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Abstract	For the Hydrogen economy, cost-effective and safe storage of hydrogen assumes importance. Among the various modes of storage, solid-state storage has definite advantages for mobile and stationary applications. However, among the probable solid-state materials, the desired levels of storage (~6 Weight %) can be possible in carbon materials suitably modified with activation centers. A variety of modifications of carbon materials have been examined for hydrogen storage. However, the search has to continue till the desired levels of storage under ambient conditions are achieved.		
Keywords (separated by '-')	Hydrogen storage - Carbon nanomaterials - Graphene - Activation centers - Carbon allotropes - Solid-state storage		

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Carbon Nanomaterials as One of the Options for Hydrogen Storage



B. Viswanathan

Abstract For the Hydrogen economy, cost-effective and safe storage of hydrogen

- ² assumes importance. Among the various modes of storage, solid-state storage has
- ³ definite advantages for mobile and stationary applications. However, among the prob-
- ⁴ able solid-state materials, the desired levels of storage (~6 Weight %) can be possible
- ⁵ in carbon materials suitably modified with activation centers. A variety of modifica-
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⁹ Keywords Hydrogen storage · Carbon nanomaterials · Graphene · Activation

¹⁰ centers · Carbon allotropes · Solid-state storage

11 **Introduction**

12 The energy needs of the world have been centered around carbon-based materials for

a number of centuries. There are various forms of carbon materials that are available.

14 It appears that the variety of carbon forms may be endless, and a simple classification

15 is given in Table 1.

Five decades ago, a concept based on the Hydrogen Economy has been evolving 16 and the transformation towards this economy essentially depends on the three compo-17 nents namely generation, storage, and delivery of hydrogen. There are consistent 18 attempts to generate hydrogen from the decomposition of water from renewable 19 sources) and this pursuit is an ongoing exercise to make this process economically 20 feasible, though the desired level of success is still to be achieved. Commonly avail-21 able methods for the storage of hydrogen, namely in high-pressure gas cylinders (up 22 to 800 bar) and as liquid hydrogen in cryogenic tanks (at 21 K) are energy inefficient 23 and not safe for certain applications, especially for mobile applications. Hydrogen 24

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Sp ³	Sp ²	Sp ¹
Diamond	Graphite	Carbyne
Cubic	Hexagonal	A-Carbyne
Hexagonal	Rhombohedral	B-Carbyne
C C		Chaoite
$Sp^3 + Sp^2 + Sp^1$	$Sp^n (3 < n > 1, n \neq 2)$	
Mixed forms of carbon	Intermediate forms of carbon	Intermediate forms of carbon
Amorphous carbon	3 > n > 2	2 > n > 1
Glassy carbon	Fullerene C _x	Cyclo (N) carbons
Carbon black	X = 60, 70, 84	N = 18, 24, 30
Adamantine carbon	(When $x = \infty$, $n = 2$)	(When $N = \infty$, $n = 1$)
	Carbon onions	
	Carbon nanotubes	

Table 1 A classification scheme for carbon allotropes, molecular crystals, and derived forms

adsorption/absorption on solid materials with high specific surface area and chem-25 ically bonded in covalent and ionic bonds appears to be attractive. Materials such 26 as metal hydrides, alloys, complex hydrides, and high surface area porous materials 27 are showing an affinity for ab(ad)sorbing large amounts of hydrogen. However, each 28 of these storage materials suffers from some particular drawbacks. These materials 29 crystalize in closed packed configurations giving rise to tetra-/octahedral- or other 30 voids thus possibly restricting one hydrogen species per void space at room temper-31 ature and atmospheric pressure, which means the weight percent of storage capacity 32 depends on the atomic or molecular weight of the nature of solid storage medium. 33 In the solid-state hydrogen storage, hydrogen is bonded by either physical forces 34 (Physical adsorption or van der Waals forces), e.g., MOF and carbon-based mate-35 rials, or chemical forces (chemisorption involving chemical bonds), e.g., hydrides, 36 imides, and nitrides. Physisorption has the advantages of higher energy efficiency 37 and faster adsorption/desorption cycles, whereas chemisorption results in the adsorp-38 tion of larger amounts of gas but in some cases, is not reversible and requires higher 39 temperatures to release the adsorbed gas which is not suitable for mobile applications. 40 The development of hydrogen storage materials with lightweight, high capacity, 41 high stability, and better safety is imperative for portable electronics and transport 42 applications. Carbon-based materials with different (allotropic) forms and chemical 43 affinity are favorable for hydrogen storage application due to their low atomic weight, 44 high surface area, porous nature, and higher safety characteristics. Carbon can be 45 hybridized in Sp, Sp², and Sp³ bonds, and exist in 0D, 1D, 2D, and 3D forms. Addi-46 tionally, chemical and physical properties of these materials can be altered by using its 47 different sources, composites, functional derivatives, or doping with other elements. 48 It may possess pores of different shapes and sizes, which are ideal physisorption/ 49 ad(ab)sorption sites for hydrogen storage. In the following, the hydrogen storage 50 capacities of different carbon nanostructures, including activated carbon, graphite, 51 graphene, CNTs, fullerene, and CNFs, are discussed. The attempts to store on modi-52 fied (especially metal loaded) carbon materials also will be considered. Recent and 53

relevant efforts for improving the hydrogen storage capacities are also considered. There are a variety of these materials and nature has shown that this is one of the potential storage media for hydrogen (in the form of hydrocarbons) and also shown that storage can go up to 25 weight percent (as in methane). However, one must be aware these hydrocarbon molecules are covalent in nature and thus may not readily release hydrogen when required as in transport applications. In this presentation, the scope of carbon materials for hydrogen storage application is considered.

