# **Rapid Water Transportation through One-dimensional Carbon** Nanospaces

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ABSTRACT : Water transportation through one-dimensional (1D) and quasi-1D internal nanospaces w by the rates of water vapor adsorption in the internal nanospaces of highly pure single\_ and double-wall carbon nanotubes (SWCNTs and DWCNTs), respectively, was directly investigated from the experimentally measured rates of water vapor adsorption in these nanospaces as well as those determined by molecular dynamics (MD) simulations that have quasi 1D and 1D nanospaces, respectively. Water iswas found to be rapidly transported more rapidly through 1D nanospaces rather than through quasi-1D nanospaces. Molecular dynamic simulation of water through carbon nanotubes shows the same tendency as the experimental adsorption rates. This faster transportation through 1D nanospaces is attributed to the formation of In 1D system, fewer water molecules rarely form hydrogen bonds bonding between water molecules of water adsorbed in a 1D system. The Therefore, the rapid water transportation is a result of less hydrogen bonding formation of water adsorbed in 1D anospaces.adsorption rates determined through the MD simulations were in agreement with the experimentally determined rates.

Water is one of the most ubiquitous naturally occurring chemical substances and well known to havehas anomalous physical properties owingdue to its constituent hydrogen bonds onding. Hence, the anomalous properties of water have been <u>continuously</u> studied <u>extensively</u>.<sup>1</sup> Water transportation through a water channel in a biomembrane is another important research topicissue, because a water channel is involved in many physiological processes. Water channels have narrow hydrophobic nanospaces.-2 As these channels are flexible and complex structures, a nanoporous material having a model structure for a water channel is required in order to understand water mechanism via a water channel. <u>A Ccarbon nanotube</u> (CNT) is believed to be an example of one such material an el, because it has having hydrophobic one-dimensional (1D) nanospaces -3 Thus, the behavior of water properties in <u>CNTsearbon nanotubes arehas</u> recently been studied actively studied recently <sup>4</sup> For xample, The water permeation through membranes fabricated byusing CNTscarbon nanotube ishas been demonstrated towith the aim of understanding study the working of a water channel HoweverNevertheless, the mechanism of water transportation through the internal nanospaces of CNTs-carbon nanotubes remains to be elucidatedis still not clear.

Ideal-Model single\_ and double-walled CNTsearbon nanotubes (SWCNTs and DWCNTs)-were synthesized by Hata et al. and Endo et al., respectively<sub>5</sub>, those havewere found to have extremely low metal catalysts, <u>long-high</u> aspect ratios, and hydrophobic internal nanospaces.<sup>-5</sup> As mentioned above, they have high potential for being used to conduct These carbon nanotubes have the great advantage of-fundamental <u>studies</u> study-on water transportation through-the their internal nanospaces. Therefore, we employedHere we directly show water vapor transportation through these <u>SWCNTs</u> and <u>DWCNTsearbon</u> nanotubes to investigate water transportation through their 1D nanospaces fromby the adsorption rates of water vapor-adsorption in <u>SWCNTs</u> and DWCNT, and molecular dynamics (MD) simulations.

The SWCNTs and DWCNTs were observed by Hhighresolution transmission electron microscopy (HRTEM) vere performed on SWCNT and DWCNT using JEM-2100F (JEOL Co.) at 120 kV. Adsorption rates of water vapor were measured at 303 K every 50 ms using a homemadecustom-designed volumetric apparatus. these These CNTscarbon nanotubes arewere heated at a pressure of less than < 0.1 Pa for 2 h prior to the adsorption rate measurements. MD simulations with the leapfrog time integration scheme were performed for understanding the water transportation through carbon nanotubes mechanism; the simulation model comprised a single SWCNT having an infinite length and diameters of 1 or 2 nm, placed in a unit cell of dimensions  $4 \times 4 \times 6$  nm<sup>3</sup>. The interaction potentials between water molecules and between a water molecule and a carbon wall were calculated-by using TIP5P and Lennard-Jones potential models, respectively. Here, tThe TIP5P potential model is a -model that combines of the e dispersion and electrostatic interactions, and it was proposed by Ma <u>al -</u> A complete MD simulation (100 ps) was performed conducted atclose to ~303 K usingwith an integration time of 0.1 fs (see The -Supporting Information- for details of the simulation procedure)

