Unique Hydrogen Adsorption Properties of Graphene

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Graphene showed an unusually high hydrogen storage capacity as well as a unique, slow hydrogen adsorption process compared with a variety of carbon materials (carbon nanotubes, activated carbons, mesoporous carbons, templated carbons, and metal organic frameworks). Catalytic dissociation of hydrogen on graphene is observed for the first time. The hydrogen dissociation rate on graphene is also significantly faster than the adsorption rate. This leads to the conclusion that the most active sites on graphene for hydrogen dissociation are not the only sites where the enhanced adsorption occurs. The mechanistic differences in adsorption on graphene when compared with the other carbons are further demonstrated by differences in temperature-programed desorption results and heats of adsorption.

Keywords: absorption, materials, adsorption/gas, surface chemistry/physics, catalysis

Introduction

Since the first isolation of graphene from graphite,1 graphene has been synthesized by various routes2–5 and has shown a wide variety of fascinating properties, such as the quantum Hall effect at room temperature (RT), extremely high carrier mobility, ambipolar field effect, and sensitive response to single molecules.6,7 The recent theoretical calculation results indicate that graphene may also work as a sorbent for hydrogen storage.8–10 Various nanostructured and microporous carbon-based materials including carbon nanotubes (CNTs), activated carbons (ACs), mesoporous carbons (MCs), templated carbons (TCs), and metal organic frameworks (MOFs) have been studied intensively because of their light weight, high surface areas, and relative chemical stabilities.11–17 However, recent results showed that these carbon-based materials cannot store a sufficient amount of H2 required for transportation applications merely by physical adsorption at ambient temperature. Novel storage sorbent materials and ways to further enhance the storage capacities need to be explored.

The promising results predicted for hydrogen storage on ideal graphene have stimulated experimental studies.8–10 However, experimental studies have been limited to hydrogen adsorption on low-surface-area graphene-like materials18–20 and the hydrogenation of supported graphene membrane by plasma exposure or the use of a hydrogen atom beam source.21,22 An in-depth study of hydrogen adsorption on graphene and an understanding of the adsorption process are desired. In this work, we studied the hydrogen adsorption properties of graphene with a high surface area of 2139 m²/g and compared it with the most studied carbon materials (for H2 storage). Among all the tested carbons, graphene showed a superior hydrogen storage capacity as well as displaying a uniquely slow hydrogen adsorption process. We have also observed catalytic hydrogen dissociation on graphene for the first time and compared its similarities to metal-doped carbons.

Experimental

Synthesis

The graphene was prepared by following a published solvothermal and sonication procedure.4 Typically, 2 g of sodium...
and 5 mL of ethanol (molar ratio, 1:1) were sealed in a teflon-lined reactor vessel and heated at 493 K for 72 h. After that the reactor vessel was cooled to RT and a solid solvothermal product (graphene precursor) was obtained. This material was then transferred to a horizontal quartz tube in He atmosphere and rapidly pyrolyzed at 1323 K for 2 min. The resulting black product was washed with copious amounts of deionized water and sonicated in ethanol. The final product was collected by filtration and dried in vacuo at 373 K for 24 h.

Mesoporous carbon, templated carbon 1 (TC-1), templated carbon 2 (TC-2), templated carbon 3 (TC-3), and MOF-177 were prepared according to the published procedures.23–28 The superactivated carbons AX-21 and Maxsorb were obtained from Anderson Development Company and Tokyo Zairyo, respectively. CNT-1 was obtained from Shenzhen Nanotech Port. The storage capacity of CNT-2 was cited from previous literature.29 A total of 6 wt % Pt-doped AX-21 was synthesized according to previous literature.30

**Characterization**

Nitrogen adsorption isotherms and low-pressure H₂ adsorption isotherms (0–1 atm) were measured by using a Micromeritics ASAP 2020 Analyzer. Nitrogen adsorption isotherms were measured at 77 K. Low-pressure H₂ adsorption isotherms were measured at 273, 298, and 323 K. Circulating baths were used to maintain the sample at 298 or 323 K. An ice-water filled dewar flask was used to maintain 273 K, and liquid nitrogen was used for 77 K. High-resolution transmission electron microscopy (HRTEM) images of the materials were obtained on a JEOL 3011 analytical electron microscope equipped with EDX analysis operated at 300 kV. The elemental ratio of graphene was approximately C: 93%, O: 6.5%, and Na: 0.5% as analyzed by a Kratos Axis ultra XPS spectrometer (Supporting Information Figure S1).