The microporosity of carbon materials is not directly related to hydrogen storage 61 application but a nearly linear relationship has been realized with respect to BET 62 surface area, typical plots can be seen in the following reference and references 63 cited therein [1]. There have been many attempts to modify the surface of carbon 64 materials so that the storage capacity can be increased. One such attempt deals 65 with the presence of heteroatoms on the surface of carbon materials for various 66 reasons like the possibility of dissociating molecular hydrogen or facilitating the 67 spill-over process However, the current situation with respect to hydrogen storage 68 in carbon materials with heteroatoms can be stated that typically N-doping is only 69 apparent when considering the hydrogen uptake as a function of microporosity (rather 70 than total porosity). This possibly leads to the conclusion that pores larger than the 71 micropore size range have a lesser role in hydrogen storage capacity. 72

The advanced design and testing of carbon materials for energy storage devices 73 appear to be important. The main shortcomings of these materials are related to irre-74 versible capacity loss, big voltage crosstalk, and low density [2]. Novel composites 75 containing multifunctional nanostructured carbon and other dopants can synergisti-76 cally take advantage of the combination of ordered building block units with other 77 desired properties. Since most physisorption is involved in this process of hydrogen 78 storage, only a small amount of hydrogen could be stored even at a pressure of 90 bar. 79 Obviously, temperature (~room temperature) will have an almost negligible effect 80 on hydrogen storage capacity. 81

82 2 Activated Carbon

Activated carbon is one of the widespread and commercially used porous carbon 83 materials, consisting of graphite crystallites and amorphous carbon. It has been used 84 for centuries as an adsorbent due to its good adsorption capacity, high specific surface 85 area (up to ~3000 m²/g), porous structure, inertness, surface reactivity, and thermal 86 stability. The pores range from micro to macro size with an abundance of micro-87 sized pores of <1 nm. Due to its widespread pores, it is a potential material for 88 hydrogen storage. In general, the hydrogen adsorption on such materials is propor-89 tional to the specific surface area (SSA) and pore volume. Several research efforts are 90 devoted to finding the hydrogen uptake in activated carbon materials with different 91 pore sizes, pore volume, and specific surface area. Most of the experimental findings 92 have demonstrated the hydrogen storage capacity of activated carbon as 2.5 and 5.5 93 wt.% at low pressure (1-10 bar) and high pressure (up to 60 bar), respectively. It 94

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was observed that at 100 bar and room temperature, the hydrogen uptake of acti-95 vated carbon was <1 wt.%, even with a highly porous structure and high specific 96 surface area. It has been observed by different researchers that the optimum pore 97 size in activated carbon for hydrogen storage is 0.6 nm. Table 2 shows the experi-98 mental results reported complied) of hydrogen storage capacity of activated carbons 99 at different temperatures and pressure conditions. Different chemical treatments and 100 doping of activated carbon were also utilized to improve the hydrogen uptake of acti-101 vated carbon materials. However, they do not meet the U.S. Department of Energy 102 (DOE) target, and further efforts are needed to realize their potential for enhanced 103 hydrogen storage capacity. 104

One can assess that the highest value of excess adsorption of hydrogen reported 105 for a commercial activated carbon is around 5 wt.% at 77 K and pressures from 3 to 106 6 MPa. In the case of activated carbons prepared on a laboratory scale, the best value 107 obtained is ~7 wt.% at 77 K and 20 MPa. Most of the studies reported in the literature 108 mainly concentrated on increasing the surface area of the activated carbon (by an 109 appropriate method for (mostly chemical) activation) and the porosity of the material. 110 The linear correlations connecting adsorption capacity and physical parameters like 111 surface area and pore volume of activated carbons have been attempted but the results 112 available indicate only limited applicability. It is also to be noted that activated 113 carbons show hydrogen adsorption capacities at cryogenic conditions and this is not 114 very suitable for the applications envisaged for hydrogen economy. 115

Adsorbent employed	Reported hydrogen storage	Conditions (temperature and	
Ausorbent employed	(wt.%)	pressure)	
Activated carbon	0.67	303 K 10 MPa	
Activated carbon	1.4	77 K 0.1 MPa	
Activated carbon	1.6	298 K 13 MPa	
Activated carbon	4.5	77 K	
Activated carbon	5.7	77 K 3 MPa	
Activated carbon	3.2	298 K 20 MPa	
Activated carbon	6/5	298 K 50 MPa	
Chemically activated carbon	8.0	77 K 4 MPa	
Activated carbon	2.3–5.8	77 K 4 MPa	
Activated carbon (treated with KOH)	6.6	4 MPa	
Activated carbon (activated with Ni)	1.8	77 K 0.1 MPa	
Activated carbon (activated with Pt)	2.3	298 K 10 MPa	
Activated carbon (activated with Pd)	5.5	298 K 8 MPa	

Table 2 Data on hydrogen storage	pacities of different activated carbon materials (Data	a from
literature)		

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116 3 Graphite

Graphite has Sp² hybridized carbon atoms arranged in a hexagonal lattice with 117 a layered structure such that the layers themselves are held together by van der 118 Waals forces of interaction. Graphite can be naturally obtained or can be artifi-110 cially prepared. It is also widely explored for hydrogen storage applications. Various 120 studies have modified their porosity, chemical bonding, and surface area by using 121 different activation methods as well as doping them with different metals. Ball-milled 122 graphite and intercalated graphite are widely explored for hydrogen storage applica-123 tions. Ball-milled graphite contains tremendous defect sites and high surface area, 124 which lead to chemisorption resulting in high hydrogen storage capacity. The ball-125 milling-induced rupture of graphite sheets and the emerging active edges and defects 126 on the fragmented part are also confirmed by neutron scattering and spin resonance 127 studies. 128

