



Article Effect and Characterization of Stone–Wales Defects on Graphene Quantum Dot: A First-Principles Study

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Abstract: A first principles based density functional theory (DFT) has been employed to identify the signature of Stone-Wales (SW) defects in semiconducting graphene quantum dot (GQD). Results show that the G mode in the Raman spectra of GQD has been red shifted to 1544.21 cm⁻¹ in the presence of 2.08% SW defect concentration. In addition, the intensity ratio between a robust low intense contraction-elongation mode and G mode is found to be reduced for the defected structure. We have also observed a Raman mode at 1674.04 cm^{-1} due to the solo contribution of the defected bond. The increase in defect concentration, however, reduces the stability of the structures. As a consequence, the systems undergo structural buckling due to the presence of SW defect generated additional stresses. We have further explored that the 1615.45 cm⁻¹ Raman mode and 1619.29 cm⁻¹ infra-red mode are due to the collective stretching of two distinct SW defects separated at a distance 7.98 Å. Therefore, this is the smallest separation between the SW defects for their distinct existence. The pristine structure possesses maximum electrical conductivity and the same reduces to 0.37 times for 2.08% SW defect. On the other hand, the work function is reduced in the presence of defects except for the structure with SW defects separated at 7.98 Å. All these results will serve as an important reference to facilitate the potential applications of GQD based nano-devices with inherent topological SW defects.

Keywords: density functional theory (DFT); Raman spectroscopy; Stone–Wales defect; electronic properties; infra-red spectra

1. Introduction

The successful experimental fabrication of the two dimensional (2D) nanostructure graphene was a path breaking event in the field of material science. The pure graphene sheet exhibits several unique optical, electronic and mechanical properties [1,2]. However, the implementation of pure graphene in the opto-electronic industry is restricted by its zero band gap nature. Various efforts have been made to tune the Dirac-like dispersion relation by means of external perturbations [3–7]. It is well established that electronic and optical properties are interdependent. Therefore, the modulation of band structure can also be realized from the optical responses [8] for different graphene like systems [9]. Another favorable approach in band gap engineering is to reduce the dimension of graphene by proper structural tailoring [10–13]. Graphene quantum dots (GQDs) are very small fragments of graphene, where the motion of electrons is confined in three spatial directions. The properties of these GQDs are very much different from that of graphene. The underlying reasons for such difference are generally localization of electrons because of the quantum confinement effect and the existence of edge states. The GQDs are generally semiconducting in nature and the band gap can

be tuned by varying the shape, size and surface chemistry [14–16]. Because of these tunable band gaps, GQDs can be used in photovoltaic devices [17,18]. Clearly, a graphene sheet does not allow photovoltaic applications because of the zero band gap nature. In addition, GQDs have also been observed to facilitate potential applications in organic LEDs [19], fuel cells [20], bio-imaging [21,22], bio-sensing [23,24], and environmental monitoring devices [25].

Point defects in graphene have drawn enormous interest because of their unavoidable existence in sample preparation [26–30]. The Stone–Wales (SW) defect [31,32] is a customary topological disorder in the graphene systems and its derivatives i.e., carbon nanotubes [33], nanoribbons [34] and fullerenes [35]. This involves rotation of carbon-carbon (C-C) bonds with respect to the midpoint by an angle $\pi/2$. As a result, four neighbouring hexagons are transformed into two heptagon-pentagon (7-5) pairs. These types of defects can also be artificially engineered in the laboratory [36]. The specific position of the defects can be identified with the help of transmission electron microscope (TEM) [36,37]. It is evident from previous reports that concentration as well as the specific positions of the defects crucially determine the hybridization and stability of the systems. These modifications have immense impact on the electronic band structure [38], and, hence, on the conductivity of the system [39,40]. In this context, it is worth mentioning that Lu et al. [41] have successfully transformed C₆₀ molecules into equal sized graphene quantum dots. Although SW defects are extremely common in the fullerene molecules [35], the effects of SW defects on graphene quantum dots have been less explored.

Raman spectroscopy [42] is a widely used non-invasive characterization technique for the carbon networks. The nature and concentration of defects on graphene like systems can be efficiently identified using this tool [43–46]. For example, the first order Raman spectrum of graphene exhibits single predominant stretching like mode called G band [42]. This mode is a signature of sp^2 hybridized C atoms in the system. The other defect related bands (D band) are due to second order Raman scattering process and it does not have any impact on the first order vibrational spectra. In addition, infra-red (IR) spectroscopy is another efficient tool for the characterization of carbon bonded materials. One can easily comment on the vibrational states of any system by studying the variation of IR frequencies of the incident electromagnetic radiation.

