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Hydrogen Storage in High Surface Area Carbon Nanotubes Produced by Catalytic Chemical Vapor Deposition

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Carbon nanotubes, mostly single- and double-walled, are prepared by a catalytic chemical vapor deposition method using H$_2$–CH$_4$ atmospheres with different CH$_4$ contents. The maximum hydrogen storage at room temperatures and 10 MPa is 0.5 wt%. Contrary to expectations, purification of the carbon nanotube specimens by oxidative acid treatments or by heating in inert gas decreases the hydrogen storage. Decreasing the residual catalyst content does not necessarily lead to an increase in ASH. Moreover, increasing the specific surface area does not necessarily increase the hydrogen storage capacity. There seems to be a correlation between the pore volume at low pore diameters (<3 nm) and the hydrogen storage capacity. Contribution from nanoscale disordered carbon to the hydrogen storage cannot be ruled out.

Introduction

Because of their large surface area, low density, and hollow structure, carbon nanotubes (CNTs) have been considered to be potential materials for hydrogen storage. Dillon et al. reported for the first time excellent hydrogen storage properties for a sample containing unpurified single-walled CNTs at 273 K. Theoretical estimates based on Monte Carlo simulations in a grand canonical ensemble for closed and open CNTs have been reviewed by Zuttel et al. who emphasized that well-characterized samples combined with appropriate experimental techniques for measurement of hydrogen storage were necessary to obtain reproducible data.

In this context, we report the measurement of amount of stored hydrogen (ASH, in wt %) at room temperature and 10 MPa by a volumetric method on high specific surface area CNTs prepared by a catalytic chemical vapor deposition (CCVD) route. The advantage of this synthesis technique is the formation of predominantly individual, as opposed to bundled, CNTs which results in near-theoretical specific surface areas. The effect of standard purification treatments for CNTs on the hydrogen storage capacity was also tested. The present results show that the ASH values for a mixture of single- and double-walled CNTs of diameter in the range 0.7–3 nm and specific surface areas of 550–850 m$^2$/g lie in the range of 0.3–0.5 wt % at room temperature and 10 MPa. It was observed that the ASH values decrease after purification of CNTs with nitric acid treatment or by heating in an inert gas. The samples were characterized by chemical analysis, Raman spectroscopy, electron microscopy and porosity measurements. The observed ASH values seem to depend on pore volumes in the less than 3 nm pore diameter range as calculated from adsorption isotherms for N$_2$ gas at liquid N$_2$ temperatures.

Experimental Section

Synthesis of Carbon Nanotubes. An oxide solid solution of the type Mg$_{0.95}$Co$_{0.05}$O was produced by combustion of the corresponding nitrates with urea, the details of which are given elsewhere. On heating the solid solution in H$_2$–CH$_4$ atmosphere, cobalt particles in the size range of 1–6 nm were produced in situ at a high temperature (1000 °C) which catalyzed the decomposition of CH$_4$ and the formation of CNTs. The obtained carbon nanotube–Co–MgO composite powder was dropped into 1 M HCl at 25 °C to separate the CNTs without damaging them. The separated CNTs were washed with water and dried at 80 °C. It was possible to produce more than 1 g of separated CNTs in every run.

The important parameters in the above-mentioned synthesis are the concentration of active cobalt particles at a particular temperature and the availability of methane gas at this temperature. For the present study, the cobalt content in the oxide solid solution was fixed (Mg$_{0.95}$Co$_{0.05}$O) and the CH$_4$ partial pressure in the H$_2$–CH$_4$ atmosphere was varied in the range 20–50 mol %. The samples thus prepared are named D1, D2, D3, and D4 for CH$_4$ contents of 20, 30, 40, and 50 mol %, respectively.
**Characterization.** The CNTs specimens were characterized by scanning electron microscopy and high-resolution transmission electron microscopy (TEM). The carbon content was analyzed by flash combustion, and the cobalt content was calculated therefrom. These values were confirmed by EDX spectra recorded on selected areas at a magnification of 100 000 during TEM observations. Raman spectroscopy was used to assess the nature of carbon. Raman spectra were recorded using a Micro-Raman setup (DILORXY), using backscattering geometry at 488 nm (3–8 mW). The sample was placed on a bilayer substrate to enhance the Raman signal or alternatively on a microscope slide. Low-frequency breathing modes were used to estimate the CNTs diameter distribution in the less than 1.8 nm range. Comparison of the relative intensities of the G band (ca. 1580 cm\(^{-1}\)) and the D band (ca. 1360 cm\(^{-1}\)) were used to qualitatively estimate the amount of disordered carbon.