Hydrogen adsorption at 298 K and pressures greater than 0.1 MPa and up to 10 MPa was measured using a static volumetric technique with a specially designed Sieverts-type apparatus. The apparatus was previously tested and proved to be leak free and accurate through calibration by using LaNi₅, AX-21 carbon, zeolites, and MOFs at 298 K.31 As shown in our previous work, the high-pressure isotherms of the commercially available superactivated carbons (i.e., AX-21, GX-31, and Maxsorb) are known and have been used as standard materials for testing the measurements. For a superactivated carbon with a BET surface area of 2800 m²/g, the isotherm at 298 K should be slightly concave with an uptake of 0.6 wt % at 10 MPa.31 Approximately 200 mg of sample was used for each high-pressure isotherm measurement. Analysis of evolved gases during TPD experiments was performed with an AeroVac 1200 Magnetic Sector mass spectrometer (VTI).

**H₂/D₂/HD isotope equilibrium reaction and temperature-programmed desorption**

The H₂/D₂/HD equilibrium on graphene was measured using a slightly modified procedure proposed by Ishikawa et al.32 In this new procedure, H₂ and D₂ were mixed in a 1:1 ratio at 4.3 MPa. This mixture was allowed to equilibrate for 2 h at RT before being exposed to the sample at 1.6 MPa. The small sample cell was then closed and allowed to react for a given amount of time before being quenched in liquid nitrogen. The gas in the sample cell was analyzed using mass spectrometer (MS) after 5 min from initial sample cell submersion in liquid nitrogen. The MS was set in sweep mode to analyze (m/z) peaks 1–6 to fully differentiate hydrogen, deuterium hydride, and deuterium. Twenty-milligram samples of graphene, AX-21, and 6 wt % Pt-doped AX-21 were subjected to this treatment for a series of different reaction times and compared to the results obtained using a blank sample cell. All samples were degassed at 523 K for 8 h under vacuum, except Pt-doped AX-21, which was degassed at 623 K and an annealed graphene degassed at 823 K. A reaction rate constant was also obtained for the blank cell, which was made of stainless steel.

Temperature-programmed desorption (TPD) was performed on the samples once all physisorbed hydrogen was removed (20 min) and the m/z 2 peak had fallen to baseline. Heating at 10 K/min (±1 K/min) was performed while analyzing (m/z) peaks 1–6 to determine total hydrogen species desorbed.

**Results and Discussion**

The published solvothermal and sonication procedure3 was selected because the production of graphene can be in gram-quantities, which is beneficial for achieving accurate storage measurements and for use in further applications. The TEM image of the products (Figure 1) clearly shows the wrinkled graphene sheets and overlap between different sheets, which are in agreement with previous literature. A N₂ isotherm at 77 K on graphene showed an increase at a low relative pressure and a hysteresis loop at pressures from 40 to 100 kPa (Supporting Information Figure S2). The graphene had a BET surface area of 2139 m²/g, which was less than the 2630 m²/g for ideal single-layer graphene. This indicates some partial overlap of graphene sheets rather than completely separated single-layer graphene.

