Efficient hydrogen storage in defective graphene and its mechanical stability: A combined density functional theory and molecular dynamics simulation study

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HIGHLIGHTS

- A combined DFT/MD approach for designing graphene for H₂ storage is presented.
- H₂ desorption from the hydrogenated defective graphene V₂₂₂ is exothermic.
- H₂ adsorption/desorption is more reversible in V₂₂₂ than in pristine graphene.
- Tensile strength of V₂₂₂ shows slight reduction with respect to pristine graphene.

ABSTRACT

A combined density functional theory and molecular dynamics approach is employed to study modifications of graphene at atomistic level for better H₂ storage. The study reveals H₂ desorption from hydrogenated defective graphene structure, V₂₂₂, to be exothermic. H₂ adsorption and desorption processes are found to be more reversible for V₂₂₂ as compared to pristine graphene. Our study shows that V₂₂₂ undergoes brittle fracture under tensile loading similar to the case of pristine graphene. The tensile strength of V₂₂₂ shows slight reduction with respect to their pristine counterpart, which is attributed to the transition of sp² to sp³-like hybridization. The study also shows that the V₂₂₂ structure is mechanically more stable than the defective graphene structure without chemically adsorbed hydrogen atoms. The current fundamental study, thus, reveals the efficient recovery mechanism of adsorbed hydrogen from V₂₂₂ and paves the way for the engineering of structural defects in graphene for H₂ storage.

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Introduction

The demand for energy globally has significantly escalated in the past years. Energy consumption has increased in all major sectors including industry [1], transportation [2] and household [3]. Presently, most of this energy demand is met by burning fossil fuels, and only a small fraction from renewable energy sources [4]. Attempts to limit the fossil fuel consumption [5] to prevent the climate change caused by the emissions of green-house gases have stimulated the research for alternative, cleaner, sources of energy. In this context, several sources of clean energy e.g., biofuels [6], solar [7], hydrogen gas [8,9], have been proposed. Hydrogen (H$_2$) is one of the most promising candidates for transmitting energy that can be used as a power source. However, there are some inefficiencies when designing cryo-adsorption-based H$_2$ storage systems. For instance, the fast refueling is hindered due to the longer cooling down times required to reach operating temperatures. One of the feasible ways to enhance the H$_2$-graphene interaction is by creating a single vacancy on the graphene surface [44–47]. A plausible way to create atomic vacancies is by irradiating the graphene surface with Ar$^+$ ions with an ion energy of 100 eV and an exposure time of 3–4 s [48,49]. Various types of hydrogenated atomic vacancies can be created by the hydrogenation treatment of defective graphene [48].

Hydrogen uptake in graphene via chemisorption is observed to be up to 8.3 wt% [50], which corresponds to the formation of a completely saturated graphene sheet, i.e. the graphene. The stability of graphene was first hypothesized in a DFT study [51], and subsequently experimentally validated by Elias and co-workers [52]. In the case of graphene sheets chemically modified on both sides, the hydrogen uptake can be as high as 12.8 wt% [53] since both sides can be used for hydrogen adsorption. High-capacity hydrogen storage can also be ensured by metallized graphene [53]. Several studies have shown that the introduction of defects on the graphene sheet reduces the modulus of elasticity, enhances the chemical reactivity, and improves the adsorption tendency [54–56]. DFT calculations with a van der Waals correction showed a gravimetric capacity of 5.81 wt% for graphene with high-defect density Stone–Wales and vacancies [57]. Denis et al. [58] showed that adsorption of H$_2$ at the dangling carbon of a single vacancy resulted in improved binding energies due to concentrated negative electronic charge leading to high hydrogen uptake. The first principles study by Kim et al. [59] showed that the presence of vacancy defects enhances the metal binding on graphene and increases the hydrogen uptake capacity. The defective graphene is found to be a potential candidate for vehicular applications since 6 wt% H$_2$ uptake is required for automobiles according to the U.S. Department of Energy (DOE) guidelines [60,61]. A comparison of H$_2$ uptake by graphene-based systems with other H$_2$ storage materials (e.g., MOFs [62]) shows H$_2$ uptake to be similar in both classes of materials. For H$_2$ storage materials, equally important to the maximum H$_2$ uptake is the H$_2$ uptake rate. For more details on both these aspects the reader is referred elsewhere [22,61,63].

