Carbyne Ring Activated Using ZnCl₂ for Hydrogen Adsorption: DFT Study

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ABSTRACT: We have studied the feasibility of activated carbyne as a good hydrogen storage material. Density functional theory (DFT) simulations through van der Waals interactions have been applied to investigate calcium sorption on activating carbyne with zinc dichloride (ZnCl₂) and also interactions of molecular hydrogen with pristine carbyne and Ca functionalized on an activated carbyne C₁₂-ring. The obtained results showed that (i) the chemical activation of the C₁₂-ring with ZnCl₂ increases its area by 5.17% with respect to pristine carbyne. (ii) Ca atoms at small concentrations tend to get atomically sparse on carbyne, donating +0.94e and +1.05e to the ring, according to Mulliken population analysis and the electrostatic potential concentrations tend to get atomically sparse on carbyne, donating +0.94e and +1.05e to the ring, according to Mulliken population analysis and the electrostatic potential charging, respectively. Furthermore, in the presence of calcium, hydrogen sorption increases by 21.8% in comparison with Ca-decorated pure carbyne. (iii) Seven hydrogen molecules per Ca atom have adsorption energy close to the range of ∼0.3−0.5 eV per H₂, which is necessary for effective charge/discharge cycles. (iv) Theoretical uptake (7.11 wt %) with a single Ca atom is higher than the U.S. Department of Energy target (5.5 wt %). Therefore, an activated C₁₂-ring can bind three Ca atoms with its seven H₂ molecules reaching 13.8 wt %. (v) Equilibrium pressure for CaC₁₂−7H₂ and Ca₃C₁₂−21H₂ systems (5−15 MPa) by means of adsorption isotherm calculations. The calculated van’t Hoff desorption temperatures exceed considerably the boiling point of liquid nitrogen. In addition, we also performed DFT-based molecular dynamics simulations for the C₁₂, CaC₁₂, CaC₁₂−7H₂, and Ca₃C₁₂−21H₂ systems to study thermal stability. Our results confirm the potential of Ca-decorated carbyne for hydrogen storage.

INTRODUCTION

The aim here is to determine hydrogen storage properties on a carbyne C₁₂-ring structure through chemical activation with zinc dichloride (ZnCl₂) and decorated with calcium atoms, by density function theory (DFT) calculations. In order to know the increasing demands necessary to hold common living standards while at the same time avoiding resource reduction and environmental pollution, there is a necessity for the growth of high-efficiency, low-cost, and eco-friendly energy storage systems.1 As we all know, hydrogen is an ideal clean energy source that could one day replace fossil fuels, particularly for transportation applications, and combat global warming. One of the principal challenges in the growth of this technology is a compact, safe, and accessible storage system. A desirable system has to be capable of storing hydrogen with upper gravimetric density (HGD) under ambient conditions.2−4 The U.S. Department of Energy (DOE) establishes a goal for ideal hydrogen storage materials: they ought to reach 4.5−5.5 wt % gravimetric density by the 2025 year.5 In general, most of the studies have been dedicated to pristine carbon-based nanomaterials, such as activated carbons,6,7 fullerenes,8−10 graphene,11 and nanotubes,12,13 because most candidates for H₂ storage are explored owing to their low density, high thermal and chemical stability, and plainness of production; nevertheless, it has been established that the hydrogen storage capacity in these systems considerably diminishes at room temperature and ambient pressure,14 being attributed to weak interactions between hydrogen molecules and carbon-based materials, due to physical adsorption (∼0.05 eV).15 Metal doping is one of the effective methods to upgrade the strength of binding between hydrogen molecules and carbon-based nanomaterials. Alkaline-earth metal dopants, especially Ca atoms, show better hydrogen storage performance.16−24 Ca atom is selected as the main dopant because of not only its lower cohesive energy (1.84 eV) compared with the transition metals (∼4 eV) but also its lightness in weight, its lower trend to aggregate on the host material whenever they are deposited, and its ability to retain the hydrogen content after doping.20−22 Carbyne is composed of sp-
hybridized carbon atoms. The material has been proposed as nanoelectronic and also is likely to be used for hydrogen storage owing to an effective surface area around 13,000 m² g⁻¹, four times larger than theoretical graphene values. The carbyne-ring structure is the base state of small carbon clusters (up to about 20 atoms) and an alternative form of linear chains, which are obtained by laser vaporization of graphite. Between carbon clusters that result beginning with this technique in which carbyne has been found, ring structures are comparatively the most stable compared to other configurations and are the principal precursors of fullerenes and nanotubes. Two carbyne structures might be defined as cumulene (equal double bonds) and polyyne (alternative single/triple bonds). Previous research has shown theoretical estimation of Ca-decorated pristine carbyne for hydrogen storage on linear chains and rings, which meet the U.S. Department of Energy target requirement of 5.5 wt %. Nevertheless, a single calcium atom can bind with binding energy higher than the cohesive energy, which makes cluster formation energetically favorable in these systems. To increase hydrogen storage properties on pristine carbyne molecule and enhance the binding energy of subsequent Ca atoms on carbyne, the carbyne ring has been activated through chemical activation with ZnCl₂ to upgrade the superficial area, and enhance pore size distribution (PSD).