**Purification of Carbon Nanotubes.** Various methods have been reported in the literature for the purification of CNTs. The principal impurity in the present CCVD CNTs specimens consists of poisoned cobalt particles encapsulated in carbon shells which resist the HCl attack. To decrease the proportion of this metal impurity, two methods were used: a nitric acid treatment followed by a heat-treatment in air and a heat-treatment in argon at elevated temperatures followed by another HCl attack.

Following experimental methods reported in the literature, selected CNTs specimens (D1 and D3) were boiled in concentrated HNO\(_3\) at 120 °C for 4 h. After centrifugation, washing and drying, the specimens were heated (heating rate 4 °C/min) in air up to 580 °C and maintained at this temperature for 10 min. The total weight loss after the two treatments was 80% in some cases, showing that some fraction of the CNTs were also destroyed. The chemical purity of the sample was assessed by EDX spectra on selected area. The amount of disordered carbon was assessed using Raman spectroscopy. The so-produced samples are named D1* and D3*. Carbon element analysis of nitric acid purified samples did not yield conclusive results due to the presence of oxygen in the form of functional groups on the surface of the CNTs.

In the second purification method, selected specimens (D1, D2, and D3) were heated in flowing argon atmosphere to 1000 °C, maintained for 30 min, and then cooled in argon. After the heat-treatment, the samples were dropped in 6 M HCl. Effervescence due to evolution of H\(_2\) was seen and the solution turned rose showing the dissolution of cobalt metal. These samples were termed D1′, D2′, and D3′. The carbon content was determined by flash combustion.

**Determination of the Amount of Stored Hydrogen (ASH).** The amount of stored hydrogen (ASH, in wt %) was measured by a volumetric method with hydrogen pressured up to 10 MPa at room temperature. The measuring device is shown in Figure 1. The ASH is obtained from the desorbed gas amount, which is measured by the accurate flow-meter with 0.1 wt % of measurement error. Pressure is corrected every time by measuring the gas amount in the blank holder before measuring the gas amount in the sample holder. The line unit temperature is controlled uniformly.

**Figure 1.** Schematic description of the device for the measurement of the amount of stored hydrogen (ASH).

**Figure 2.** Specific surface area (SSA) and amount of stored hydrogen (ASH) for CNTs samples prepared using different CH\(_4\) content in the H\(_2\)–CH\(_4\) atmosphere.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Carbon (wt %)</th>
<th>Cobalt (wt %)</th>
<th>BET SSA (m(^2)/g)</th>
<th>Pore vol &lt; 3 nm (cm(^3)/g Å)</th>
<th>ASH (wt %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>D1</td>
<td>74</td>
<td>21.0</td>
<td>587</td>
<td>0.098</td>
<td>0.34</td>
</tr>
<tr>
<td>D2</td>
<td>83</td>
<td>15.8</td>
<td>763</td>
<td>0.123</td>
<td>0.42</td>
</tr>
<tr>
<td>D3</td>
<td>90</td>
<td>9.0</td>
<td>881</td>
<td>0.242</td>
<td>0.51</td>
</tr>
<tr>
<td>D4</td>
<td>91</td>
<td>9.0</td>
<td>885</td>
<td>0.127</td>
<td>0.46</td>
</tr>
<tr>
<td>D1*</td>
<td>86</td>
<td>0.2</td>
<td>612</td>
<td>0.118</td>
<td>0.25</td>
</tr>
<tr>
<td>D3*</td>
<td>85</td>
<td>9.7</td>
<td>746</td>
<td>0.125</td>
<td>0.25</td>
</tr>
<tr>
<td>D2*</td>
<td>86</td>
<td>2.9</td>
<td>736</td>
<td>0.120</td>
<td>0.41</td>
</tr>
<tr>
<td>D3*</td>
<td>89</td>
<td>2.4</td>
<td>659</td>
<td>0.110</td>
<td>0.27</td>
</tr>
</tbody>
</table>

* See text for details on the purification procedures.