The underlying mechanism of chemisorption and desorption of hydrogen in H$_2$ storage materials such as...
graphene remains elusive because of the formidable challenge posed by the experimental imaging of H2 at nano-scale. Atomistic simulations, contrary to experiments, can provide the necessary insight into the hydrogen interaction with the graphene structure at the fundamental level. For instance, the reaction pathways associated with the H2-graphene interaction can be obtained within the Density Functional Theory (DFT) framework [37,42,43]. DFT based studies can provide the necessary understanding of the underlying adsorption and desorption processes of H2, thereby complementing experiments towards creating structure-property relationships which are needed to design H2 storage processes. To this purpose, Sunnardianto et al. [64] performed DFT simulations to investigate the reaction pathways associated with the interaction of molecular hydrogen with hydrogenated single vacancy (i.e. V11). This study revealed an energy barrier of 0.5 eV for the dissociative chemisorption of H2 on V11 in order to reach to the stable hydrogenated vacancy state V211. The energy barrier associated with the desorption of H2 from the V211 state (i.e. V211 + H2) was found to be as high as 3 eV. Thus, the H2 adsorption is found to be energetically feasible on the hydrogenated defective graphene structures proposed so far, however, the energy barrier associated with the desorption is predicted to be high [64,65] thereby hindering easy recovery of adsorbed H2. The irreversible adsorption-desorption process, therefore, demands search for specific hydrogenated defective graphene structures, which exhibit a reduced desorption barrier. The success with identifying such kinds of structures is crucial for improving the applicability of graphene as a H2 storage medium.

Along with the understanding of the H2 adsorption-desorption processes, it is important to get atomistic insights into the mechanical behavior of the hydrogenated defective graphene structure. Earlier works have demonstrated significant impact of the presence of defects such as vacancies on the mechanical properties of the graphene [66–71]. For instance, low concentrations of defects (e.g. single or double vacancies) in graphene have been found to be nucleating points for subsequent fracture in the material [67,68]. Studies have been carried out to explore mechanical properties of both pristine and defect-containing graphene based on first principles, molecular dynamics (MD), and continuum-mechanics simulations. Sun et al. [69] performed MD simulations to probe the effect of a single vacancy on the mechanical behavior of graphene. The study revealed that the existence of a defect significantly reduces the strength of graphene, which is consistent with the findings of Wang et al. [70]. Based on a coupled approach of quantum with MD simulations, Khare et al. [71] investigated the fracture behavior in defective graphene and showed that the fracture stresses can be described by the Griffith formula for defect sizes smaller than 1 nm. A wide range of vacancy coverage in graphene has also been investigated in the past [66], which showed that the ultimate strength gradually degrades with the increase in the vacancy coverage. It is observed that the brittle fracture for low vacancy coverage transforms to ductile fracture for high vacancy coverage.

In the present work, we carried out a comprehensive study to identify a specific hydrogenated defective graphene structure, which has the required characteristics i.e. efficient recovery of adsorbed H2, reversible adsorption-desorption behavior and mechanical stability comparable to that of pristine graphene. Our study is based on a multi-scale approach where the quantum mechanical DFT calculations, coupled with MD simulations, revealed that the two-dimensional graphene structure with hydrogenated vacancy, V222, satisfies the above requirements. The reaction pathway of V211 + H2 ⇔ V222 obtained within the DFT framework revealed that the desorption process is exothermic, which is desirable for H2 storage. To the best of our knowledge, the mechanical properties of hydrogenated defective graphene structure within MD simulations framework have not been yet reported in the literature. MD simulations performed within this work on V222 yielded the tensile strength and fracture strain, which are found to be comparable to those of the pristine graphene. Our study also revealed V222 structure to be mechanically more stable than the defective graphene structure without chemically adsorbed hydrogen atoms. It is important to note that the computation of maximum H2 uptake and uptake rate [61] of the graphene system under investigation is beyond the scope of the present work. Our focus to obtain fundamental insights into H2 adsorption and desorption processes on defective graphene and not on developing a fully optimized graphene-based H2 storage system. For this, more research at multiple length scales is required. Nevertheless, we believe that our study will motivate further research in the area of H2 storage using modified graphene structures.