Chemical activation with KOH, NaOH, H₃PO₄, or ZnCl₂ is a process where an activating agent is aggregated into the carbon precursor prior to pyrolysis at a temperature normally in the 450—900 °C range. The chemical agents help to develop porosity by means of dehydration (ZnCl₂ or H₃PO₄) and degradation of the biomass structure, especially when the activation agent is highly alkaline (KOH). Among the various activation agents, ZnCl₂ activation reacts with lignocellulosic precursor at T < 500 °C, producing a template effect and inducing a uniform micropore formation. The surface areas are normally between 1500 and 2000 m² g⁻¹ with pore volumes <
1.5 cm$^3$ g$^{-1}$, and the broad pore size distribution increases with the concentration of ZnCl$_2$. The highest hydrogen adsorptions at 77 K and 1 bar reported for any natural and synthetic activated carbon material are 3.28 wt.% from hemp stem, 2.85 wt.% from petroleum pitch, and 2.96 wt.% from NAC-1.5-600 when the activating agent is KOH. For ZnCl$_2$, the storage capacities obtained on CH$_4$ and CO$_2$ are 13 cm$^3$ g$^{-1}$ from oil palm shell and 1.3 mmol g$^{-1}$ from rice husk, respectively. Overall, the specific surface area (SSA) is key to not just enhancing H$_2$ storage but also achieving a higher electrochemical capacitance in terms of power delivery rate and energy storage. So, in this study, we have investigated how functionalization with Ca atoms on activated carbyne with ZnCl$_2$ could influence the hydrogen storage ability. The results show adsorption energy which corresponds to chemisorption between activated carbyne and dopant agents and agreed with gas adsorption on a solid surface; with clarity we found physical adsorption between Ca atoms and hydrogen molecules. In addition, we have studied hydrogen desorption temperatures with respect to the equilibrium pressure by using the van’t Hoff equation, and we determine the equilibrium pressure for the Ca-decorated carbyne by means of adsorption isotherms, to thoroughly evaluate the potential of carbyne as a hydrogen storage material.

**Results**

To carry out the analysis of this work, we take the pristine C$_{12}$-ring previously investigated and then we activate with zinc dichloride (ZnCl$_2$) in order to grow the surface area and enhance pore size distribution, with the aim of improving its hydrogen storage properties. The C$_{12}$-ring corresponds to a C$_{4}$ structure with N = 3; with a D$_{4h}$ symmetry. Figure 1a shows geometry optimization of pristine carbyne C$_{12}$-ring with alternating single/triple bonds, called polyyne, where the most general case is linear acetylene (H$\equiv$C$\equiv$C$\equiv$H). First, to estimate the H$_2$ adsorption capacity on the pristine carbyne without calcium attachment, we use two measures: specific surface area (SSA) and accessible surface area (ASA). The SSA method is based on the geometrical calculation of the area, whereas the ASA method is based on the Monte Carlo integration technique where the probe molecule is “rolled” over the framework surface. The SSA calculations of pristine and activated carbyne C$_{12}$-ring have been carried out by inserting triangles on the ring using the Heron formula, eq 1.45

$$A = \sqrt{P(P-a)(P-b)(P-c)}$$

where $P$ is the perimeter of a triangle with $a$, $b$, and $c$ sides, whereas the PSD is calculated as an approximation to the circle area.

Using eq 1, we calculated the area and pore diameter showing values of 18.37 Å$^2$ and 4.83 Å, respectively. With these results, we calculated the SSA of pristine carbyne, showing a value of 4606.31 m$^2$ g$^{-1}$, whereas ASA shows a value of 13000 m$^2$ g$^{-1}$, according to Biovia Materials Studio Software and the Monte Carlo integration technique. Second, the average hydrogen binding energy on pristine and activated carbyne was calculated from the following, eq 2.46

$$E_b = (E_{C_{12}+nH_2} - E_{C_{12}} - N_H E_{H_2})/N_H$$

where $N_H$ is the number of hydrogen atoms physisorbed on the inner and outer surfaces of pristine and activated carbyne C$_{12}$-ring, $E_{C_{12}+nH_2}$ and $E_{C_{12}}$ are the total energies of the hydrogenated carbyne ring and corresponding pristine and activated C$_{12}$-ring, respectively, and $E_{H_2}$ is the energy of an isolated hydrogen atom. As in the cases of graphene, CNTs, and C$_{60}$, hydrogen adsorption on pristine C$_{12}$-ring is due to weak van der Waals (vdW) interactions. The adsorption energy of a single H$_2$ molecule to the outer and inner surfaces of pristine carbyne only reaches adsorption energies of 0.077 and 0.028 eV, respectively. The next hydrogen molecules added to the host material on the inner and outer surfaces diminish the binding energy until 0.01 eV; this means that the pristine carbyne is not a good candidate for hydrogen storage directly. The interactions are evident through the potential energy surfaces (PESs), which give the minimum $E(r)$ and correlate with the equilibrium point later to geometry optimization of the interacting system (C$_{12}$ + H$_2$) on the outer and inner surface of pristine carbyne. Figure 1b shows the bond strengths of C$_{12}$ + H$_2$, where the H$_2$ molecule is adsorbed at the outer and at the inner surfaces. The resulting values show the dissociation energy to form C$_{12}$ + H$_2$ corresponding to physisorption, which involves binding hydrogen molecules to the host material and requires very low temperatures and high hydrogen gas pressure.