Figure 1 shows the HRTEM images of the\_DWCNT and SWCNT, with The average internal diameters of DWCNT and SWCNT are 1 and 2–3 nm, respectively, which are also shown in the preceding papers.<sup>-7</sup> Water vaporvapors can be adsorbed mainly in the internal nanospaces, as these <u>CNTs</u> carbon nanotubes form bundles. Thus, the SWCNTs and DWCNTs act <u>asproduce</u> typical 1D and quasi-1D systems, respectively, for water, respectively. Figure 2(a) shows <u>changes in</u> the fractional filling <u>changes</u> of water with time progress. Water vaporvapors are adsorbed <u>only frombelow</u> 0.1 s <u>onward</u>, then begin to be adsorbed at that time, and finally, achieve equilibrium fractional fillings is achieved. The fractional fillingfillings of water vapor for the 1D system were increases rapidly-significantly increased **Comment [A1]:** You can include the telephone number and fax number of the corresponding author here.

**Comment [A8]:** This expression appears confusing. In particular, it is unclear what "low" means in this context. Are you referring to the small amount of metal catalyst required for their preparation? Or are you referring to metals with a smaller atomic number? Please revise this expression suitably according to your intended meaning.

**Comment [A9]:** This expression appears to be written more commonly as simply "TIP5P model." Please check whether this is indeed correct. If yes, then make this revision here.

**Comment [A10]:** This information does not seem to be relevant here; simply giving the citation number here is sufficient. Therefore, I have deleted this part from here.

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**Comment [A2]:** Here, it would be helpful to give a few examples of the physiological processes being referred to. This would aid readers' understanding.

**Comment [A3]:** The meaning of this part is not entirely clear to me. Do you mean "it is essential to use a nanoporous material having a model water channel structure to determine the mechanism of water transport through the channels"?

**Comment [A4]:** I have made this revision to avoid repetition of text. Please ensure that the revision is correct.

**Comment [A11]:** This part appears slightly ambiguous. Do you mean "as determined previously"?

**Comment [A5]:** For better clarity, I suggest that this part be revised as "hydrophobic one-dimensional (1D) nanospaces that water can permeate."

**Comment [A6]:** I have added this introductory term here to improve the flow from the previous sentence to this one.

**Comment [A7]:** I have edited this part to make it grammatically correct. However, should it appear as "CNT membranes"?

**Comment [A12]:** Here, it is not entirely clear what "equilibrium fractional filling" means. Do you mean "fractional filling factor stabilizes after approximately 10 s"?

from 0.5 to 10 s<sub>7</sub> and that for while these for the quasi-1D system increases gradually in the same timewere gradual increased. Therefore, we can say that water molecules are transported more rapidly-transport through 1D nanospaces-rather than through quasi-1D nanospaces. The Aadsorption rates of water vapor were evaluated from the differential fractional filling change, as shown in Figure 2(b); these rates which are directly representative of show the transportation of water vapor transportations through 1D or quasi-1D nanospaces. Furthermore, we observed Eextremely fast adsorption of water vapor below 0.2 s, relatively fast adsorption from 0.2 to 10 s, and gradual-progress of adsorption above 10 s-were observed. Fast adsorption below 0.2 s is caused by diffusion of a result of water diffusion into the internal nanospaces. The relatively fast and gradual adsorption-process should behaviors are believed to result from water vapor transportation of water vapor through the nanospaces. In comparison with the quasi-1D system, faster adsorption in the 1D systemnanospaces is actually observed from 0.2 to 10 s.