**Computational methods**

**DFT calculations**

We performed calculations based on quantum mechanical DFT approach described in Ref. [72,73] and implemented in the Quantum ESPRESSO (QE), an open source package available under a GNU license code [74]. The total energies and forces were calculated using the projector augmented wave method [75] and an ultra-soft pseudo potential [76] together with the local density approximation (LDA) for the exchange-correlation potential parameterized by Perdew-Zunger [77]. The energy cut off of 40 Ry was chosen for the plane wave expansion of the wave function and 400 Ry for the expansion of the augmented charge. A Monkhorst-Pack scheme was used with distribution of k-points on a 12 × 12 × 1 mesh [78] for a rectangular supercell of graphene with a single vacancy comprising 63 atoms. A spacing of around 10 Å was used in the direction normal to the graphene layers in order to avoid interlayer long-range van der Waals interactions. The two-dimensional graphene structure with hydrogenated vacancy (abbreviated as V222) used for the present DFT study is shown in Fig. 1. The three hydrogen orientations i.e. Up (U), Slightly Down (SD) and Down (D), are defined based on the hydrogen position relative to the graphene plane. Single hydrogen atoms form out-of-plane bonds such that their position dynamically switches between above and below the graphene plane during the reaction with a vacancy. When a second hydrogen atom is adsorbed at the mono-hydrogenated
vacancy site, steric hindrance does not allow it to come close to the first one, causing the “SD” and “D” orientations. This can be clearly seen in Fig. 1 (b). Structural relaxations were performed until the forces on each atom were below $10^{-4}$ Ry/a.u. The nudged-elastic-band (NEB) method [79,80] was employed to calculate the minimum energy pathway and the energy barriers corresponding to the adsorption-desorption processes. This method takes into account the effect of the local chemical environment on the energy barriers with the adsorption and desorption of the H$_2$ molecules.

In order to check the accuracy of the parameters chosen within our DFT study, we performed structural optimization of graphene with a single hydrogen adsorbed on its surface. Our study yielded C–H bond length to be approximately equal to 1.12 Å, which is in good agreement with the existing DFT-based works [81] and also with the experimentally observed C–H bond length for a free C–H molecule [82]. We also performed some benchmark tests on the buckling which is the distance by which the hydrogenated carbon atom at the vacancy site is shifted in the z direction. We found the buckling of the C atom for the monomer-hydrogenation case to be 0.19 Å, which is in reasonable agreement with previously calculated values reported in the literature, i.e. 0.20 Å [83] and 0.18 Å [84]. All atomic visualizations within our study were created using XCrySDen (Version: 1.5.60) [85].

**MD simulations**

The mechanical stability of the proposed hydrogenated defective graphene structure, V$_{222}$, was studied using MD simulations. The graphene monolayer sheet in our simulations consisted of 2303 carbon atoms with a single carbon vacancy having 6 hydrogen atoms adsorbed at the vacancy edges. Such a system size ensures that possible finite-size effects due to periodic boundary conditions are avoided [66,86]. The choice of this system-size in our MD simulations results in a different defect density compared to the system considered in the DFT calculations. Nevertheless, the purpose of the MD study is to show how the tensile strength of the graphene sheet is locally affected by the hydrogen-defect interaction. The length and width of the graphene sheet were chosen to be 105 Å and 62 Å respectively. The x-axis and y-axis were set along the armchair and zigzag directions, respectively, with the z-axis being normal to the graphene sheet. Periodic boundary conditions were applied along the x- and y-directions of the graphene sheet. As with the DFT simulations, a spacing of ca. 10 Å was applied in the z-direction to avoid interlayer long-range van der Waals interactions. The MD simulations were performed using the large-scale atomic/molecular massively parallel simulator (LAMMPS) software package [87]. The Adaptive Intermolecular Reactive Empirical Bond Order (AIREBO) potential [88] was used which is a reliable interatomic potential for studying covalent bond formation, bond breaking and failure of functionalized graphene [89,90].

Prior to the uniaxial tensile loading, the hydrogenated defective graphene structure was energetically minimized using the conjugate gradient method. The system was then equilibrated for 65 ps in the isothermal-isobaric ensemble (NpT) at 300 K and zero bar using Nose-Hoover thermostat and barostat. A time step of 1 fs was used for the integration of equations of motion. After the equilibration period, a uniaxial tension was applied along the x-direction to both the pristine and hydrogenated defective graphene keeping the temperature equal to 300 K. The strain rate of loading was set as 0.001 ps$^{-1}$. This stepwise straining method has been previously used in MD simulations for studying the mechanical properties of various materials [91,92]. To ensure good statistics, 10 independent runs for the pristine graphene and the hydrogenated defective graphene structure were performed. The Open Visualization Tool (OVITO) [93] was used to visualize the MD data and to generate the MD snapshots.