Subsequently, through chemical activation with zinc dichloride (ZnCl$_2$), we activate the pristine carbyne to increase the surface area and enhance PSD. In widespread terms, these characteristics are central to not only enhancing hydrogen storage but also achieving the best work in power distribution rate and energy storage. We place a ZnCl$_2$ molecule to the center of the pristine C$_{12}$-ring at 2.529 and 2.410 Å distances as shown in Figure 1c. By applying geometry optimization and removing ZnCl$_2$, the area and pore size diameter result in 19.32 Å$^2$ and 4.96 Å, respectively, using eq 1, representing increases of 5.17% and 2.69% with respect to pristine C$_{12}$-ring (Figure 1d). In addition, we calculate the SSA and ASA of activated carbyne C$_{12}$-ring as a function of their geometrical characteristics. The obtained values show 4844.53 m$^2$ g$^{-1}$ (SSA) and 15047.88 m$^2$ g$^{-1}$ (ASA), which represents a macroscopic parameter that might be the kind to modify the synthesis condition of carbyne molecules. Even more, from literature, activated carbon shows a surface area of fewer than 2000 m$^2$ g$^{-1}$ and cannot be used for hydrogen storage. Activated carbyne ring represents an alternative due to high specific surface area as a hydrogen storage material. Then, we investigate the hydrogen adsorption of a single H$_2$ molecule as much at the outer as the inner surface of activated C$_{12}$-ring (Figure 1e). The results show that only in the center of the C$_{12}$-ring the H$_2$ molecules might be adsorbed; however, only two H$_2$ molecules may be adsorbed with an adsorption energy higher than -0.1 eV. The outer zones of the activated C$_{12}$-ring only reach an adsorption energy of -0.030 eV (Figure 1f), which corresponds to lower adsorption energies for hydrogen storage at environmental conditions. These results are compared against toroidal carbon nanostructure C$_{120}$ where hydrogen adsorption energies are lower by 0.1 eV per H$_2$; a full hydrogen storage uptake of 2.05 wt % is for 15 H$_2$ molecules adsorbed at the inner surface of the toroidal carbon C$_{120}$. The same case is observed for the activated C$_{12}$-ring, where the full hydrogen storage capacity is 2.72 wt % with only two H$_2$ molecules adsorbed at the inner surface, which do not meet the goals established by the DOE. Although there is an upgrade in hydrogen adsorption on activated carbyne, this makes it impractical for mobile applications, just like a pristine carbyne ring. However, this is a good start for future research to address this.
Subsequently, we studied the behavior of Ca-doped activated carbyne with ZnCl₂. The modeling of the Ca−carbyne complex considers several starting configurations at the inner and outer surfaces (in front of single and triple bonds and at the center of the activated carbyne ring) separated at 2.48 Å, and we examine the Ca−carbyne stability by determining the binding energy ($E_{b,\text{Ca}}$) using:

$$E_{b,\text{Ca}} = E^{\text{Ca}}_{\text{C12}} - E^{\text{Ca}}_{\text{C12}} - x E_{\text{C12}} - E_{\text{C12}}$$

where $E_{\text{C12}}$ is the total energy of the activated carbyne molecule, $E_{\text{Ca}}$ is the total energy per calcium atom, and $E_{\text{C12},\text{C12}}$ is the total energy of one carbyne molecule with $x$ Ca atoms. For a single Ca atom, we find three optimal positions around the C12-ring to be located in front of either single or triple bonds and also at the center of the ring, with binding energies of 2.79 eV (single and triple bonds) and 3.34 eV (center of the ring), which with respect to the pristine CaC12 complex represents increases of 25.11% and 49.77%, respectively, which means it is strongly chemisorbed.

Table 1. Average Energy, Adsorption Energy of the $n$th H₂ Molecule on the Doped Complex (eV) and Binding Energy of CaC₁₂ (eV)

| position | functional | energy (eV) | $E_{b,\text{Ca}}$ (eV) |
|----------|------------|-------------|------------------------|
| triple   | DFT-D      | 2.79        | 0.5362 0.4346 0.4143 0.4132 0.4103 0.4018 0.3938 |
| single   | DFT-D      | 2.79        | 0.4515 0.4278 0.4124 0.4062 0.4019 0.4034 0.3983 |
| center   | DFT-D      | 3.34        | 0.3818 0.3454 0.3502 0.3176 0.3293 0.3929 0.3706 |

As a note, unlike the pristine CaC₁₂ complex, only the single bond C₁−C₂, C₅−C₆, or C₉−C₁₀ was the better zone for calcium atoms; these zones present the blue lobes for HOMO−LUMO spatial distribution. For activated C₁₂-ring we found a better HOMO−LUMO distribution (Figure 2a) around the ring, where the Ca atom prefers to bind indistinctly in any zone of carbyne C₁₂-ring. Here, the positive value of $E_{b,\text{Ca}}$ means that the doped complex is thermodynamically stable. In addition, our results of binding energy values were higher than those of fullerenes, carbon nanotubes, carbyne chains, graphene, and heterofullerenes. Therefore, this indicates that cluster formation is energetically unfavorable, so this factor will not diminish the possible hydrogen capacity on activated C₁₂-ring. Panels b−d of Figure 2 show the geometry optimization of the activated doped complex. The calcium atom tends to extend the double and single bonds to the activated carbyne to 1.404 and 1.405 Å, respectively, and how the Ca atom is adsorbed at the center of the ring.