Results of the MD simulation of water in the internal nanospaces are shown in Figure 3; they demonst stratesalso clarify the mechanism theof water vapor transportation through 1D and quasi-1D nanospaces, as shown in Fig.3. The absolutely different experiment and MD simulation n tho different length of carbon nanotubes. This Ffigure-3 also shows the snapshots of water transportations through the 1D and quasi-1D nanospaces. At 0 ps, Wwater molecules are locatedsituated on the left side of in the CNTscarbon nanotubes, after which at 0 and then<u>they</u> begin movingto flow to the right side. Fractional filling of water is calculated from the number of water moleculesar number in on the right side. In the 1-nm diameter DWCNT, Waterwater transports rapidly transports to the right side in the carbon nanotube of 1 nm diameter, in contrast to while rater transportation for that in the 2- to 3-nm diameter diffusivity of water vapor in the DWCNTfor 1 nm diameter increases rapidly quickly increased with oscillation. On the other handsIn contrast, the increase of thermal diffusivity for in the 2nm--diameter SWCNT increases moderatelyis moderate. After 10 <u>ps, The the thermal diffusivities of water vapor are  $28 \times 10^{-9} \text{ m}^2 \text{ s}^{-1}$ </u> for the 1D nanospace and  $4 \times 10^{-9} \text{ m}^2 \text{ s}^{-1}$  for the quasi-1D anospace in 10 ps. However, after 70 ps, they define a state of the s and  $28 \times 10^9$  m<sup>2</sup> s<sup>1</sup>-in 70 ps, respectively, see Supporting Information). These valuestendencies agreecorrespond with the experimental adsorption rates, as shown in Figure 2. The number of Numbers of hydrogen bondsbonding are evaluated from the snapshots, in Floure 1 as shown in Figure 4. In the 1D system, the number of hydrogen bondsbonding is constant at (0.4) during the hadronic sterr. the calculation time step. In the quasi-1D system, however, the number of hydrogen <u>bonds</u> bonding is increasesd from 1 and to more than over 3 in 50 ps. <u>Thus</u>, That is, it is clear that the water molecules are unable tocannot form sufficientadequate hydrogen bonds-bonding in the 1D system; this lack of bond formation Hence, hydrogen bonding formation hindersprevents from water transportation through 1D nanospaces, because these 1D nanospaces provide a path for rapid water transportation, as mentioned above. Thus, the experimental and simulation results are in good agreement; the minor difference in the experimental and MD-simulation time scales of water transportation is attributed to the difference in the lengths of the CNTs considered in the experiment and simulation.

In this <u>studypaper</u>, <u>we investigated</u> direct water transportation through 1D and quasi-1D nanospaces <u>of CNTs</u>-were observed. Water vapor <u>was found to transport more</u> rapidly-<u>transports</u> through 1D nanospaces <del>rather</del> than <u>through</u> quasi-1D nanospaces, which is in agreement with<u>supported</u> by results of MD simulations. The rapid transport of water molecules in 1D nanospaces is attributed to the restricted formation of hydrogenHydrogen bonds bonding formation of in water molecules in these is restricted by 1D nanospaces. Therefore, as a result of restriction of hydrogen bonding formation water molecules rapidly transport through 1D nanospaces.

#### Associated conten

**SUPPORTING INFORMATION AVAILABLE** N2 adsorption isotherms at 77 K for SWCNT and DWCNT, and the <u>analysis of</u> nanospace structure analysis. Water vapor adsorption isotherms at 303 K and thermal diffusivities of water by MD simulation. This material is available free of charge via the Internet at

http://pubs.acs.org.

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## ACKNOWLEDGMENT

This research was supported by a Research Fellowship from the Foundation foref the Promotion of Ion Engineering, Kurata Memorial Hitachi Science and Technology Foundations, Nippon Sheet Glass Foundations, Global COE program, MEXT, Japan, and a Grant-in-Aid for Scientific Research (A) (No. 21241026) by the Japan–Society for the Promotion of Science. TEMobservations were performed at the Chemical Analysis Center Chiba University. **Comment [A13]:** Again, this expression is slightly confusing. Do you mean "differential change in the fractional filling factor"?

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**Comment [A14]:** This part is slightly confusing. Do you mean "taken as"? Or is the fractional filling actually *calculated* from the number of water molecules? If yes, then how exactly is it calculated? Please revise this part suitably according to the intended meaning.

**Comment [A15]:** Whose oscillation is being referred to here? Oscillation has not been mentioned prior to this in the text. Please clarify.

**Comment [A16]:** It is not clear what "time step" refers to here. Please explain what time step means in the current context.

**Comment [A17]:** Please check whether this revision is correct. If yes, then consider specifying the exact value (in decimals) greater than 3 (e.g., 3.4).