**Results and discussion**

**Reaction pathway of H$_2$ desorption from the V$_{222}$ surface**

In this study, we chose to study a single vacancy graphene since this system has been experimentally synthesized [94]. Higher defect concentrations can also be experimentally obtained by tuning the ion energy and exposure time in a
controlled manner [95,96]. The ion energy and exposure time are controlled in the experiments in order to avoid high defect concentrations, which can deteriorate the performance of graphene-based devices. This is because high defect concentrations result in the loss of the surface area of graphene, which may result in a reduced hydrogen storage capability [33]. Furthermore, the reconstruction around defects such as a double vacancy makes the graphene structure non-reactive because of the complete saturation of dangling bonds. These kinds of defects, unlike the mono-vacancy systems, are therefore not appropriate for hydrogen storage. Here, we focus on the reaction of H2 molecule at the V211 structure since such a hydrogenated defective structure is experimentally observed for a wide range of hydrogen gas pressures [48]. For instance, V211 is experimentally observed to exist in the chemical potential range –3.66 to –3.20 eV with the total hydrogen pressure in the ultrahigh-vacuum (UHV) chamber adjusted to ~10^{-2} Pa [48]. The surrounding environment is not considered within our study since H2 molecules in the surrounding environment have no effect on the desorption process of V222 → V211+H2. This is primarily because the dangling bonds around the vacancy are completely saturated with hydrogen atoms for the V222 structure as can be seen in Fig. 1.

Using DFT, we investigated the minimum energy pathway of V222 ⇔ V211+H2, which yielded the adsorption and desorption of H2 from the hydrogenated defective graphene surface. As shown in Fig. 2, in the initial state (IS) the two adsorbed hydrogen atoms in the up orientation (indicated by “U” in Fig. 1) at the hydrogenated carbon site of V222 start to desorb, causing the energy of the system to rise. The reaction proceeds via the transition state (TS), during which the two desorbed hydrogen atoms start forming a hydrogen molecule, and ends up at the final state (FS) state, comprising a hydrogenated graphene vacancy V211 with H2 in the vacuum. Our calculations yield an energy difference of approx. 0.63 eV between IS (V222) and FS (V211+H2). Importantly, the FS is found to be at a lower energy than the IS, indicating that the process of H2 desorption is exothermic. This finding is reported for the first time since the hydrogenated defective graphene structures proposed earlier [37,64] lacked this exothermic nature of H2 desorption. In order to substantiate this fact, in Fig. 2 the reaction pathway corresponding to V221 ⇔ V111+H2 computed in a previous work by us [37] is presented. It can be clearly seen that the desorption process for V221 is not exothermic, with a much higher energy barrier for desorption (i.e. 1.32 eV) compared to the desorption barrier of the newly proposed hydrogenated defective graphene structure V222 (i.e. 0.95 eV). This desorption barrier is also much smaller than the corresponding desorption barrier for pristine graphene which is as high as ca. 3 eV [97]. The fact that the desorption process is exothermic makes the corresponding reaction pathway extremely relevant to H2 storage since there is no need of an external catalyst to recover adsorbed hydrogen from the graphene surface. Thus, it is a possible solution to the current problems associated with the difficulty of extracting adsorbed hydrogen from the surfaces of H2 storage materials [98,99].

It is noteworthy that the H2 adsorption and desorption associated with the newly proposed reaction pathway V222 ⇔ V211+H2 is more reversible compared to the pristine graphene. While the energy difference between the adsorption and desorption processes for this pathway is 0.63 eV, the corresponding difference for the pristine graphene is ca. 2 eV [97]. The reaction pathway V221 ⇔ V111+H2 previously reported by us (see Refs. [37]) does not show an exothermic desorption behavior and the nature of the reversibility is not significantly better than the reaction pathway proposed in this study. Thus, the combined characteristics of exothermic H2 desorption and fairly reversible H2 adsorption-desorption, make the graphene with V222 defect more efficient for H2 storage than the previously proposed hydrogenated defective graphene structures.