In addition, we have also achieved DFT-based molecular dynamics (MD) simulations for both C₁₂ and CaC₁₂ systems in their forms pristine and activated with ZnCl₂. All of the...
Figure 3. continued
simulations were achieved in an NVT ensemble (constant number of atoms, volume, and temperature) with a specific temperature of 300 K. The molecular dynamic simulations were run for 6 ps, with 1 fs as a time of step, using massive GGM and Nosé–Hoover thermostat to address the structural and thermal stabilities of the system (Figure 3a–h). Our results after

Figure 3. Molecular dynamics (MD) production run of the C12-ring and calcium-decorated carbyne after 6 ps with massive GGM and Nosé–Hoover thermostat. (a, b) Pristine carbyne C12-ring. (c, d) Activated carbyne C12-ring with ZnCl2. (e, f) Calcium-decorated carbyne CaC12 (pristine). (g, h) Calcium-decorated carbyne CaC12 (activated with ZnCl2).
6 ps of MD production show better thermal stabilities of all activated structures with Nose–Hoover thermostat at 300 K. Every system was equilibrated for 3–6 ps, and after 5 ps of production, no breaking of bonds was observed, which implied the thermal stability of C12 and CaC12 systems. The average temperature of the MD production run for pristine carbyne C12-ring is 327.11 K and 299.11 K for massive GGM and Nose–Hoover thermostat, respectively, along the 6 ps MD production run. For activated C12-ring the average temperature was 327.46 K and 300.04 K for massive GGM and Nose–Hoover thermostat, respectively. The mean square displacement (MSD) is shown in Figure 3a–h for all of the systems (C12-ring and Ca-decorated carbyne). To summarize, the run of MD production showed that activated CaC12 is a good candidate since this structure after 6 ps retains its initial properties without too much variation in bond lengths and considerably lower temperatures, which implied good thermal stability of activated carbyne. In addition, using the Nose–Hoover thermostat we have carried out an MD production run at 200 and 100 K for the carbyne C12-ring and doped complex CaC12 (pure and activated) that is shown in the Supporting Information.

Through Mulliken population analysis and electrostatic potential (ESP)-fitted charges, we observe positive charge on the Ca atom toward the carbyne ring, which results in (+0.949e and +1.464e) for a single bond, triple bond, and center of carbyne molecule, respectively. As shown in the Supporting Information, we added the charge-transfer mechanism of carbyne C12-ring and doped complex CaC12−nH2 with n = 1–7 H2 molecules adsorbed around the Ca atom for pristine and activated carbyne with ZnCl2.

Once the CaC12 activated complex reaches equilibrium, the next step is the adsorption analysis of H2 molecules on the decorated complex. Using eqs 4 and 2, we calculate the average binding energy and adsorption energy of nH2 molecules adsorbed on the doped complex.

\[
E_{av} = \frac{[E_{(Ca_{12})} + nE_{(H_2)} - E_{(Ca_{12}−nH_2)}]}{n} \tag{4}
\]

\[
E_{ad} = E_{(Ca_{12}+(n-1)H_2)} + E_{(H_2)} - E_{(Ca_{12}−nH_2)} \tag{5}
\]

where \(E_{(Ca_{12})}\) and \(E_{(H_2)}\) are the total energies of the CaC12 complex and an isolated H2 molecule, respectively. The \(E_{(Ca_{12}−nH_2)}\) is the total energy of the CaC12 system with nH2 molecules and \(E_{(Ca_{12}+(n−1)H_2)}\) is the total energy of the CaC12 system with \((n−1)\) H2 molecules adsorbed on the doped complex. The next H2 molecules were placed one by one around the Ca atom until there were seven H2 molecules. The average energy and adsorption energy results of the nth H2 molecule

Figure 4. Geometry optimization scheme for activated CaC12−nH2 with n = 1–7 H2 molecules adsorbed onto the doped complex. (a–d) Hydrogen adsorption on DFT-GGA-PBE with the empirical correction scheme of Grimme (DFT-D), where the first six H2 molecules are adsorbed around the Ca atom and the seventh H2 molecule is on top of the Ca atom. The configuration of H2 molecules in the doped complex is observed as gray color for carbon atoms, white color comprises H2 molecules, and green color corresponds to the decoration Ca atoms.
Adsorption of H2 by CaC12 complex using eqs 4 and 5 are summarized in Table 1. In Figure 4, we legibly illustrated the process of attaching molecules to the CaC12, performed by setting one by one until a maximum of seven H2 molecules.