**Comment [A18]:** I have revised this part to make it grammatically correct. However, it does not make sense logically. How can hindering of water transportation result in rapid water transportation? Do you mean "this lack of bond formation promotes rapid transportation of water through 1D nanospaces, as mentioned above"?



Figure 2. <u>Changes in fractional + Inling changes</u> and rates of water vapor adsor<u>ptionbed</u> in the internal nanospaces of <u>CNTs.carbon nanotubes.</u>

REFERENCES

- (1) (a) Gragson, D. E.; Richmond, G. L. J. Phys. Chem. B 1998, 102, 3847. (b) Koop, T.; Luo, B.; Tsias, A.; Peter, T. Nature 2000, 406, 611. (c) Matsumoto, M.; Saito, S.; Ohmine, I. Nature 2002, 416, 409. (d) Headrick, J. M.; Diken, E. G.; Walters, R. S.; Hammer, N. I.; Christie, R. A.; Cui, J.; Myshakin, E. M.; Duncan, M. A.; Johnson, M. A.; Jordan, K. D. Science 2005, 308, 1765. (e) Pal, S. K.; Peon, J.; Bagchi, B.; Zewail, A. H. J. Phys. Chem. B 2002, 106, 12376. (f) Werder, T.; Walther, J. H.; Jaffe, R. L.; Halicioglu, T.; Koumoutsakos, P. J. Phys. Chem. B 2003, 107, 1345. (g) Lee, H. M.; Suh, S. B.; Lee, J. Y.; Tarakeshwar, P.; Kim, K. S. J. Chem. Phys. 2000, 112, 9759. (h) Bellissent-Funnel, M.-C. J. Chem. Phys. 1996, 22, 10023. 22. (i) Do, D. D.; Do, H. D. Carbon 2000, 38, 767. (j) Ohba, T.; Kanoh, H.; Kaneko, K. J. Phys. Chem. B 2004, 108, 14964. (k) Alcaniz-Monge, J.; Linares-Solano, A.; Rand, B. J. Phys. Chem. B 2001, 105, 7998. (1) Slasli, A. M.; Jorge, M.; Stoeckli, F.; Seaton, N.A. *Carbon* 2003, 41, 479. (m) Desbiens, N.; Demachy, I.; Fuchs, A. H.; Kirsch-Rodeschini, H.; Soulard, M.; Patarin, J. Angew. Chem. Int. Ed. 2005, 44, 5310. (n) Striolo, A.; Gubbins, K. E.; Chialvo, A. A.; Cummings, P. T. Mol. Phys. 2004, 102, 243. (o) Ohba, T.; Kaneko, K. Mol. Phys. 2007, 105, 139. (p) Ohba T.; Kaneko, K. J. Phys. Chem. C 2007, 111, 6207.
- (a) Agre, P.; Brown, S.; Nielsen, S. Curr, Opin. Cell. Biol. 1995, 7, 472. (b) Murata, K.; Mitsuoka, K.; Hirai, T.; Walz. (2)T.; Agre, P.; Heymann, J. B.; Engel, A.; Fujiyoshi, Y. Nature 2000, 407, 599. (c) Agre, P.; Kozono, D. FEBS Lett. 2003, 555.72.
- Sansom, M. S. P.; Biggin, P. C. *Nature* 2001, *414*, 156.
  (a) Hummer, G.; Rasaiah, J. C.; Noworyta, J. P. *Nature* 2001, *414*, 188. (b) Kalra, A.; Garde, S.; Hummer, G. *Proc. Natl. Acad. Sci. USA* 2003, *100*, 10175. (c) Holt, J. K.; Park, H. G.; (3)(4)

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- (a) Hata, K.; Futaba, D. N.; Mizuno, K.; Namai, T.; Yumura, (5)M.; Iijima, S. Science 2004, 306, 1362. (b) Ohba, Matsumura, T.; Hata, K.; Yumura, M., Iijima, S.; Kanoh, H.; Kaneko, K. J. Phys. Chem. C 2007, 111, 15660. (c) Endo, M.; Muramatsu, H.; Hayashi, T.; Kim, Y. A.; Terrones, M.; Dresselhaus, M. S. *Nature* 2005, 433, 476. Mahoney, M. W.; Jorgensen, W. L. J. *Chem. Phys.* 2000, 112. (6)
  - 8910.

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