4 Graphene: The Hydrogen Adsorption/Desorption Isotherm

The hydrogen adsorption/desorption isotherm of the nitrogen-doped graphene, 131 Graphitic Oxide and Graphite powder is shown in Ref. [3]. The hydrogen adsorption 132 isotherm has been carried out at 298 K and 90 bar pressure. The nitrogen-doped 133 graphene material showed nearly ~ 1.5 wt.% hydrogen storage capacity at room 134 temperature and 90 bar pressure. In this context, the graphitic oxide (GO) showed 135 ~ 0.21 wt.% hydrogen storage capacity at room temperature and 90 bar. This value 136 is less than that of nitrogen-doped graphene material. However, nitrogen doping of 137 graphene materials takes up substitution positions in the carbon lattice, there is trans-138 portation of hydrogen atoms on to the graphene surface. Furthermore, these results 139 reveal that the nitrogen doping on graphene materials can extensively modify the 140 catalytic effect of the graphene materials for hydrogen dissociative adsorption, fore-141 most for the improvement of the dissociative hydrogen adsorption. This observation 142 suggests that the nitrogen atoms possibly take part in a role in the hydrogen adsorption 143 capacity at room temperature. A previous study recommended that the presence of 144 nitrogen atoms in graphene sheets increases the enthalpy of hydrogen adsorption [4]. 145 Potentially almost all the adsorbed amount can be desorbed which is an interesting 146 characteristic expected for hydrogen storage materials. Various attempts have been 147 made to modify or adopt different preparation procedures for graphene materials and 148 it has been shown that N and P doping in graphene does not improve the materials' 149 hydrogen storage capacity [5]. It has been proposed that the hydrogen binding energy 150 can be tuned as a function of the graphene local curvature. The adsorption is facili-151 tated by convex sites and desorption is facilitated on concave sites. The corrugation 152 of graphene flakes and its control with large variety of functionalization may make 153 these materials can impact in the search for hydrogen storage [6]. 154

5 Modification of Activated Carbon

For conventional activated carbon materials, the hydrogen uptake is proportional to 156 the surface area and pore volume; and normally the data are fitted well with the 157 Langmuir isotherm model (monolayer adsorption). High adsorption capacity is only 158 obtained at extremely low cryogenic temperatures and high pressures. Hydrogen 159 adsorption on various types of commercial and modified activated carbon products 160 has been extensively studied. Experimental results show that products with micropore 161 volumes greater than 1 mL/g are able to store ca. 2.2 wt.% of hydrogen due to 162 physisorption and it is expected that optimization of the adsorbent and sorption 163 conditions could lead to a storage capacity of 4.5-5.3 wt.%. Agricultural waste 164 materials such as coconut shells, coconut fibers, jute fibers, nut shells, oil seeds, 165 etc., are popular raw materials for producing activated carbon materials. Carbon 166 materials and their activation have been extensively discussed in Ref. [7]. Jin et al. 167 [8] prepared activated carbons with different porosities using chemically activated 168 coconut shells. They reported a maximum hydrogen adsorption capacity of 0.85 169 wt.% at 100 bar and 298 K. Sharon et al. [9], produced activated carbon fibers (ACF) 170 using soybean and bagasse. The authors measured hydrogen storage capacities of 171 1.09–2.05 wt.% at a pressure of 11 Pa and room temperature. Another form of AC, 172 the advanced AC monoliths, with good mechanical strength (maximum compression 173 strength of 22 MPa), high volume of micropores (up to 1.04 cm³/g), and high density 174 (up to 0.7 g/cm³) have been shown to adsorb 29.7 g/L of hydrogen at 77 K and 175 4 MPa [9]. Mechanically milled AC consists of some form of defective nanostructure, 176 which increases the specific surface area. Research findings have revealed that after 177 10 h of milling, the hydrogen storage capacity increases from 0.90 wt.% to ca. 1.7 178 wt.%. Studies have shown that the loading of precious metals, e.g., Pt, onto AC, 170 increases the adsorption capacity. The merging of the two adsorption phenomena, 180 i.e. chemisorption (on the Pt surface) and physisorption (on the carbon surface) gives 181 rise to a significant amount of spillover hydrogen. 182

183 6 Carbon Nanotubes

Ever since the discovery of carbon nanotubes was reported in 1991, there have been various attempts to use this new type of carbon material for hydrogen storage. These studies have led to some unexpected levels of storage up to nearly 60 wt.% or even more. However, the consensus now is that these reports claiming over 60 wt.% are flawed by experimental aberrations.

Only limited data are given in Table 3. For more extensive compilations, one is directed to references [35]. Various nanotubes like carbon nanotubes, boron nitride nanotubes, silicon carbide nanotubes, carbon nano-scrolls, pillared Graphene, and porous nanotube network materials have been extensively investigated and the final suggestion is that one should design novel materials with the following key 105

parameters namely high accessible surface area, large free pore volume and strong 194 interactions between the surface-active sites and the substrate hydrogen.

In a recent review, Lyu et al. [36] propose that "a detailed study of the optimum 196 number of metal atoms without aggregation on CNT should be performed. (1) At 197 the same time suitable preparation methods for realizing controllable doping sites 198 and doped configurations should be devised; (2) The material synthesis, purification, 199 and activation methods have to be optimized; (3) Active sites, molecular configura-200 tions, effectively accessible surface area, pore size, surface topology, the chemical 201 composition of the surface, applied pressure and temperature, defects, and dopant, 202

Sample	Temp (K)	P (MPa)	Hydrogen storage (wt.%)	Refs.
Herring bone GNFs	RT	11–35	67.5	[10]
Platelet GNFs	RT	11–35	53.68	[10]
Graphitic nano fibers	RT	101	10	[11]
Graphitic nano fibers	RT	8-120	10	[12]
SWNTs (low purity)	273	0.4	5-10	[13]
SWNTs (high purity)	80	70–80	8.25	[14]
SWNT (high purity + Ti alloy)	300-600	0.7	3.5–4.5	[15]
Li-MWNTs	473–673	1	20	[16]
Li-MWNTs (K0MWNTs)	473–673	1	2.5 (1.8)	[17]
MWNTs	RT	Ele-	<1	[18]
SWNTs	300-520	1–100	0.1	[19]
Various CN	RT	1	<0.1	[20]
SWNTs (+ Ti alloy)	RT	35	0	[21]
SWCNT	RT	0/8	4.5	[22]
SWCNT	295	10	0.93	[23]
SWCNT	RT	0.1	1.2	[24]
SWCNT	323	4.8	4.77	[25]
SWCNT	-	-	-0.8	[26]
MWCNT	-	3	2	[27]
MWCNT	77	0.005	0.54	[28]
MWCNT	298	0.1	0.2	[29]
NWCNT	425	10	3.8	[30]
MWCNT	143	3	3.5	[31]
MWCNT	RT	7.5	1.5–2.1	[32]
MWCNT	-	-	2.7–3.8	[31]
Carbon nano onion	77	1.5	>18.2	[33]
g-C ₃ N ₄ nanotubes	RT	37	0.78	[34]

 Table 3
 Selected data (for comparison) on the storage of hydrogen by carbon nanotubes

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which are some of the important factors that strongly affect the hydrogen absorption
 in carbon nanotubes" should be elucidated.