We determine that the first six H2 molecules tend to be adsorbed around the calcium atom, and the seventh H2 molecule is physically adsorbed on the upper side of the calcium atom. The optimized systems of CaC12–7H2 are rather similar for single and triple bonds; therefore, we only take the triple bonds and center to properly put each one of the seven H2 molecules. Therefore, in this study with correction of Grimme calculations, we determine that the CaC12 complex might adsorb until seven H2 molecules per Ca atom with enough energy and quantity of hydrogen for using it as a storage material. The weight percent (wt %) of H2 molecules in a carbyne CaC12 are also calculated with eq 6.

\[
\text{wt} \% \text{H}_2 = \left[ \frac{m_{\text{H}_2}}{m_{\text{H}_2} + m_{\text{CaC}_{12}}} \right] \times 100
\]

where \(m_{\text{H}_2}\) is the mass of H2 molecules adsorbed on the decorated complex, and the mass of CaC12 decorated complex is \(m_{\text{CaC}_{12}}\). The hydrogen storage capacity obtained in this study is 7.11 wt % (Table 1). Thus, the maximum hydrogen storage capacity is greater than the capacity ~6 wt % of Ca-decorated boron hetero-fullerenes, ~5 wt % Ca-decorated carbon nanotubes, by theoretical DFT calculations, and other activated carbons by using several activating agents such as KOH, ZnCl2, H2SO4, and H3PO4 (see Table 2).
Subsequently, we saturate the doped complex by placing up to three Ca atoms on the mentioned positions (single and triple bonds and the center of the ring) and we observe binding energies above 2.8 eV for the second and third Ca atoms, which show stability in the system, indicating that the cluster formation is energetically unfavorable as an increase in the calcium atoms concentration in the system. We repeat the same methodology of adding H₂ molecules to the Ca atoms, even to which 7H₂ molecules per Ca atom might bind, with an average binding energy of ~0.36 eV per H₂, obtaining a Ca₃C₁₂–21H₂ system. This study reaches 13.8 wt % for the gravimetric density, fulfilling DOE requirements. However, this gravimetric density requires experimental investigation to be validated and makes it very feasible that the system can only absorb up to one Ca atom.

As the next step, we built potential energy surfaces for doped complex CaC₁₂–nH₂ with n = 1–7 hydrogen molecules adsorbed on it. The methodology to accomplish potential energy curves for the hydrogen molecules adsorbed by Ca atom is to perform a geometry optimization for each H₂ molecule added to the doped complex as a first step, which provides the minimum energy and distance corresponding to the equilibrium point of each system. Then, single-point calculations by oscillating ±4 Å around the minimum energy with steps of 0.02 Å to calculate energies at each point and build potential energy surfaces, E(r). Figure 5 shows PES for the seven H₂ molecules adsorbed on activated doped complex when the Ca atom is placed in front of a single bond of carbyne molecule. All the minimum energies obtained and calculated by eq 4 are equivalents and provide information about the 7H₂ molecules that were physically adsorbed by calcium atoms (see Table 1 and Figure 5).

We carry out HOMO−LUMO calculations for the CaC₁₂–7H₂, which is observed in Table 3. We clearly observed that values of energy difference for the doped complex are about 0.926–0.960 eV for the single bond, 0.925–0.965 eV for the double bond, and 0.13–0.6 eV for the center of the carbyne molecule, indicating that the doped complex is stable enough.

We determine the isothermal curves at three temperatures (274, 298, and 322 K) for the CaC₁₂–7H₂ and Ca₃C₁₂–21H₂ systems using the Sorption program as described in Computational Methods. The equilibrium pressure with the fitting curve when we have 7.11 wt % for the CaC₁₂–7H₂ system and 13.8 wt % for the Ca₃C₁₂–21H₂ system, since they present energies in the desirable range of 0.2–0.6 eV, for hydrogen storage. Therefore, the equilibrium pressure lies in the range of 5–15 MPa as shown in Figure 6.

Together with the gravimetric densities, the thermal stability of adsorbed H₂ on activated complex should be investigated as it plays an indispensable role in predicting the effectiveness of hydrogen charge/discharge cycles. The thermal stability correlates with the binding energy of hydrogen to the storage

![Figure 5. Potential energy surfaces (PESs) corresponding to the CaC₁₂–7H₂ system.](image)

![Figure 6. Isothermal curves at three temperatures (274, 298, and 322 K), when we have 7.11 and 13.8 wt %, which corresponds to CaC₁₂–7H₂ and Ca₃C₁₂–21H₂ systems. The equilibrium pressures from the fitting curve when we have 7.11 and 13.8 wt % are in the range of 5–15 MPa.](image)

| Table 3. HOMO–LUMO Energy Difference or Gap (Δ, eV) of CaC₁₂ and CaC₁₂–7H₂ |
|---|---|---|---|---|---|
| system | HOMO (eV) | LUMO (eV) | Δ | HOMO (eV) | LUMO (eV) | Δ | HOMO (eV) | LUMO (eV) | Δ |
| CaC₁₂ | –4.292 | –3.332 | 0.960 | –4.295 | –3.33 | 0.965 | –5.206 | –4.601 | 0.605 |
| CaC₁₂–H₂ | –4.222 | –3.290 | 0.932 | –4.238 | –3.301 | 0.937 | –5.240 | –4.782 | 0.458 |
| CaC₁₂–2H₂ | –4.137 | –3.211 | 0.926 | –4.196 | –3.271 | 0.925 | –5.240 | –5.103 | 0.137 |
| CaC₁₂–3H₂ | –4.211 | –3.261 | 0.950 | –4.182 | –3.233 | 0.949 | –5.272 | –5.061 | 0.211 |
| CaC₁₂–4H₂ | –4.201 | –3.251 | 0.950 | –4.215 | –3.261 | 0.954 | –5.275 | –5.079 | 0.195 |
| CaC₁₂–5H₂ | –4.207 | –3.273 | 0.934 | –4.224 | –3.283 | 0.941 | –5.246 | –5.050 | 0.196 |
| CaC₁₂–6H₂ | –4.233 | –3.292 | 0.941 | –4.232 | –3.297 | 0.935 | –5.181 | –5.013 | 0.168 |
| CaC₁₂–7H₂ | –4.223 | –3.281 | 0.942 | –4.241 | –3.301 | 0.940 | –5.194 | –5.025 | 0.169 |