Motivated by the above discussions, we have systematically investigated the effect of SW defects on the vibrational and electronic properties of the graphene quantum dot (GQD). The effects have been explored for different concentrations and different positions of the SW defects. This paper has been organized as follows. First, the computational methodologies have been presented. In the next section, we have critically commented on the structural modifications due to the presence of SW defects in the system. Afterwards, we have extensively discussed the impact on the Raman and IR spectra of the GQD, followed by the variation in electronic properties i.e., dipole moment variation, effect on conductivity, work function, energy gap, etc. Finally, conclusions are drawn in Section 7. We expect that these results will broaden the possibilities of GQDs to be used in nano-electronic devices.

2. Computational Details

All the calculations have been performed under the framework of density functional theory (DFT) as implemented in the software Gaussian 09W [47]. The self-consistent ground state (true minima) has been achieved using Becke three parameters functional with Lee–Yang–Parr non-local electron correlation (B3LYP) [48,49] and the basis set 6-31G(d). We have used the optimized geometry for vibrational analysis with the same functional and basis set. The Hessian matrix (*f*) contains the second order partial derivatives of the potential (Φ) with respect to atomic displacements (*x*) from equilibrium positions as described in Equation (1)

$$f_{p,q} = \left(\frac{\partial^2 \Phi}{\partial x_p \partial x_q}\right)_{equilibrium}.$$
(1)

Order of the matrix depends on the number of atoms (N) in the system. These force constants are further rewritten in terms of mass (m) weighted Cartesian coordinates (f_{mwc}) as shown in Equation (2)

$$f_{mwc;p,q} = \left(\frac{\partial^2 \Phi}{\partial X_p \partial X_q}\right)_{equilibrium}.$$
(2)

Here, $X_p = \sqrt{m_p} x_p$ and $X_q = \sqrt{m_q} x_q$. The square roots of the eigenvalues of the above 3 N × 3 N matrix give rise to the fundamental frequencies of the system. The translational and rotational modes are then separated out and Schmidt orthogonalization gives rise to the remaining 3 N–6 N vibrational modes. All these frequencies are verified to be real for the stability purpose. Raman intensities (I_i) have been calculated from the Raman scattering activity (S_i) for the i-th normal mode using Equation (3)

$$I_{i} = \frac{f(\bar{v}_{0} - \bar{v}_{i})^{4}S_{i}}{\bar{v}_{i}e^{-\frac{h\bar{v}_{i}}{kT}}}.$$
(3)

Here, $\bar{v_0}$ is the wavenumber (WN) of the incident photon, $\bar{v_i}$ is the WN of i-th normal mode, *h* is the Planck's constant, *c* is the speed of light, *k* is Boltzmann constant and *f* is the normalization factor. This calculation has been performed at temperature 293.15 K and an exciting wavelength 785 nm. The Raman and IR spectra are plotted with Lorentzian envelope with full width half maximum (FWHM) equal to 5 cm⁻¹ for better understanding. Besides, a scaling factor 0.9614 has been employed to obtain the actual WN from the DFT generated data in accordance with our basis/functional pair [50].

It is worth noting that the electronic ground state of the systems gets perturbed due to the first order Raman scattering process. However, the electronic state returns to the same ground state with different vibrational level after each event. Therefore, we have also explored the density of states (DOS) of the systems with the help of GaussSum 3.0 software [51]. HOMO and LUMO are the highest occupied molecular orbital and lowest unoccupied molecular orbital of the systems and the difference between their energies ($E_{LUMO} - E_{HOMO}$) is essentially the band gap of the systems. We have additionally calculated the activation energy or work function of the systems from the difference between energies of the vacuum level and the HOMO ($W = E_{vacuum} - E_{HOMO}$).

3. Structural Modifications

It is well established that electron confinement effect reduces with increasing size of graphene clusters and, for a system with 48 carbon (C) atoms, the bulk property predominates [16].