Hydrogen storage on graphene was investigated at 298 K and up to 10 MPa. As shown in Figure 2, graphene exhibited a surprisingly high storage capacity of 0.90 wt %. The hydrogen storages of a series of carbon materials including...
CNT, AC, MC, TC, and MOF were measured to compare with graphene. In Figure 3, there is a clear relationship between the storage capacity and the BET surface area for all other tested carbons (with surface areas ranging from 810 to 3840 m²/g), as expected for physical adsorption. This correlation is in agreement with previous reports. However, the storage capacity of graphene was well above the correlation. The capacity of 0.90 wt % for graphene was the highest compared with all the other carbons (0.62 wt % for AX-21, 0.72 wt % for Maxsorb, 0.66 wt % for MOF-177, and 0.82 wt % for TC-3). If we consider the correlation between the capacity and surface area and extrapolate linearly, a carbon with a BET surface area of 2139 m²/g should have a capacity of 0.56 wt % at 10 MPa. This showed that the storage capacity on graphene was enhanced by 60%. The reversibility was assessed by measuring the desorption branch down to 1 atm. It can be seen from Figure 2 that the desorption branch nearly coincided with the adsorption branch although there appeared to be a slight hysteresis. However, the next two adsorption branches were in agreement with the first adsorption branch indicating no observable loss in adsorption sites during adsorption–desorption cycles at 298 K (Figure 4).

To evaluate the interactions between graphene and H₂, the isosteric heats of adsorption were calculated from the adsorption isotherms at different temperatures by using the Clausius–Clapeyron equation (Figure 5). The hydrogen adsorption isotherms were measured at 273, 298, and 323 K, and the heat of adsorption values were determined using the slope of a plot of ln(P) vs. (1/T). The heats of adsorption were ~12 kJ/mol at low surface coverage and leveled off to ~8 kJ/mol at higher surface coverages. The high values of heat of adsorption at low surface coverage can be attributed to the adsorption of H₂ on the more energetic sites on graphene. Defect sites and edge sites (i.e., armchair and zigzag edge sites) are strong sites for adsorption. The heat of adsorption (~8 kJ/mol) on graphene at high coverages is clearly higher than that of H₂ on microporous carbons such as activated carbon (~5 kJ/mol).
To understand the high capacity of graphene and the adsorption mechanism, we studied the adsorption kinetics on graphene and compared it with other carbons (superactivated carbons AX-21 and Maxsorb, TC, and CNTs). The kinetics of uptake were studied by increasing the pressure in steps and following the fractional completion vs. time during each step. Figure 6 shows the adsorption kinetics of hydrogen on graphene and other carbons at ~6 MPa final pressure and 298 K. It is seen that the adsorption rates on graphene were clearly slower than on all other carbons. For all other carbons, the adsorption was completed within 10 min, whereas for graphene the completion for total uptake took ~25 min. This indicates a unique hydrogen uptake process that occurs on graphene. A detailed comparison of adsorption kinetics on graphene and AX-21 carbon at various pressures was carried out further. As shown in Figure 7, the adsorption on AX-21 at all pressures was completed within 10 min. For graphene, the completion time ranged from 12 to more than 30 min, and the kinetics became slower with the increase in pressure. The difference between graphene and AX-21 is more clearly shown by comparing their fractional uptake completion during individual pressure steps (Figure 8). Within 0.5 min, over 90% completion was accomplished on AX-21, whereas the remaining 10% was completed in less than 10 min. This can be understood because the fast adsorption of H₂ on AX-21 was via van der Waals interactions. In 0.5 min, the adsorption on graphene had completion fractions of 76% at 2.0 MPa, 53% at 6.3 MPa, and 33% at 7.9 MPa. The slower adsorption rates on graphene compared with AX-21 and other carbons indicate that a chemical process may be involved in addition to physical adsorption.