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Here, we are using eq 7 (van’t Hoff equation) to estimate the desorption temperature.

\[
T_D = \frac{E_{ad}}{k_B} \left(\frac{\Delta S}{R} - \ln p\right)^{-1}
\]

where \(E_{ad}\) is the hydrogen adsorption energy, \(k_B\) is the Boltzmann constant, \(\Delta S\) is the change in the hydrogen entropy from molecular gas to dissolved solid hydrogen (standard hydrogen entropy, 130 J K\(^{-1}\) mol\(^{-1}\)).\(^{14}\) \(R\) is the gas constant, and \(p\) is the equilibrium pressure (in our calculations we used the range of 0.1–1 MPa with respect to the standard atmospheric pressure).

The activated carbyne complex (shown in Figure 3) is considered as the highest gravimetric density to estimate the hydrogen desorption temperatures. Panels a–c of Figure 7 show van’t Hoff desorption temperatures in the range of equilibrium pressures (0.1–1 MPa) corresponding to hydrogen gravimetric densities for the chosen structure. So, all \(T_D(p)\) dependencies were obtained by employing hydrogen adsorption energies (in the case of Ca\(_{12}\)–7H\(_2\) activated complex) in eq 6.

Average \(T_D\) is calculated by using \(E_{av}\) (average binding energy of seventh H\(_2\) molecule adsorbed per Ca atom). The onset desorption temperature (min \(T_D\)) is obtained to describe adsorption energy (\(E_{ad}\)) of the seventh hydrogen molecule per Ca atom, and it corresponds to the minimal temperature, which is necessary to start hydrogen release. By using the adsorption energy of the first H\(_2\) molecule adsorbed by Ca atom on a doped complex, we calculate the highest desorption temperature, max \(T_D\), which is necessary to fully discharge the considered system. At normal atmospheric pressure (\(p = 0.1\) MPa), the average \(T_D\) is 292 and 295 K (Figure 7a,b) when the Ca atom is placed in front of triple and single bonds and can adsorb until seven H\(_2\) molecules, in the case of the center of the ring the maximum desorption temperature at 0.1 MPa is 275 K (Figure 7c), which is much higher than the critical point of hydrogen (33 K) and more than triple the boiling point to liquid nitrogen (77 K). As a note, desorption temperatures could be further increased by the increase of equilibrium pressure. On the basis of this, we take the equilibrium pressure obtained with the isothermal curves in the range (5–15 MPa) for Ca\(_{12}\)–7H\(_2\), Ca\(_2\)C\(_{12}\)–14H\(_2\), and Ca\(_3\)C\(_{12}\)–21H\(_2\) activated complexes, and we compare the thermal stability using eq 7. Here we used the average adsorption energy of all hydrogen molecules adsorbed by the doped complex, and we give an estimate of desorption temperatures (Figure 7d). At pressures of 50, 100, and 150 MPa, the average \(T_D\) is 384, 407, and 423 K for the Ca\(_{12}\)–7H\(_2\), Ca\(_2\)C\(_{12}\)–14H\(_2\), and Ca\(_3\)C\(_{12}\)–21H\(_2\) doped complex, respectively. The highest gravimetric densities for each of the systems are 7.11 wt % (Ca\(_{12}\)–7H\(_2\)), 11.17 wt % (Ca\(_2\)C\(_{12}\)–14H\(_2\)), and 13.8 wt % (Ca\(_3\)C\(_{12}\)–21H\(_2\)).
Figure 8. continued
In addition, we also performed DFT-based molecular dynamics for CaCl$_2$$^{-}$7H$_2$ (pristine and activated carbyne) and Ca$_3$Cl$_2$$^{-}$21H$_2$ systems with massive GGM and Nose–Hoover thermostats at 300 K. We observed unified thermal stability after 6 ps at 300 K for all molecular dynamics’ simulations when the Nose–Hoover thermostat was used. The average temperature was 300.94 K, and 300.23 K along the MD-production run for pristine and activated structures (Figure 8a–f). For doped complex Ca$_3$Cl$_2$$^{-}$21H$_2$, the average temperature was 301.44 K with the Nose–Hoover thermostat. Every system was equilibrated for 3–6 ps, and after 6 ps of production, no breaking of bonds was observed, which implied the thermal stability of the systems. In addition, using the Nose–Hoover thermostat, we have carried out an MD production run at 200 and 100 K for the CaCl$_2$$^{-}$7H$_2$, and Ca$_3$Cl$_2$$^{-}$21H$_2$-doped complex (pure and activated), which is shown in the Supporting Information.