**Results and Discussion**

As-prepared (unpurified) carbon nanotubes. Carbon element analysis of specimens D1–D4 (Table 1) showed that the carbon content is increased as the CH\(_4\) content is increased from 20 to 50 vol % (74, 83, 90, and 91 wt % for D1, D2, D3, and D4, respectively). The cobalt content accordingly decreases (Table 1). The BET SSA and ASH of samples D1–D4 are reported in Table 1 and Figure 2. The BET SSA increases from D1 (587 m\(^2\)/g) sharply up to D3 (881 m\(^2\)/g) thereafter, showing a plateau. The ASH at room temperature and 10 MPa increases up to D3 and sharply decreases thereafter. The ASH for all samples are less than 1 wt %. This value, while being considerably lower than that reported by Dillon et al., is in general agreement with the values reported in the review by Züttel et al. for CNTs samples from various sources.

Samples D1–D3 were studied more closely in order to investigate the influence of the CH\(_4\) content during the synthesis.
Increasing it will tend to increase the carbon content in the samples, as shown above, but two distinct reasons, or a combination of both, can account for this result. Indeed, a key point of the synthesis is to match the carbon source (here CH₄) to the metal content so as to produce as many CNTs as possible through the catalytic decomposition of CH₄ at the surface of the Co particles, while avoiding the thermal decomposition of CH₄ which will produce disordered, nontubular carbon. So the carbon content increase could reflect a higher CNTs content in the sample and/or the formation of nontubular carbon.

Typical TEM images are shown in Figure 3, revealing CNTs and cobalt particles that appear with a dark contrast (Figure 3a, Figure 3b), a double-walled nanotube (Figure 3c), and two side-by-side double-walled nanotubes (Figure 3d). The distributions of the number of walls of CNTs in D1 and D3 as well as the distribution of the inner and outer diameters, derived from the measurements of 128 individual CNTs for each sample on high-resolution TEM images, are shown in Figure 4. About 90% of the CNTs are either single- or double-walled. The inner and outer diameter distributions are in the ranges 0.5—3.0 and 0.75—4.0 nm, respectively. TEM images also showed a lower content of poisoned (i.e., inactive) cobalt particles with increasing CH₄ content. Thus, it appears that the increased partial pressure of methane helps in preventing the poisoning of the cobalt particles, thereby accounting at least partly for the formation of more CNTs. Poisoned cobalt particles are the main impurity in the case of D1, D2, and D3 show in addition some disordered carbon which appear as flakes, the detailed structure of which is difficult to resolve. The flakes apparently consisted in one or two graphene sheets, and the CNTs were seen embedded on the flakes, forming an intimate composite structure (Figure 5).

Raman spectra of D1—D3 (Figure 6) show an increasing D band, revealing that the proportion of disordered carbon increases as the methane content is increased, in good agreement with the TEM study. The low-frequency radial breathing modes for specimen D1 (inset) shows several peaks corresponding to CNTs with different diameters in the range 0.75—1.8 nm, showing that no one diameter is preferred. In fact, the very low-diameter CNTs are found to be mostly the internal wall of double-walled tubes.¹⁹

**Purified Carbon Nanotubes.** After nitric acid treatment followed by oxidation in air, the cobalt contents could be reduced to less than 1% in D1* and D3*. However, EDX analysis revealed that the purified samples showed the presence of oxygen, as compared to none at all in the as-prepared samples. It is possible therefore that extensive CNTs opening and functionalization takes place during the purification. IR spectra (not shown) have revealed the presence of carbonyl groups.

Figure 7a shows a typical image of D1*, revealing a very clean specimen. A closer look at the TEM images of purified samples showed that many of the open CNTs were filled with disordered carbon (Figure 7b). Thus, the nitric acid treatment cuts the CNTs and probably creates fragments of disordered carbon and these fill the CNTs during the thermal treatment in air. Raman spectra showed that little or no disordered carbon is present after purification. The Raman spectrum for D1* (Figure 8) shows a low-intensity D band. The BET SSA of D1* is of the order of that of D1, while the BET SSA of D3* (313 m²/g) is much lower than that of D3 (881 m²/g). This could reflect the initial proportion of nontubular carbon in D3.

The purification treatments, contrary to expectation, decrease the ASH values (Table 1). The decrease is particularly evident in the case of D3 and D3*, but despite that all characterization show the high chemical purity and structural homogeneity of D1*, the ASH value also show a decrease. Thus, the presence of metal impurity has little or no influence on the ASH values. The presence of open CNTs also does not influence the ASH values. It is probable that the opening of the present CNTs is accompanied by their functionalization which blocks potential adsorption. Indeed, Hirscher et al.¹⁸ have studied the influence of cutting CNTs on hydrogen adsorption and concluded that cutting has to be clean and hydrogen has to be introduced in...
situ in order to increase adsorption. The sharp reduction of ASH values from D3 to D3* indicates that the presence of disordered carbon could also play an important role in the adsorption of hydrogen.