In contrast, Lobo et al. [37], propose carbon nanostructures are promising mate-205 rials for hydrogen storage applications. They emphasize that hydrogen can be 206 physisorbed in carbon nanotube bundles on various sites such as external wall 207 surfaces, grooves, and interstitial channels. Therefore, it can have a large energy 208 density (as required for mobile applications). It is also known that by tuning the 209 adsorption conditions, hydrogen can be either chemisorbed or physisorbed in carbon 210 nanotubes. In a review, Lee et al. [38] and others [39] propose that a more detailed 211 understanding of the interfacial interactions between adsorbent and adsorbate should 212 be evolved and the phenomenon of spill-over can contribute to adsorbent surfaces to 213 achieve the desired levels of hydrogen storage. 214

7 Impact of Structures on Hydrogen Storage in Different Carbon Materials

Increasing hydrogen storage to the levels advocated by DOE through materials in 217 the nanoscale has been proposed and intensively researched but the success appears 218 to elude us. Among the several options available, various allotropic forms of carbon, 219 like CNTs, graphene, and activated carbons have been considered to be appropriate 220 systems owing to their unique properties like high surface area, (this parameter has 221 limited linear variation with absorbed hydrogen) porous nature (but pore volume is 222 limited for each material) and high thermal and mechanical stability, and all these 223 have been so far considered though questionable? As vital factors for hydrogen 224 storage. Emphasis has been given to the hollow and porous structure of CNTs which 225 is supposed to give the possibility of hydrogen storage both in the inner and outer 226 surfaces that are well investigated by many theoretical as well as experimental studies. 227 These studies though have relevance did not answer the stumbling block in achieving 228 the goal of reaching the DOE standards. CNTs are termed to be one of the promising 220 nanostructures for hydrogen storage however this expectation also has not yielded 230 the desired result. 231

Moreover, the simultaneous presence of defects, both topological and structural 232 types, and other irregularities affect the activation and adsorption and hence the 233 amount of hydrogen stored in the system. In addition to this, doping also leads to 234 structural and electronic property variations. However, these modifications so far 235 have not shown enormous improvement in hydrogen storage capacity. It is however 236 believed that estimated values of hydrogen storage still show the high impact of 237 structure on the storage capacity. Nanomaterials are also analyzed and found that 238 some of these strongly affect the storage capacity. But still, the DOE target could not 239 be achieved which makes this issue an interesting topic of study even today. 240

Different types of nanotubes like metallic and semiconductor (also single or multi-241 walled) with varying structures were considered. It has been observed that the adsorp-242 tion binding energy values as a function of different orientations of adsorbed species 243 and at different sites of nanostructures are compared and the results show the strong 244 impact of CNT structure on the storage capacity of hydrogen. Hydrogen can be 245 adsorbed on graphene in two different ways: physisorption or chemisorption. While 246 the first one is due to Van Der Waals interaction, the second is by forming a chemical 247 bond with the C atoms. Physisorption usually happens with hydrogen in molecular 248 form and in chemisorption, dissociation of H2 into atomic hydrogen takes place and is 249 a rather favorable process unlike in CNTs where physisorption is the most preferred 250 way of storage. A single graphene layer is a quasi-2D system, and its VD is not well 251 defined, thus in the evaluation of the potentialities of graphene for hydrogen storage, 252 different forms of graphene such as multilayers, three-dimensional assemblies, or 253 nanostructures of graphene are considered. However, it should be remarked that the 254 standards for the necessary hydrogen storage for the projected applications should 255 be realized at the earliest. 256

257 8 Modified Carbon Materials for Hydrogen Storage

Modification of the equipotential surface of carbon materials has been attempted 258 for various reasons. Usually, the activation is aimed at introducing active centers on 259 supported phases. It is usually to disperse active metals so that the exposed metal 260 surface area is maximum. Hydrogen adsorption involves dissociation of molecular 261 hydrogen and any storage medium should have dissociating centers for molecular 262 hydrogen. Recognizing this aspect, heteroatom substitution has been proposed as one 263 of the options. DFT calculations have also shown that molecular hydrogen dissocia-264 tion energy is considerably reduced on heteroatoms like N, P, or B substituted carbon 265 surfaces (Table 5), however, these studies so far have not led to the levels of storage 266 of hydrogen on these materials. 267

268 8.1 Nitrogen Doping in Carbon Materials

Nitrogen is an abundant (80%) element in the terrestrial atmosphere. Molecular 269 nitrogen is stable and has a minor role in the lower atmosphere. N-doped carbon 270 materials are the much-studied area in energy storage. Depending on the type of 271 N bonding within the carbon matrix, nitrogen can share one to two π -electrons 272 with the π -electron system of the carbons. This sharing of electrons causes an *n*-273 type doping if N atoms directly substitute the C atoms in the graphitic lattice. N-274 doping in carbon generally manifests itself in three different forms namely, pyrrolic-275 N, pyridinic-N, and quaternary-N, and each form alters the carbon electronic band 276

gap differently. The band gaps of pyrrolic-N, pyridinic-N, and quaternary-N-doped
carbons are reported to be 1.20, 1.40, and 1.39 eV, respectively (Table 4).

Particularly, all materials obtained by these techniques have a nitrogen content
lower than 10 atoms % because of the high temperature environment that was used
in carbonization. An alternate synthesis method needs to be developed that enables
the preparation of carbon material with high nitrogen content and time they should
be stable at high temperatures. It will be beneficial if one can adopt a polymerization
and low-temperature growth process to create nitrogen-doped carbon materials.