Along with the previous studies in this work, we can determine that activated carbyne C$_{12}$-ring with zinc dichloride and decorated with Ca atoms can adsorb seven H$_2$ molecules with an average energy of $\sim$0.39 eV per H$_2$ molecule corresponding to 7.11 wt %. It is expected that activated C$_{12}$-ring might bind three Ca atoms around the inner/outer surface with their H$_2$ molecules, respectively, which represents an increase up to 13.8 wt % with respect to the pristine C$_{12}$-ring previously investigated. This storage capacity satisfies the requirements established by the U.S. Department of Energy by the end of the year 2025. Therefore, the considered material might be a promising choice for efficient hydrogen storage media, so this material certainly requires further experimental investigation.

**DISCUSSION**

The research for hydrogen storage materials is very attractive for fuel cell applications among others. Nevertheless, it is a great challenge to find hydrogen storage materials with great hydrogen gravimetric density under ambient thermodynamic conditions. Previous studies have explored that pristine nanomaterials$^{6-13}$ cannot efficiently store hydrogen, mainly due to weak van der Waals interaction between the hydrogen molecules and host material. Pristine carbyne is not an exception; for example, in the C$_{12}$-ring structure only a single H$_2$ molecule can adsorb with adsorption energies of 0.077 and 0.028 eV at the outer and inner surfaces, respectively, by DFT calculations presented in this analysis, making unsuitable its use as a hydrogen storage material. Previous research has shown that the pristine carbyne C$_{12}$-ring$^{21}$ structure is the ground state. We take this structure for new experimentation, activating with ZnCl$_2$. Before chemical activation, we calculated the area of pristine carbyne and pore diameter (18.37 Å$^2$ and 4.83 Å) presented in this work. Even more, we determined the specific surface area shows a value of 4606.31 m$^2$ g$^{-1}$, and accessible surface area shows a value of 13,000 m$^2$ g$^{-1}$. The chemical activation helps to develop the porosity by means of dehydration with ZnCl$_2$ or H$_3$PO$_4$ of the biomass structure. In this work, we take the pure C$_{12}$-ring and then we activate it with ZnCl$_2$, as shown in the methodology in this work. By applying geometry optimization and removing the ZnCl$_2$ the area and pore diameter result in 19.32 Å$^2$ and 4.96 Å, respectively, representing...
increases of 5.17% and 2.69% with respect to pristine carbyne. The SSA and ASA show values of 4844.53 m² g⁻¹ and 15.047.88 m² g⁻¹, respectively. These represent increases of 5.17% and 15.75% with respect to the pristine carbyne. However, the activated carbyne does not enhance the hydrogen storage properties, since only two H₂ molecules can be adsorbed with adsorption energy higher than 0.1 eV at the center of the activated carbyne, which does not meet the goals established by the DOE.

In addition, to improve the hydrogen storage properties, metal doping is one effective method to enhance the adsorption energies between H₂ molecules and host material. Previous research has shown that pristine carbyne C₁₂-ring decorated with Ca atoms might bind up to six H₂ molecules with average binding energies of 0.18 eV per H₂ (PW91) and 0.32 eV per H₂ with the empirical correction scheme of Grimme (DFT-D).²¹ Nevertheless, only a single Ca atom might be bound to the carbyne molecule with binding energy (E_b) higher than 2 eV, reaching 6.16 wt % for the gravimetric hydrogen storage. Even for cluster C₁₀⁰ rings in polyene and cumulenic forms, only a single Ca atom might be chemisorbed with binding energies greater than 2.5 and 2.33 eV for GGA-PW91 and GGA-PBE functionals, respectively. Up to either six or seven H₂ molecules are physisorbed by Ca atom, with average energies of 0.22 eV per H₂ (PW91) and 0.263 eV per H₂ (DFT-D) for cumulene and polyene molecules, respectively. The hydrogen storage capacity obtained corresponds to 7.02–8.09 wt %. Thus, hydrogen storage capacities using only one Ca atom in carbyne rings reach H₂ storage capacities, which are much higher than other carbon- and boron-based materials (see Table 2), where other investigations saturate the system with more dopant atoms until reaching the goal of DOE requirements. In this work, we demonstrate that activated carbyne can bind up to three calcium atoms around this surface, with binding energy greater than 2.7 eV per Ca atom, which represents up to an increase of 49.77% with respect to the pristine CaC₁₂ complex, indicating that the system is strongly chemisorbed and up to seven H₂ molecules can be physisorbed with an average energy of 0.39 eV per H₂. Reaching 13.8 wt % for gravimetric storage capacity, fulfilling the requirements by the DOE. In addition, we also performed DFT-based molecular dynamics for the C₁₂ and CaC₁₂ systems in their pristine and activated with ZnCl₂ to study the structural stability of the molecules. We determine the equilibrium pressure by means of adsorption isotherms and the van’t Hoff equation, which suggests that Ca-decorated carbyne could operate as hydrogen storage media at temperatures above the boiling temperature of liquid nitrogen presented in this work.