**Mechanical properties of hydrogenated defective graphene V222**

To study the influence of hydrogenation on the mechanical properties of defective graphene, both the pristine and hydrogenated defective graphene are subjected to uniaxial tension along the armchair direction. The corresponding tensile stress–strain curves for the hydrogenated defective graphene structure (V222) and the pristine graphene are shown in Fig. 3 for a temperature of 300 K. Both systems are found to undergo brittle fracture under tensile loading. A linear increase in the stress is observed with increasing strain (ε) at the initial deformation stage i.e. until the strain value of ca. 0.06. With the progressive increase in the strain values, the stress curves for both the cases show a non-linear behavior until the elastic limit of the strain is reached (i.e. the fracture strain), at which the stress undergoes a sharp drop due to the initiation of fracture in the graphene sheet. Fig. 3 clearly shows that the presence of 6 adsorbed hydrogen atoms on a single vacancy

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**Fig. 2 – (Color online) The red curve is the minimum energy pathway from initial state (IS) V222 to final state (FS) V211+H2 via the transition state (TS). The black curve is minimum energy pathway from initial state (IS) V221 to the final state (FS) V111+H2 which is reproduced from Ref. [37] for comparison. The activation barriers are indicated in the figure for both the cases. The blue and red atoms represent the carbon and hydrogen atoms, respectively. The lines connecting the NEB images are guide to the eye. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)**
has an insignificant effect on the stress-strain curve of graphene until a strain value of ca. 0.14. This is evident from the overlapping stress-strain curves for both the systems until \( \varepsilon = 0.14 \). The maximum value of the stress, i.e. the tensile strength, is computed to be 100 GPa for the hydrogenated defective graphene. The corresponding value for the pristine graphene is found to be 106 GPa, which is in good agreement with the result obtained in other simulation works [70,100], thereby validating our modelling approach. The computed fracture strain values for the hydrogenated defective graphene and the pristine one are 0.169 and 0.173, respectively.

Our results, thus, show that the tensile strength and the fracture strain of the pristine graphene are reduced by 5.7% and 2.3%, respectively, in the presence of a hydrogenated single vacancy. This reduction can be attributed to the conversion of local carbon bonding from sp\(^2\) to sp\(^3\)-like hybridization for the di-hydrogenated carbon atoms. Such a transition of local carbon bonding from sp\(^2\) to sp\(^3\) has been reported earlier in MD studies of hydrogenated graphene [100,101]. It is reported in Refs. [70] that the presence of a single vacancy leads to 17% reduction in the tensile strength of graphene. Thus, the qualitative comparison of the tensile strength of the hydrogenated defective graphene at 300 K with the one corresponding to graphene with a single vacancy [70] reveals that the former is mechanically more stable than the latter. Furthermore, the tensile strength of hydrogenated graphene without vacancy is also found to be significantly lower (i.e. ca. 55 GPa) than that of pristine graphene at 300 K [101].

Our results, thus, show that the adsorption of hydrogen atoms in a defective graphene tends to improve the mechanical properties of graphene as compared to graphene with a single vacancy [70] or hydrogenated graphene without vacancy [101].

In earlier MD work [69,100,101], the atomistic origin of the failure of pristine, hydrogenated graphene and vacancy containing graphene under uniaxial tensile load have been systematically investigated. However, atomistic insight into the hydrogenation-induced altering of local binding configurations and its subsequent impact on the failure of defective graphene structure are still lacking. To this end, we investigate the atomic stress distribution in hydrogenated defective graphene (V\(_{222}\)) sheet subjected to uniaxial tensile loading along the armchair direction at 300 K. The snapshots of atomic stress distributions in the hydrogenated defective graphene sheet are illustrated in Fig. 4. This figure illustrates a bond breaking, crack nucleation and growth scenario in hydrogen functionalized defective graphene as observed in our simulations. To get deeper insights into the bonds having maximum stresses and the initiation of bond breaking, the zoomed in images of atoms around the vacancy have been included in Fig. 4(a) and (b). Fig. 4(a) shows the atomic stress distribution on the carbon and hydrogen atoms in the defective graphene sheet before the breaking of bonds. The orientations of the hydrogen atoms are denoted by “U” for up, “D” for down and “SD” for slightly down orientations in Fig. 4(a). The dashed circle in the zoomed in image denotes the position of the single vacancy in the graphene sheet. It is evident that the highest stresses occur on carbon atoms C\(_1\)–C\(_4\) (inside the two black squares) above and below the hydrogen functionalized region, where the C–C bonds are still sp\(^3\)-hybridized. From Fig. 4(a), it can also be seen that the two carbon atoms each bonded with “U” and “D” orientated hydrogen atoms are under intermediate stress. However, the carbon atom bonded with “U” and “SD” orientated hydrogen atoms is found to have minimal stress. Our results indicate that the sp\(^3\)-like hybridization of the two carbon atoms (each bonded with “U” and “D” orientated hydrogen atoms) leads to the highest stress concentration on their two nearest neighbor carbon atoms i.e. C\(_1\) and C\(_3\) located just above and below the vacancy. However, the two sp\(^2\) bonded nearest neighbor carbon atoms of the third sp\(^3\)-like hybridized carbon atom (bonded with “U” and “SD” orientated hydrogen atoms) are under intermediate stress.