Motivated by this, we have chosen the graphene cluster with 48 C atoms to investigate the effect of SW defects on its vibrational and electronic properties. Note that only the middle bond has been rotated for the SW defect concentration 2.08% and, for 4.17% defect concentration, different possible positions have been indicated in Figure 1a. In Figure 1a, we have intentionally avoided the terminal bonds as defect positions for stability reasons. Therefore, the only choices to obtain 4.17% defect are 'ab', 'ac', 'ad', 'ae', 'af' and 'dd''. Other possibilities have been declined due to symmetry arguments. In addition, the ground state geometry reveals that, similar to 'ae', the 'af' system (indicated in Figure 1a) also introduces tetragon into the structure (shown in Table 1), which also reduces the stability of the system.

Table 1. Table for the number of non-hexagonal polygons in the systems, distance between two defect positions and the total energy E of the systems.

Systems	Tetragon	Pentagon	Heptagon	Octagon	Distance (Å)	E (eV)
pristine	0	0	0	0	-	-50,074.06
s-SW	0	2	2	0	-	-50,070.92
ab	0	2	2	0	2.36	-50,069.59
ac	0	3	3	0	3.00	-50,069.01
ad	0	4	2	1	3.99	-50,068.48
ae	1	2	4	0	2.82	-50,067.24
dd′	0	4	4	0	7.98	-50,068.52



Figure 1. (a) The schematic diagram of the optimized GQD with 2.08% defect concentration. Any one of the bonds b, c, d, e, f, d' can be additionally rotated to achieve 4.17% defect concentration; (b) Bond lengths and angles of the 2.08% SW defect concentration; (c) The structure ae has 4.17% defect concentration and possesses tetragon in the structure.

It is clear from Figure 1b that the optimized length of the middle bond reduces from 1.41 Å to 1.35 Å after rotation, whereas the bond 'd' or 'd'' gets a little elongated to 1.43 Å during this process. The SW defects, however, do not alter the total number of electrons in the system. Therefore, the total

energy will be a crucial parameter to compare their relative stabilities. We have observed that the total energy of the system increases by an amount of 3.15 eV after the incorporation of a 2.08% SW defect into the pristine cluster. We have further compared their respective dimensions. It is notified that the distance between terminal C atoms along the direction parallel to 'a' has been increased from 9.91 Å to 10.54 Å, while that along 'd' contracted from 15.56 Å to 15.15 Å. This change is due to the generation of tensile and compressive stresses into the system along respective directions as a secondary effect of SW defects. This SW defect generated stress is responsible for the increase of the total energy of the system with a 2.08% SW defect. The stress in the system is further enhanced for 4.17% defect concentration. As a result, the systems undergo buckling while trying to minimize the effect of this additional stress. This supports the fact that presence of SW defects in the system has the tendency to form non-planar nanostructures due to the intrinsic rumpling in the system [52]. The effects of topological SW defects on the systems are listed in Table 1 for better understanding. It is to be noted that pristine structure exhibits the highest stability and the stability decreases with increasing defect concentrations. Two structures 'ab' and 'ac' possess the least buckling and have the highest stability, whereas the 'ae' structure shown in Figure 1c contains tetra-ring in the network and is the least stable structure among all.

4. Raman Spectra

In the 1st order Raman scattering process, an incident photon interacts with the system and the system radiates a photon with some other frequency. As a result, the system changes its initial vibrational state. The frequency difference between the incident and emergent photon is thus a measure of long wavelength lattice vibration. Using these frequencies, popularly known as Raman shift, the systems can be efficiently characterized. In this context, it is important to note that the IR absorption spectra also provide the same information as we will see in the next section. However, Raman spectra are more handy tools for the high frequency lattice vibrations. The dangling bonds of the terminal C atoms are passivated with H atoms for stability purposes [53]. The Raman spectra of the graphene cluster exhibit a prominent C-C stretching mode at 1562.22 cm⁻¹. This mode is similar to the characterizing G mode vibration of an sp^2 hybridized graphene sheet. In addition, there are some other modes present in the Raman spectra of the pristine system. These are the stretching modes at 1303.37 cm⁻¹, 1326.44 cm⁻¹, 1336.40 cm⁻¹, 1337.82 cm⁻¹ etc. In addition, we have observed a low frequency mode at 228.79 cm⁻¹ attributed to the out of phase contraction and expansion of the whole system. The edge atoms as well as the terminal C-H bonds also contribute in the Raman pattern. We have intentionally avoided the discussions related to the less important high frequency modes caused by the vibration of C-H bonds only. We have further explored the effect of 2.08% SW defect on the Raman spectra as depicted in Figure 2.