CONCLUSIONS

We performed the analysis of activated carbyne C₁₂-ring with zinc dichloride (ZnCl₂) and decorated it with Ca atoms by DFT calculations. First, the pristine carbyne C₁₂-ring used in this work corresponds to the C₂₀₉ structure with N = 3, with a D₉₀ point group symmetry. The pristine C₁₂-ring exhibits area and pore diameter of 18.37 Å² and 4.83 Å, respectively. According to this, our results using activated carbyne C₁₂-ring show an increase in area and pore diameter of 5.17% and 2.69%, respectively. In the case of several carbon-based nanomaterials, the hydrogen adsorption on pristine carbyne is impractical due to weak interactions of H₂ molecules on the host material, since the adsorption energies of a single H₂ molecule at the outer and inner surfaces of pristine carbyne are only 0.077 eV per H₂ and 0.028 eV per H₂, respectively. In order, to increase the hydrogen storage properties on pristine C₁₂-ring, we activate the ring through chemical activation with zinc dichloride (ZnCl₂) to increase the surface area and enhance pore size distribution. These theoretical specific surface area on activated carbyne results in an increase from ∼4606.31 m² g⁻¹ (pristine C₁₂-ring) to ∼4844.33 m² g⁻¹. The accessible surface area shows a value of 13000 m² g⁻¹ for pristine carbyne, whereas activated carbyne shows a value of 15045.88 m² g⁻¹, which shows an increase of 15.76% with respect to that of the pure carbyne. These values represent a macroscopic parameter that can be helpful to adjust the synthesis condition of the carbyne molecule. Unlike pristine C₁₂-ring, the activated ring can adsorb two H₂ molecules at the center of the molecule with a desirable binding energy of ∼0.1 eV per H₂. However, the hydrogen storage capacity obtained is only 2.72 wt % which does not fulfill objectives established by the DOE. Subsequently, we studied the case of a single Ca atom adsorbed on a carbyne surface (outer and inner), and we calculate the binding energy of the CaC₁₂ system, showing values of 2.79 eV for single and triple bond zones, where Ca atom was placed, and of 3.34 eV for the center of the activated ring, which represents an increase of 49.77% with respect to the pristine carbyne. In addition, we determine the zones that Ca atoms prefer to bond on the carbyne molecule (in front of single and triple bonds and the center of the ring), which makes it more attractive in comparison with pristine C₁₂-ring, which only has an area for placement of the calcium atom in front of single bonds of the ring. The activation of the carbyne ring produces higher stability on the ring, causing more Ca atoms that can bind to the host material. We showed that Ca adatoms at small concentrations stay atomically dispersed on carbyne, donating +0.94 e and +1.05 e to the ring, for Mulliken population analysis and ESP-fitted charges, respectively. Furthermore, in the presence of Ca, hydrogen adsorption increases 21.8% in comparison with Ca-decorated pristine carbyne. We determine that up to seven H₂ molecules can be physically adsorbed with an average energy of ∼0.39 eV per H₂ molecule. The first six H₂ molecules tend to adsorb around the Ca atom and the seventh H₂ molecule is adsorbed on the top of the Ca atom. The hydrogen storage capacity obtained in this study is 7.11 wt % and therefore represents an increase of 15% with respect to the pristine carbyne. However, it is expected to reach 13.8 wt % with three Ca atoms, which represents an increase of 124% with respect to pristine carbyne, and satisfactorily meets the target set by the DOE for the year 2025. We determine the equilibrium pressure for CaC₁₂−7H₂ and CaC₁₂−21H₂ systems (5–15 MPa), by isotherm calculations. Furthermore, the Tₚ₀ which is calculated using the van’t Hoff equation, suggests that Ca-decorated carbyne could operate as hydrogen storage media at temperatures above the boiling temperature of liquid nitrogen. The molecular dynamics after 6 ps at 300 K show unified thermal stability when the Nosé−Hoover thermostat was used. The average temperatures were 300.94 and 300.23 K for CaC₁₂−7H₂ (pristine and activated carbyne). For doped complex Ca₅C₁₂−21H₂, the average temperature was 301.44 K with the Nosé−Hoover thermostat. Every system was equilibrated for 3−6 ps, and after 6 ps of production, no breaking of bonds was observed, which implied the thermal stability of the systems. Therefore, the activated carbyne decorated with Ca atoms attains H₂ storage capacity which is much higher than other carbon and carbon-based materials reported in the literature (see Table 2), fulfilling DOE requirements, so this material certainly requires further experimental investigation.
Density functional theory calculations are carried out to activate carbyne C_{12}-ring through chemical activation with zinc dichloride (ZnCl₂) and decorating with Ca atoms, by means of Biovia Materials Studio Dmol³ software⁵⁵,⁵⁸ to determine its capability of hydrogen storage. To calculate adsorption energies, with one C_{12}-ring are calculated by means of a set of double numerical plus polarization basis (DNP). For occupied orbital, we consider two atomic orbitals in the basis set. For C and H atoms, d- and p-polarization functions are respectively used. The employed basis set has the advantage of being equivalent to the analytical basis set 6-31G**. All presented geometry optimizations are obtained for a tolerance on which the maximum forces are lower than 0.002 Ha/Å. Here the effect of van der Waals interactions is included explicitly by using the empirical correction scheme of Grimme (DFT-D) for periodic systems.⁶⁰

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