Nitrogen is essentially introduced into the carbon matrix in two ways, either
by the carbonization of N-containing precursors or by post-modification methods.
The common nitrogen-containing precursors are urea, melamine, cyanide, polyacrylonitrile, and ammonia. Another resourceful approach for the synthesis of N-doped
carbon materials is based on naturally (sustainable) nitrogen-containing precursors

Substitution	Total energy (Hartrees)	Bond length H_1-H_2 (Å)	Dissociation energy (eV)
Hydrogen	-1.175	0.708	4.76
CNT	-3686.5502	0.776	
CNT + H ₂	-3687.7161		4/51
NCNT	-3702.5908		
NCNT + H ₂	-3703.5989	0.815	0.22
PCNT	-3989.1694		
PCNT + H ₂	-3990.2550	0/815	2.33
SCNT	-4046.0020		
SCNT + H ₂	-3047.0067	0.817	0.13
BCNT	-3671.7254		
BCNT + H ₂	-3672.9440	0.818	5.95
2BCNT (adjacent)	-3658.6666		
2BCNT (adjacent) + H ₂	-3659.8092	0.813	3.88
2BCNT (Alternate) 2BCNT (alternate) + H ₂	-3659.3491 -3660.3594	0.928	0.28

 Table 4
 Bond length and dissociation energy of hydrogen on the CNTs calculated using B3LYP with 6.31G (P,D) basis set on the UFF optimized structures [40]

Table 5 Technical system targets for on-board hydrogen storage for light-duty fuel cell vehicles

	2020	2025	Ultimate
Usable specific energy from H ₂ [kWh/kg]	1.5	1.8	2.2
Net usable energy/mass system mass [kg H ₂ /kg system]	0.045	0.055	0.065
Usable energy density from H ₂ [kWh/L]	1.0	1.3	1.7
Net usable energy/max system volume [kgH2/L system]	0.030	0.040	0.050
System cost [USD/kWh net]	10	9	8

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like amino-carbohydrates or other N-enriched polymers, amino acids, proteins, N-200 ionic liquids, and waste crab shells. Another possible way is to pyrolyze the nitrogen 201 and carbon-containing precursors, such as heterocycles or melamine, by which direct 292 incorporation of nitrogen atoms into the forming of carbon backbone becomes 293 possible. One of the procedures for deriving N-doped carbon is hydrothermal 201 treatment of carbohydrate-rich biomass. Using nitrogen-containing biomass-related 205 precursors and hydrothermally treating them yields nitrogen-containing carbona-296 ceous materials that offer tremendous possibilities for further treatments and energy 297 applications. The application of nitrogen-doped carbon materials has been investi-298 gated as a material for hydrogen storage at room temperature and ambient pressure 299 [41]. It has been reported that nitrogen-enriched graphitic carbon material exhibits 300 a hydrogen storage capacity of 0.34 wt.% at 298 K under 100 bars [42]. It has also 301 been reported that the addition of N-species in mesoporous carbons showed hydrogen 302 adsorption capacity of 1.1 wt.% at 298 K and 100 bar pressure [43]. It has also been 303 reported that the microwave plasma CVD process enables the growth of specific 304 nanostructured nitrogen-doped carbons. Nitrogen incorporation into these forms of 305 carbon is approximately 1 at.%. It shows a gravimetric hydrogen storage capacity of 306 0.7–0.8 wt.% under 300 K and 0.1–7 MPa [44]. Hydrogen adsorption on nitrogen-307 doped carbon xerogels showed maximum hydrogen uptake of 0.28 wt.% at 308 K 308 [45]. 300

8.2 Phosphorus-Doping in Carbon Materials

The changes in the physical properties of Sp² carbon motifs after the addition of 311 phosphorous into their lattice are considered. However, P has a larger atomic radius 312 and higher electron-donating ability, which makes it an option as a dopant. Phos-313 phorus is not a common element in carbons, although it is present in carbons obtained 314 using phosphoric acid activation. Due to the addition of P in the carbon matrix, the 315 density of states near the Fermi level is also found to increase, which increases with 316 the increase in the P-doping level. In these reactions, the formation of phosphate 317 and polyphosphate bridges provokes the expansion and cross-linking of the carbon 318 matrix, driving to an accessible pore structure after the removal of the acid. The 319 chemical state of phosphorus in carbons is a rather controversial issue. Some exper-320 imental evidence using different analytical techniques (FTIR and XPS) has shown 321 that the most abundant P species introduced in carbons by phosphoric acid activa-322 tion are -C-P- or -C-O-P bonds in phosphate and phosphonate-like structures. The 323 XPS analysis further allowed for an insight into the binding states, proving the true 324 incorporation of the phosphorus atoms into the graphite sheets, besides some P-O 325 binding sites, most likely on the surface of the material. The existence of pentava-326 lent phosphorus and elemental phosphorus is very infrequently detected, except when 327 high temperatures are applied. Recent studies have reported that P-containing groups 328 might be significant for the progress of graphitic crystallites which contrasts with far 329 reported role of P as an inhibitor of carbon graphitization. Yang et al. [46] reported that 330

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phosphorous-doped ordered mesoporous carbon was synthesized by co-pyrolyzing a 331 phosphorus-containing source and a carbon source collectively using ordered meso-332 porous silica (SBA-15) as a template without the use of any metal components. More 333 recent approaches have recognized dissimilar synthetic pathways in the direction of 334 phosphorus-doped carbon materials, proving themselves as promising candidates for 335 energy storage applications. It has been shown in the previous section that carbon 336 materials with substitution by heteroatoms like N, S, and B, show hydrogen sorption 337 capacity. However, the effect of substitution of phosphorus in carbon materials has 338 not been investigated to the same extent. 339

