their obtained results. The current work recommended the N-doped SWSiCNT as good photocatalysts for hydrogen production.	[58]

Metal decorated SiCNT	nitrogen concentrations on the doped SWSiCNT which was found to make a significant effect on the electronic and optical properties of the nanotube. Results revealed that metal-decorated SiCNT can actively adsorb hydrogen in the visible region. In the current work, a non-metal doped (N-doped) SWSiCNT was found to absorb solar energy very well in the visible range.	Ram et al. recommended metal-doped SiCNT as a good candidate for hydrogen storage while the current work found N-doped (non-metal) SWSiCNT as a good candidate for hydrogen production. [59]
Hydrogen boride nanotubes with C, N and O decoration	Results by Nontawat et al. showed that N-doped hydrogen boride could be used to store hydrogen. However, the scope of our work is entirely on a different material and for hydrogen production not for hydrogen storage.	Doped hydrogen boride by Nontawat et al. was found worthy for water-sensing applications while the current work found N-doped SiCNT for water-splitting applications. [60]
Metal decorated SiCNTs	Results by Singh revealed that CO ₂ adsorption on Cu-decorated SiCNT undergoes spontaneous exothermic reaction.	Singh reported the possible potential of metal-decorated SiCNT for CO ₂ storage. Their work, however, found the material unsuitable for such application. In our work, we studied non-metal doped SiCNT as a catalyst for producing hydrogen only and it was found worthy applicable. [61]

Novelty statement

- This work reports fundamental photo performance properties of a new system, i.e., nitrogen-doped single-walled silicon carbide nanotubes (N-doped SWSiCNTs) as photocatalysts for overall water splitting for the first time. The observed structural, mechanical, electronic and optical behaviors demonstrated by the N-doped SWSiCNT exposed it as a better photocatalyst for hydrogen production under visible light, which indicates it a better candidate for photocatalytic applications such as water splitting, biodegradation, and wastewater purification.
- 3.6% N conc. has been found to be the suitable photocatalyst for water splitting.

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