The G mode of the pristine structure has been observed to be slightly red shifted to 1544.21 cm⁻¹ in the presence of the defect. Similar to the pristine structure, this mode is also prominent. Therefore, the SW defect in the system acts as an active scattering center for the Raman process. For better understanding, the nature of vibration corresponding to this G mode has been well described in Figure 3a.

Similar to the previous case, there are many other additional stretching modes at 970.36 cm⁻¹, 1276.38 cm⁻¹, 1413.21 cm⁻¹, 1575.89 cm⁻¹, etc. in the Raman spectra. In addition, a mode at 1504.42 cm⁻¹ needs special attention due to the atomic vibrations perpendicular to unrotated bonds. The low frequency mode associated with the out of phase contraction and expansion of the whole system has also been identified for the defected structure. This mode gets softened to the value 224.77 cm⁻¹ for a 2.08% SW defect.

We have calculated the intensity ratio of the low frequency mode to the G mode before and after the introduction of defect. This ratio for the defected structure (1.18) is smaller than that of the pristine structure (1.34). There is a mode at 1138.31 cm⁻¹ corresponding to the pendulum like oscillation of the rotated bond. In addition, there is a mode due to the solo contribution of the rotated bond at 1674.04 cm⁻¹ as depicted in Figure 3b. This mode is the maximum frequency mode due to the vibration of C atoms of the structure. The same phenomenon has been previously observed in the case of graphene sheet with a 2.04% SW defect [54]. Absence of out of plane vibrational modes in Raman spectra strongly supports the stability of both the systems. The Raman spectra have been considerably modified with increasing defect concentration (4.17%). In addition, the Raman pattern is also very sensitive to the specific position of the defect in the system. It is evident from Figure 4 that the mode near 224.77 cm⁻¹ is almost robust for all the systems.



Figure 2. Comparison between the Raman spectra of pristine graphene cluster and 2.08% SW defect.

On the other hand, the G like stretching mode with the maximum scattering activity are situated at 1541.29 cm⁻¹, 1513.73 cm⁻¹, 1528.78 cm⁻¹, 1537.24 cm⁻¹ and 1535.51 cm⁻¹ for 'ab', 'ac', 'ad', 'ae' and 'dd'', respectively. It is clear from the above data that the G band for the most stable structure 'ab' remains almost unaltered. On the contrary, the Raman spectra of the least stable structure 'ae' exhibit several low intense modes with comparable intensities [16]. The mode (1674.04 cm^{-1}) due to the solo contribution of the SW defect has been found to be blue shifted to 1830.66 cm⁻¹ for the 'ae' system. This shift is because of the change in electrical polarizability of the system in the presence of the tetragon. The structure 'dd' has however been obtained by creating two 2.08 % SW defects at large (7.98 Å) distance as shown in Table 1. Therefore, we have compared the Raman spectra of 2.08% SW defect with 'dd'. The G band of the structure is at 1535.51 cm⁻¹, which is red shifted compared to the s-SW structure. In addition, two new low frequency modes at 22.65 cm⁻¹ and 43.08 cm^{-1} have been observed in case of dd'. It is worth noting that these modes are torsional and bending modes, respectively. Therefore, the appreciable number of non-stretching modes for the dd' system establishes relatively less stability of the structure. In addition, a 224.77 cm^{-1} mode of the s-SW system is completely suppressed in dd' because of large buckling produced in dd' structure. Most interestingly, a mode at 1615.45 cm⁻¹ is identified as a solo stretching mode of two rotated bonds. This mode is red shifted compared to the similar mode of s-SW structure. These results help us to interpret that 7.98 Å is the minimum distance between two SW defects required to get the signature of two distinct defects separately. The assignments of different Raman active vibrational modes are listed in the supplementary document for convenience.



Figure 3. (**a**) Signature of the G mode vibration and (**b**) Solo contribution of SW defect at 1674.04 cm⁻¹ of graphene cluster with 2.08% SW defect.



Figure 4. Raman spectra of different possible systems with 4.17% SW defect concentration. The Raman spectra for 2.08% SW defect is also given for comparison.