340 8.3 Boron-Doping in Carbon Materials

Boron is an element with unique properties. It is thus an interesting candidate 341 for doping of carbon materials, modifying the properties of pure carbons. Several 342 researchers have started focusing not only on basic studies on B-doping, but also on 343 applying the obtained materials and exploiting their favorable properties in energy-344 related applications. Due to its three valence electrons, B is well thought-out as a 345 good dopant. Substitutional boron enhances the graphitization of carbon. It has been 346 found that boron atoms are favored to be substituted in the graphite lattice. The 347 existence of B-C bonds in the carbon framework can lower the Fermi level of the 348 structure and then tune the properties of oxygen chemisorption and electrochemical 349 redox reactions. The synthetic procedure, in which elemental B and graphite powder 350 served as precursors, yielded a mixture of different B-containing carbon nanostruc-351 tures, such as thin graphitic sheets, tubes, and filaments. The substitutional doping 352 of carbon atoms in Sp² and Sp³ configurations with boron can modify the elec-353 tronic and structural properties of the resulting carbon. Over the years boron-doped 354 carbons have been synthesized by standard CVD process using BCl₃. Substituted 355 boron atoms in the carbon lattice accelerate the graphitization and suppress the 356 oxidation of carbon materials, which seems promising for their use as reinforce-357 ment materials in aerospace applications. The positive effect of boron doping on 358 diamond and carbon electrodes and in the field of hydrogen storage has also been 359 reported, although further optimization of the boron doping environment seems yet 360 to be needed. 361

The storage of hydrogen in carbon nanomaterials requires appropriate chemical 362 activators in suitable geometry. Sankaran et al. reported different types of carbon 363 materials employed for the hydrogen sorption capacity. The storage capacity of 364 2 wt.% at 298 K and 80 bar pressure is obtained for boron boron-substituted carbon 365 nanotube. However, a maximum storage capacity of 2 wt.% is attained at 80 bar 366 and 300 K for boron-containing carbon nanotubes (BCNT) whereas pure carbon 367 nanotubes (CNT) show only 0.6 wt.% at 300 K and 80 bar and B-doped bulk 368 carbon material (PBC) shows only 0.2 wt.%. Mike Chung et al., reported the microp-369 orous boron-substituted carbon (B/C) materials show a significantly higher hydrogen 370 binding energy and reversible hydrogen physisorption capacity of 0.6 and 3.2 wt.% 371

at 293 and 77 K, under 40 bar of hydrogen pressure [47]. B-containing polymeric 372 precursors and pyrolysis were employed to synthesize microporous B/C materials 373 with a high B content (7.2%) and high surface area (780 m^2/g). The substitutional 374 B elements in B/C material serve as internal p-type dopants and polarize the C 375 surface, which exhibits a significantly higher hydrogen binding energy [48]. For 376 efficient hydrogenation and hydrogen storage, these boron atoms should be incor-377 porated geometrically and chemically into the carbon network. Wang et al. reported 378 that B- and N-doped microporous carbon had a hydrogen storage capacity of 0.55 379 wt.% at 298 K and 10 MPa. By doping 6.0 wt.% Ru metal on the B- and N-doped 380 microporous carbon, the hydrogen uptake at 10 MPa was increased to 1.2 wt.%, i.e. 381 The improvement of hydrogen storage was due to the spillover of atomic hydrogen 382 from the Ru metal particles to the B-and N-doped microporous carbon [49]. 383

384 9 Perspectives

³⁸⁵ The standards that one wishes to achieve in the storage of hydrogen are: [54–56].

- Gravimetric H density in the range of 5–10 wt.% H₂, and energy density of 1.6–3.2 kWh/kg.
- Volumetric H density >50 kg H_2 m⁻³ and energy density >1.6kWh/L.
- Thermodynamics: T 0 < 85 °C (transport applications) or <200 °C (stationary applications).
- Kinetics (tank level): fill time 3-5 min; H₂ release flow 1.6 g/s.
- Durability: 1500 cycles (1/4 tank to full).

One of the recent specifications and the time to achieve them are assembled in Table 5.

Of all the available hydrogen storage materials, why carbon materials are preferred 395 option? What is the maximum hydrogen storage capacity that can be expected and 396 what will be the limit that can be practically achieved? It may be remembered that 397 nature mostly provides hydrogen source in combined form with carbon and oxygen 398 though other elemental compositions are also possible. If carbon materials can be 399 obtained in an atomic state, then the maximum storage capacity can be expected to 400 be around 25 wt.%. However, since it is not possible to get atomic hydrogen, the 401 carbon materials can be obtained at the limit with one vacant valency in carbon two-402 dimensional material and the maximum storage can be expected to be 6.25 wt.%. 403 This limit is arrived at assuming that hydrogen is held by the solid by valence forces. 404 If hydrogen is stored or retained by other forces, this limit may not hold good. 405

If the stored hydrogen were to occupy the interstitial sites in carbon materials,
then the energetics of storing and releasing should also be considered for practical
application.

Since normally carbon materials are microporous in nature, hydrogen may be held
 in these pores by condensation forces and hence one can hope for higher storage

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capacity, however, the experimental variables for this process namely temperatureand pressure have to be different from normal ambient conditions.

413 **10** Conclusion

Hydrogen storage characteristics and treatments for improving the storage capacities
 of different carbon materials are presented. The following deductions can be stated:

- The hydrogen storage capacity of carbon materials depends upon surface area,
 which is affected by micropore size distribution that counts for the presence of
 narrow micropores.
- 2. Thermal treatments and metal doping on carbon nanostructures are observed to be useful for improving hydrogen storage capacities but higher storage capacities
 can be obtained at cryogenic temperature and higher pressure. Further investigations on modified carbon nanostructures may be useful to achieve the target of 7.5 wt.% for automobile applications.
- Theoretical studies show a strong structural dependence of carbon materials on
 hydrogen adsorption capacity, especially in CNTs. The defects influence the
 adsorption capacity.
- 427 4. Several theoretical studies predicted amazing storage capacities but were not 428 confirmed experimentally. Also, higher storage capacities like 20 wt.% for metal 429 doped MWCNT and 10 wt.% for CNF measured experimentally are reported in 430 the literature but are to be confirmed.
- 431 5. Experimental and theoretical studies on adsorption storage reactors are scarce
 432 in the literature. Studies on the design, material, and thermophysical properties
 433 of reactors can be done for isotherm measurements, especially for automotive
 434 applications. These vital issues will be a good technical contribution in the field
 435 of hydrogen-adsorption systems for onboard applications.