Previous theoretical studies showed that the edge sites or structural vacancies on graphene could dissociate H₂ into H atoms. Despite their activities, free edge sites (zigzag and armchair sites) and their joint corner sites could be stable enough to remain free (i.e., not hydrogenated) at ambient temperature. To gain further insight into the activities of graphene, the H₂/D₂/HD isotope equilibrium reaction was used. The reaction rate for graphene was measured using a slightly modified procedure proposed by Ishikawa et al (See experiment section). In this new procedure, a mixture of H₂ and D₂ (1:1 ratio) was allowed to mix uniformly (2 h) at RT in a stainless steel dosing cell before being exposed to the sample. The gas in the dosing cell was then introduced to the sample cell after which the two cells were then isolated. The gas in the sample cell was allowed to react with the sample for a specific amount of time (time at RT) before the sample cell was quenched in liquid nitrogen and remained there. Five minutes from the moment the sample cell was submerged in liquid nitrogen, the gas was sampled via a molecular leak valve and analyzed using a MS. Graphene (20 mg) was subjected to this treatment and compared to the results obtained on AX-21 (20 mg), 6 wt % Pt-doped AX-21 (20 mg), and the blank cell. The Pt-doped AX-21 is known to have catalytic activities and adsorbs hydrogen via a hydrogen dissociation/spillover process. The reaction H₂ + D₂ → 2HD (Kₑq ~ 3.5) was measured as a function of completion vs. time at RT, and a correction was added to the time at RT to account for the cooling process. In Figure 9, a first-order rate is shown for both the graphene and Pt-doped AX-21.
dissociation rates. Given the highly increased dissociation rates compared with AX-21 and the blank cell, it is clear that graphene and Pt-doped carbon have similar hydrogen dissociation capabilities. It is also clear that there is an additional hydrogen dissociation mechanism for graphene that was not observed by Ishikawa et al. for graphite.32 To our knowledge, this is the first time this highly increased catalytic activity for graphene (when compared with all other carbons) has been observed experimentally.

To gain insight into possible binding sites and the relation between adsorption and hydrogen dissociation, we also measured TPD spectra for graphene (as explained in Experimental section.). The TPD spectrum continues from the end of the H2/D2/HD reaction and starts by evacuating all gas from the sample while still at liquid nitrogen and heating at a specified rate all while monitoring the evolved gas using the same MS. These spectra were taken after time at RT of 0.5 min, 1.5 min, and 2 h. As shown in Figure 10, TPD for time at RT of 2-h adsorption shows a clear peak with isotopic effects above RT (hydrogen comes out in higher concentration than is present in the system). At only 1.5 min at RT, this peak disappears and a new peak (assumed to be hidden) becomes apparent at lower temperature. At 0.5 min at RT, only the peaks due to physically adsorbed species seem to be present. Previous work on hydrogen spillover suggests that reverse spillover and surface recombination of H/D will be the two probable methods for hydrogen desorption.41 We have assigned the peak at higher temperatures to surface recombination due to isotopic effects (i.e., the lightest species are in greatest concentration). A blank TPD and plain AX-21 showed no peaks except the typical rise in hydrogen when approaching the degas temperature (starting around 473 K) and the physical adsorbed hydrogen during initial heating for AX-21. TPD for AX-21 is provided (Supporting Information Figure S3).

The recombination and subsequent desorption is inherently measured in the H2/D2/HD experiment showing that the bond formed between these most active graphene dissociation sites and hydrogen is very weak. Further, the reaction rate shown here for dissociation (Figure 9) is faster than the completion rate for hydrogen uptake. This shows that the sites responsible for hydrogen dissociation are not the only sites increasing the hydrogen storage. A degas temperature of 823 K also shows that effective sites are stable at high temperatures. Theoretical and experimental studies suggest that a functional on the graphene (the zigzag edge) is the most reactive site and would be capable of dissociating hydrogen.36,38 This may be able to explain the difference between adsorption and the hydrogen dissociation rates. Other studies suggesting direct dissociation on the basal plane may be able to explain the slow adsorption rates as an alternative to a possible spillover mechanism.42 The lack of studies on other functional groups indicate they should not be ruled out, including the possibility of sodium implanted into the carbon lattice or as a salt with carboxylic groups on the edges. However, further experiments are warranted for understanding the hydrogen adsorption sites present on graphene and the effect of oxygen and sodium.

Conclusions

In summary, hydrogen adsorption properties of graphene have been studied and compared with a diverse collection of
Graphene shows an unusually high hydrogen storage capacity and a unique hydrogen adsorption process. The dissociation of hydrogen on graphene has been observed experimentally for the first time. We have also shown that the hydrogen desorption rate is significantly faster than the adsorption rate. This leads us to conclude that the most active sites for dissociation are not the only sites where the enhanced adsorption occurs. The slow adsorption rates and TPD results show adsorption mechanism differences in graphene compared with activated carbon and demonstrate the unique hydrogen storage capabilities of graphene.

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Literature Cited


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