Fig. 4(b) illustrates the initiation of the bond breaking in the hydrogenated defective graphene sheet. It can be seen in the inset of the figure that the bond breaking initiates with the debonding of the sp\(^2\)-hybridized carbon atoms, C\(_1\) and C\(_2\), which had highest stress concentration prior to the breaking of the bond. The opening of the crack is indicated by dashed curves in the inset of the figure. Our simulation results, thus, show that the fracture initiates near the hydrogenated vacancy by debonding of carbon atoms with the highest stress. Our results also point to the fact that the sp\(^3\)-like hybridization of the di-hydrogenated carbon atoms results in the reduction of the mechanical strength of hydrogenated defective graphene. The reason is that the sp\(^3\)-like hybridization of the carbon atoms leads to the highest stress concentration on sp\(^2\)-hybridized carbon atoms, which initiates bond breaking leading to the destruction of the six-membered ring configuration near the vacancy.

Fig. 4(c) shows subsequent breaking of bonds outside of the hydrogenated defective region leading to the formation and propagation of a crack along the y-direction i.e. zigzag direction. It can also be observed that there are few monoatomic carbon chains formed to connect locally torn graphene
patches. With further increase in the applied uniaxial strain (Fig. 4(d)), the hydrogenated defective graphene sheet is torn into two patches by the successive debonding of sp²-hybridized C–C bonds along the armchair direction. Few of the monoatomic carbon chains are still found to exist connecting the two patches of the graphene sheet. Our results, thus, show that the crack originated near the hydrogenated vacancy, propagates perpendicular to the direction of the applied tensile load until complete rupture of the graphene sheet.

Conclusions

A comprehensive study of the H₂ adsorption-desorption on a hydrogenated defective graphene structure and its mechanical stability were performed based on a combined DFT-MD approach. The minimum energy pathway corresponding to the H₂ adsorption at V₂₁₁ and desorption from V₂₂₂ was investigated based on DFT. Our study revealed the desorption process to be exothermic with an activation barrier of 0.95 eV. Furthermore, the value of the energy barrier associated with the H₂ adsorption was found to be comparable to that of the desorption process. The existence of the comparable energy barriers suggests that the V₂₂₂ works as a self-catalyst for both adsorption and desorption processes on the graphene surface. In addition, the exothermic nature of H₂ desorption from the V₂₂₂ makes it a better prospect for H₂ storage than the other kinds of hydrogenated defective graphene structures proposed so far in the literature. To obtain atomistic insights into the mechanical stability of the hydrogenated defective graphene structure, we performed MD simulations to compute the tensile strength and the fracture strain. The tensile strength of the proposed hydrogenated defective graphene structure showed slight reduction with respect to its pristine counterpart due to the transition of local carbon bonding from sp² to sp³-like hybridization for the di-hydrogenated carbon atoms. Moreover, our study revealed that the V₂₂₂ structure is mechanically more stable than the graphene with a single vacancy with no chemically adsorbed hydrogen atoms at the vacancy edge. Thus, we made a new finding that V₂₂₂ has special characteristics such as exothermic H₂ desorption, reversible H₂ adsorption-desorption along with mechanical properties comparable to the pristine graphene and better than the 2D graphene sheet with a single vacancy. Based on our analysis, we conclude that it is possible to improve the
suitability of graphene for H₂ storage by appropriate structural modification at the atomistic level.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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