Spill-over is one of the phenomena invoked by the transport of hydrogen from the
site of impact to other normally inactive sites. This phenomenon has been investigated
and is commonly accepted alternate way of surface transport. This transport requires a
transporting medium and in the case of hydrogen, it is usually water and its fragments.
In the case of carbon, this transporting medium can be either adsorbed water species
or the carbon species themselves. The role of spillover in hydrogen sorption and
storage needs more information.

443 **References**

- Panella, B., Hirscher, M., Roth, S.: Hydrogen adsorption in different carbon nanostructures.
 Carbon 43, 2209 (2005)
- Zhu, Z.H., Hatori, H., Wang, S.B., Lu, G.Q.: New insights into the interaction of hydrogen atoms with boron substituted carbon. J. Phys. Chem. B 109, 16744 (2005)

- Arjunan, A., Viswanathan, B., Nandhakumar, V.: Nitrogen doped graphene as potential material
 for hydrogen storage. Graphene 6, 41 (2017)
- 4. Wang, L., Lee, K., Sun, Y.-Y., Lucking, M., Chen, Z., Zhao, J.J., Zhang, S.B.: Graphene oxide as an indeal substrate for hydrogen storage. ACS Nano 3, 2995 (2009)
- Ariharan, A., Viswanathan, B., Nandhakumar, V.: Heteroatom doped multi-layered graphene
 material for hydrogen storage application. Graphene 5, 39 (2016)
- Tozzini, V., Pellegrini, V.: Prospects for hydrogen storage in Graphene. Phys. Chem. Chem.
 Phys. 15, 80 (2013)
- Viswanathan, B., Indraneel, P., Varadarajan, T.K.: A process for the preparation of activated
 carbon from botanical sources. Catal. Surv. Asia 13, 164 (2008)
- Jin, H., Lee, Y.S., Hong, I.: Hydrogen adsorption characteristics of activated carbon. Catal.
 Today 120, 399 (2007)
- Jorda-Beneyto, M., Lozano-Castello, D., Suarez-Garcia, F., Cazorla-Amoros, D., Linares-Solano, A.: Advanced activated carbon monoliths and activated carbons for hydrogen storage.
 Microporous Mesoporous Mater. 112, 235 (2008)
- Chambers, A., Park, C., Baker, R.T.K., Rodriguez, N.M.: Computer simulations of hydrogen
 adsorption on graphite nanofibers. J. Phys. Chem. B 102, 4253 (1998)
- Fan, Y.Y., Liao, B., Liu, M., Wei, Y.L., Lu, M.Q., Chang, H.M.: Hydrogenuptake in vapor
 grown carbon nanofibers. Carbon 37, 1649 (1999)
- 467 12. Gupta, B.K., Srivastava, O.N.: Synthesi and hydrogenation behaviour of graphitic nanofibers.
 468 Int. I Int. J. Hydrog. Energy 25, 825 (2000)
- 13. Dillon, A.C., Jones, K.H., Bekkedahl, T.A., Klang, C.H., Bethume, D.S., Eben, H.J.: Storage of hydrogen in single-walled carbon nanotubes. Nature 386, 377 (1997)
- 471 14. Ye, Y., Ahvi, C.C., Witham, C., Fultz, B., Liu, J., Rinzler, A.G., Colbert, D., Smith, K.A.,
 472 Smalley, R.E.: Hydrogen adsorption and cohesive energy of single-walled carbon nanotubes.
 473 Appl. Phys. Lett. **74**, 2307 (1999)
- 474 15. Dillon, A.C., Genneth, T., Jones, K.M., Alleman, J.A., Parilla, P.A., Heben, H.J.: A simple and
 475 complete purification of single-walled carbon nanotube (SWNT) materials. Adv. Mater. 11,
 476 1354 (1999)
- theorem 16. Chen, P., Wu, A., Lim, J., Tan, K.L.: High H₂ uptake by alkali doped carbon nanotubes under ambient pressure and moderate temperatures. Science 265, 91 (1999)
- 479 17. Yang, R.T.: Hydrogen storage by alkali-doped carbon nanotubes-revisited. Carbon 38, 623
 (2000)
- 18. Elzbeta Fracko Wiak and Francois Bequin: Electrochemical storage of energy in carbon
 nanotubes and nanostructured carbons. Carbon 40, 1775 (2002)
- Hirscher, M., Bechner, M., Haluska, M., Quintel, A., Skakalova, V., Choi, Y.-M., Dettlaf Weglikowska, U., Roth, S., Stepanek, I., Bernier, P., Leonhardt, A., Fink, J.: Hydrogen storage
 in carbon nanostructures. J. Alloy. Compd. 330–332, 654 (2002)
- 486 20. Tirbetti, G.G., Heisner, G.P., Olk, C.H.: Hydrogen storage capacity of carbon nanotubes,
 filaments, and vapor-grown fibers. Carbon 39, 2291 (2001)
- 488 21. Hirscher, M., Becher, M., Haluska, M., Dettlaff-Weglikowska, U., Quintel, A., Duesberg, G.S.,
 489 Choi, Y.M., Downes, P., Hulman, M., Roth, S., Stepanek, I.: Hydrogen storage in sonicated
 490 carbon materials. Appl Phys A 72, 129 (2001)
- 22. Chambers, A., Park, C., Baker, R.T.K., Rodriguez, N.M.: Hydrogen storage in graphite
 nanofibers, J. Phys. Chem. B 102(22), 4253–4256 (1998)
- Assance and the second state of t
- Smith, M.R., Bittner, E.W., Shi, W., Johnson, J.K., Bockrath, B.C.: Chemical activation of single-walled carbon nanotubes for hydrogen adsorption. J. Phys. Chem. B 107, 3752–3760 (2003)
- 499 25. Silambasaran, D., Surya, V.J., Vasu, V., Iyakutti, K.: Experimental investigation of Hydrogen
- storage in single walled carbon nanotubes functionalized with borane. Int. J. Hydrog. Energy
 36, 3574–3579 (2011)

600834_1_En_15_Chapter 🗸 TYPESET 🔄 DISK 🔄 LE 🗹 CP Disp.:6/1/2024 Pages: 18 Layout: T1-Standard