The first step of the second purification procedure is a heat-treatment in argon at 1000°C. No major weight loss was observed after the heat-treatment. However, TEM images and EDX spectra showed that some poisoned cobalt particles migrate from their carbon shells and coalesce to form micrometer-sized spheres, leaving behind empty carbon shells. However, some cobalt particles, particularly those larger than 10 nm did not migrate out of their shells. How this phenomenon was produced in the absence of oxygen (no weight loss was observed), remains unclear. Perhaps cobalt particles in the size range of 1–6 nm are highly mobile at 1000°C and hence can diffuse out of the carbon shells which surrounds them even if there is no oxidation of the carbon shell. Some CNTs are open and disordered carbon is seen inside and on the outer walls. It is interesting to note that in contrast to CNTs prepared by arc discharge and laser ablation of graphite, the present CCVD CNTs are not stable at temperatures higher than 1000°C even in inert atmosphere. This instability may be due to the presence of more defects in the as-prepared CNTs due to the lower synthesis temperature or to the presence of extremely small catalyst particles (0.8–3 nm) which may be highly mobile at temperatures much lower than the melting point of cobalt.

The second step of this purification procedure is a soaking in 6 M HCl, which dissolves the now unprotected cobalt particles, giving rise to a strong effervescence. Element analysis show that the cobalt content is much lower in the D1′–D3′ samples than in the parent D1–D3 specimens (Table 1). No functionalization of the CNTs was observed and a high BET SSA (Table 1) were maintained. The ASH values are also decreased for these purified specimens, except for D2′ which shows an ASH very similar to than of D2 (Table 1).

Thus, both the acid purification treatment and the argon treatment reduce the ASH values even though the acid treatment tends to decrease the specific surface area and the argon treatment increases it slightly. Thus, it appears that although
the trend is that the ASH increases with increasing SSA (Figure 9a), it does not depend only on SSA, and other factors are involved.

To explain these results, the adsorption properties of the different samples were studied by measuring the N\textsubscript{2} adsorption isotherm at liquid N\textsubscript{2} temperatures. The pore size distributions were calculated using the BJH equation from the adsorption branch of the isotherm. The cumulative pore volume in the range 1.8–3 nm was estimated therefrom (Table 1). The pores of higher dimensions were disregarded as being due to the secondary structure of nanotubes. The ASH values are plotted as a function of this pore volume (Figure 9b). There seems to be a correlation between the pore volume in the <3 nm range and hydrogen storage.

The pore size distributions as calculated using the BJH equation are shown in Figure 10 for D1–D3 and D1*–D3*. In the less than 5 nm range, the distribution consists of three peaks at 2.0, 2.2, and 2.8 nm. The pores at 2.2 and 2.8 nm are high in D3, which showed the highest ASH, and decrease drastically after purification for D3*, which showed the lowest ASH (Table 1). These pores could arise from nanostructured disordered carbon. However, note that D3 and D3* were also
extremes in SSA (Table 1). The case of D1−D1* is somewhat different, since the ASH for D1 is higher than for D1*, despite D1 having both a lower SSA and pore volume <3 nm (Table 1). The pore at 2.0 nm is increased after nitric acid treatment and could be related to adsorption inside the CNTs, perhaps reflecting some degree of CNTs opening.

The argon treatment does not change the pore distributions significantly (not shown in the figure) showing that the empty nanoshells do not contribute to the SSA and porosity of the samples.

Conclusions

The hydrogen storage at room temperatures and 10 MPa pressure was studied in carbon nanotubes (mostly single- and double-walled) prepared by a catalytic chemical vapor deposition method. Under these conditions, the maximum hydrogen storage obtained is 0.5 wt %. This value for hydrogen storage is much lower than that reported by Dillon et al.1 but is in good agreement with those reported in the review by Zuettel et al.8 Purification of the carbon nanotubes by oxidative acid treatments or by heating in inert gas decreases the hydrogen storage. Decreasing the residual catalyst content does not necessarily lead to an increase in ASH. Also, increasing the specific surface area does not necessarily increase the hydrogen storage capacity.

There seems to be a correlation between the pore volume at low pore diameters (<3 nm) and hydrogen storage capacity. Contribution from nanoscale disordered carbon to the hydrogen storage cannot be ruled out. An in-depth characterization of carbon nanotubes samples, as-prepared and after purification, is essential in order to try and understand the complex phenomena involved in hydrogen storage.

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References and Notes