5. IR Spectra

IR spectroscopy is also a very efficient tool to determine the structural details for characterization. The absorption bands of this spectra possess the information about the nature of vibration of different bonds. These vibrational modes are discrete and when the frequency of the incident photon matches with any one of these fundamental modes, they undergo absorption. The IR spectra can be classified in three frequency regions near IR (14,000–4000 cm⁻¹), mid IR (4000–400 cm⁻¹) and far IR (400–10 cm⁻¹). In this section, we have mainly explored the mid and far IR fingerprints of the SW defects in

graphene quantum dots. For better understanding, the IR spectra of all the systems are depicted in the supplementary document.

Most of the IR modes of pristine GQD have been observed to be red shifted in the presence of s-SW defect. For example, the mode corresponding to the maximum absorption is at 1604.75 cm⁻¹ for GQD, whereas the same for s-SW system is 1674.04 cm⁻¹. We have further explored the low intense absorption peak due to the solo contribution of the defected bond is at 1674.04 cm⁻¹. Additionally, some new IR active modes have been observed for the defected system. The maximum absorption peaks for the structures with 4.17% doping concentration are at 884.25 cm⁻¹ (ab), 873.54 cm⁻¹ (ac), 876.88 cm⁻¹ (ad), 1522.78 cm⁻¹ (ae) and 863.28 cm⁻¹ (dd'). Therefore, these systems can also be characterized with the help of IR spectroscopy. It is interesting that we have successfully identified the IR absorption frequency corresponding to the simultaneous stretching of two individual SW defects of dd' at 1619.29 cm⁻¹. Similar to the Raman case, this also supports the fact that 7.98 Å is the minimum distance for two SW defects in graphene for their distinct existence.

6. Electronic Properties

The anisotropic and electronic properties perturbed by the topological SW defects are discussed in this section. The importance of dipole moment is of immense help for determining related transport, chemical reactivity and opto-electronic properties of any system. It has been observed that the presence of SW defects affects the charge distribution and hence the dipole moment of the systems as listed in Table 2.

Table 2. Table for dipole moment (D), vacuum energy level (E_v), LUMO and HOMO energies, band gap ΔE (= E_{LUMO} (eV) – E_{HOMO}), and the work function W (= $E_v - E_{HOMO}$).

Systems	D (debye)	\mathbf{E}_{v} (eV)	E _{LUMO} (eV)	E _{HOMO} (eV)	ΔE (eV)	W (eV)
pristine	0	0.46	-2.94	-4.16	1.22	4.62
s-SW	0.0013	0.47	-2.82	-4.09	1.27	4.56
ab	0.8619	0.29	-2.86	-4.19	1.33	4.48
ac	0.9864	0.45	-2.88	-4.15	1.27	4.60
ad	0.4654	0.02	-2.84	-4.09	1.25	4.11
ae	1.4739	0.13	-2.88	-4.25	1.37	4.38
dd′	0.1537	0.54	-2.93	-4.20	1.27	4.74

It is clear from Table 2 that finite dipole moments have been induced in the otherwise non-polar system due to the presence of SW defects. The magnitude of dipole moment varies with concentration as well as distance between the defects. The charge density and direction of the dipole moment has been depicted in Figure 5. It is observed from Figure 5 that the charges are accumulated more in the left and right end of the systems.

This is because of the maximum value of local curvature near these two points. The 'ae' structure possesses the highest dipole moment. As an important note, the band gap of this structure is also maximum. In addition, 'ac' and 'ab' systems also exhibit appreciable dipole moments. Here, another important note is that the dd' structure has the minimum dipole moment among all 4.17% defected systems. This is due to the fact that the separation of SW defects has the maximum value in dd' structure. All these finite and appreciable values of dipole moment can attain the minimum requirement for any polar gas sensors [44,55,56].



Figure 5. Charge density plot of (**a**) s-SW and (**b**) as structure. The direction of the dipole moment has been shown with a red arrow. The image has been drawn with isovalue $0.01 \text{ e}^{-}/\text{au}^{3}$.

The electronic properties essentially depend on DOS near E_F of the systems. Not only that, the HOMO-LUMO gap in this regard plays a vital role in deciding appropriate opto-electronic properties and chemical stability of the systems. We have thus explored the electronic state of the systems by computing the DOS corresponding to different discrete molecular levels as shown in Figure 6. For better comparison, Fermi energy ($E_F \approx (E_{LUMO} + E_{HOMO})/2$) of the system is set to zero. Two systems ab and ac with 4.17% defect concentration possess a higher degree of stability and exhibit zero DOS at E_F similar to the pristine and s-SW defect case. Figure 6a reveals that the nature of DOS near E_F remains almost similar for pristine and s-SW structure.