- 26. Rashidi, A.M., Nouralishahi, A., Khodadadi, A.A., Mortazavi, Y., Karimi, A., Kashefi, K.: 502 Modification of single wall carbon nanotubes (SWNT) for hydrogen storage. Int. Hydrog. 503 504 Energy 35, 9489–9495 (2010)
- 27. Mosquera, E., Diaz-Droguett, D.E., Carvajal, N., Roble, M., Morel, M., Espinoza, R.: Charac-505 terization and hydrogen storage in multi-walled carbon nanotubes grown by aerosol assisted 506 CVD method. Diam. Relat. Mater. 43, 66-71 (2014) 507
- 28. Lee, S., Park, S.: Influence of the pore size in multi-walled carbon nanotubes on the hydrogen 508 storage behaviors. J. Solid-State Chem. 194, 307-312 (2012) 509
- 29. Barghi, S.H., Tsotsis, T.T., Sahimi, M.: Chemisorption, physisorption and hysteresis during 510 hydrogen storage in carbon nanotubes. Int. J. Hydrog. Energy 39, 1390-1397 (2014) 511
- 30. Lin, K., Mai, Y., Li, S., Shu, C., Wang, C.: Characterization and hydrogen storage of surface 512 modified multiwalled carbon nanotubes for fuel cell application. J. Nanomater. 1-12 (2012) 513
- 31. Rakhi, R.B., Sethupathi, K., Ramaprabhu, S.: Synthesis and hydrogen storage properties of 514 carbon nanotubes. Int. J. Hydrog. Energy 33, 381-386 (2008) 515
- 32. Rostami, S., Pour, A.N., Izadyar, M.: A review on modified carbon materials as promising 516 agents for hydrogen storage. Sci. Prog. 101, 171 (2018) 517
- 33. Sahu, S., Khan, M.S., Gupta, N., Chennakesavulu, K., Sasikumar, C.: The hydrogen storage 518 capacity of carbon nano-onions fabricated by thermal chemical vapour. Int. J. Hydrog. Energy 510 48 (2023). https://doi.org/10.1016/j.ijhydene.2023.03.156 520
- 34. Guo, R., Teng, Y.-S., Retita, I., Bakmanrokj, G., Arhurst, N., Chan, S.L.I.: A detailed exper-521 imental comparison on the hydrogen storage ability of different forms of graphitic carbon 522 nitride bulk, nanotubes and sheets with multiwalled carbon nanotubes. Mater. Today Chem. 523 30, 101508 (2023) 524
- 35. Lyu, J., Kudiiarov, V., Lider, A.: An overview of the recent progress in modifications of carbon 525 nanotubes for hydrogen adsorption. Nanomaterials, 10, 255 (2020) 526
- 36. Lyu, J., Kudiiarov, V., Lider, A.: An overview of the recent progress in modifications of carbon 527 nanotubes for hydrogen adsorption. Nanomaterials 10, 255 (2020) 528
- 37. Lobo, R., Ribeiro, J., Inok, F.: Hydrogen uptake and release in carbon nanotube electrocatalysts. 529 Nanomaterials 11, 97 (2021) 530
- 38. Lee, S.Y., Lee, J.H., Kim, Y.H., Kim, J.W., Lee, K.J., Park, S.J.: Recent progress using solid-state 531 materials for hydrogen storage: a short review. Processes 10, 304 (2022) 532
- 39. Lim, K.L., Kazemian, H., Yaakob, Z., Daud, W.W.: Solid-state materials and methods for 533 hydrogen storage: a critical review. Chem. Eng. Technol. 33(2), 213-226 (2010) 534
- 40. Sankaran, M.: On the potential of carbon materials for solid state hydrogen storage. Ph.D. thesis, IIT Madras (2007) 536
- 41. Yao, Y., Zhang, B., Shi, J., Yang, Q.: Preparation of Nitrogen-doped carbon nanotubes with 537 different morphologies from melamine-formaldehyde resin. ACS Appl. Mater. Interfaces 7, 538 7413-7420 (2015) 539
- 42. Yang, S.J., Cho, J.H., Oh, G.H., Nahm, K.S., Park, C.R.: Easy synthesis of highly nitrogen 540 enrichedgraphitic carbon with high hydrogen storage capacity at room temperature. Carbon 47, 541 542 1585-1591 (2009)
- 43. Cai, J., Bennici, S., Shen, J., Auroux, A.: The acid-bas properties of nitrogen-containing carbon 543 materials. Mater. Chem. Phys. 161, 142-152 (2015) 544
- 44. Badzian, A., Badzian, T., Breval, E., Piotrowski, A.: Nanostructured, nitrogen-doped carabon 545 materials for hydrogen storage. Thin Solid Films 398, 170–174 (2001) 546
- 45. Kang, K.Y., Lee, B.I., Lee, J.S.: Hydrogen adsorption on nitrogen-doped carbon xerogels. 547 Carbon 47, 1171–1180 (2009) 548
- 46. Yang, D.-S., Bhattacharjya, D., Song, M.Y., Yu, J.-S.: High efficient metal-free phosphorus-549 doped platelet ordered mesoporous carbon for electrocatalytic oxygen reduction. Carbon 67, 550 551 736–743 (2014)
- 47. Chung, T.M., Jeong, Y., Chen, Q., Kleinhammes, A., Wu, Y.: Synthesis of microporous boron-552 substituted carbon (B/C) materials using polymeric precursors for hydrogen physisorption. J. 553 Am. Chem. Soc. 130, 6668-6669 (2018) 554

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- Chung, T.M., Jeong, Y., Kleinhammes, A., Wu, Y.: Synthesis of micro-porous boron substituted carbon (BC) materials using polymeric precursors for hydrogen physisorption. ECS Trans. 19, 57–66 (2009)
- 49. Wang, L., Yang, F.H., Yang, R.T.: Hydrogen storage properties of B- and N-doped microporous carbon. AIChE J. 55, 1823–1833 (2009)

600834_1_En_15_Chapter 🗸 TYPESET 🔄 DISK 🔤 LE 🗸 CP Disp.:6/1/2024 Pages: 18 Layout: T1-Standard

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