In addition, the identification of HOMO-LUMO levels in the defected systems is also essential for designing opto-electronic devices. Therefore, the positions of the HOMO and LUMO orbitals of s-SW and dd' structures are given in Figure 7. It is observed that there are no LUMO orbitals in the C atoms situated on the principal axis (parallel to the bond 'a') for both the systems. On the contrary, HOMO levels are observed on the rotated C dimers and their neighbouring C atoms of both the systems. We have further calculated the electrical conductivity (σ) of the systems from the following Equation (4)

$$\sigma = \sigma_0 e^{-\frac{\Delta E}{2kT}}.$$
(4)

Here, σ_0 , *k* and *T* represent the proportionality constant, Boltzmann constant and temperature, respectively. These values of electrical conductivity essentially depend on the energy difference of the discrete molecular levels near E_F . It is really important to obtain the information about electron scattering through the medium. We have observed that the room temperature conductivity of the s-SW structure is 0.37 times that of the pristine structure, whereas, for the 4.17% defect concentration, the structures 'ac' and 'dd'' exhibit similar conductivity as s-SW because of the similar values of the HOMO-LUMO gaps, whereas the conductivity of 'ab' is 0.11 times that of the pristine system or 0.3 times that of the s-SW system. Careful observation, however, indicates that the pristine system possesses the highest conductivity and the system 'ab' shows the least conductivity. The reason for this low conductivity of 'ab' is as follows. The electrons encounter more scattering due to the structural defomation of this structure.



Figure 6. Density of states spectra of (**a**) s-SW and pristine GQD' (**b**) ab; (**c**) ac; (**d**) ad; (**e**) ae; and (**f**) dd' structures. The E_F of all the structures are set to zero for convenience. The full width at half maximum of the Lorentzian fitted DOS envelope is 0.1 eV.



Figure 7. (a) HOMO of s-SW; (b) LUMO of s-SW; (c) HOMO of dd'; (d) LUMO of dd'. The image has been drawn with isovalue $0.01 \text{ e}^-/\text{au}^3$.

To explore the possibility of thermionic emission, we have now explored the variations of work function (W) with concentration and distance between the defects. For clear understanding, these values are plotted with the conductivity data in Figure 8. The values of W are relatively high for all the structures. The value of W generally decreases in the presence of SW defect. The only exception is the 'dd'' where the value of W is greater than the pristine structure. These results will be important related to the study of possible applications of these systems in electrical devices.



Figure 8. Conductivity and work function for structures with different defects.

7. Conclusions

A systematic investigation of geometric structure, Raman spectra, IR spectra and electronic properties of the graphene quantum dot with different SW defects have been presented in this paper

under the framework of DFT. The stress has been developed due to the presence of 2.08% SW defect in the graphene network. As a consequence, the G mode $(1562.22 \text{ cm}^{-1})$ has been found to be red shifted to 1544.21 cm⁻¹ after incorporation of a defect. The specific nature of the atomic vibrations responsible for this new G mode vibration has been identified in this paper. In addition to that, low frequency out of phase vibration of the whole system is also found to be softened from 228.79 cm^{-1} to 224 cm^{-1} after incorporation of the defect. Another high frequency peak at 1674.04 cm⁻¹ has been recognized as a characterizing mode caused by the solo contribution of the defected bond. The stability of the systems decreases with increasing SW defect concentrations. In addition, the relative stabilities of the systems for a particular SW defect concentration (4.17%) also depend upon the separation between the defects. We have thus explored the most stable defect positions ('ab' and 'ac') for 4.17% doping concentration. The DOS spectra reveal the absence of DOS at E_F for all these defected structures similar to the pristine system. The electrical conductivity has been observed to be reduced in the presence of SW defects. In addition, the value of work function generally decreases in the presence of SW defects. However, 'dd' possesses a greater value of it as an exception. As a special note, the Raman spectra of the system with two SW defects separated at a distance 7.98 Å possess a mode at 1615.45 cm⁻¹ that can be identified as the signature benchmark related to the collective effect of two individual SW defects only. This result has been well supported by the existence of the same mode in IR spectra at 1619.29 cm⁻¹. Therefore, this distance can be thought of as a minimum distance (separation) for which two SW defects can exhibit independent behaviour. Our study will be very effective to characterize the different types of SW defects into the experimental systems.

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