

Confinement by Carbon Nanotubes Drastically Alters the Boiling and Critical Behavior of Water Droplets

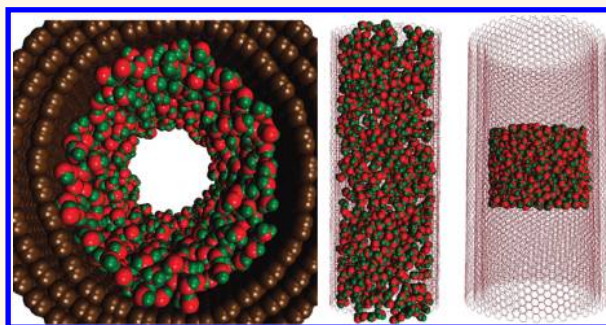
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Confinement of conventional liquids inside nanoscale structures gives rise to fascinating physical properties that form the basis for an ever-increasing number of novel applications. Significant efforts have been devoted to the study of structure, dynamics, and thermodynamics of water inside carbon nanotubes (CNTs).^{1–8} Both experimental and computational studies suggest that confined fluids exhibit a variety of new properties that differ greatly from those of bulk fluids. For instance, water readily forms ice-like structures inside CNTs even at room temperature.^{4,9} Confinement notably softens water dynamics, resulting in a drastic reduction of shear viscosity as the CNT diameter decreases to a few effective solvent diameters.¹⁰ Polar solute molecules, driven by dipole–dipole interactions, tend to aggregate,⁵ and the aggregation is essential for CNT filling.

The thermodynamics of solid/liquid phase transitions has received significant attention due to important implications in biology and materials science. Hydrogen-bonded chains of water structures in protein ion channels can greatly facilitate proton transport across cellular membranes. Uniform diffusion of ionic solutions through nanoscale carbon architectures is essential for reliable charging and discharging of supercapacitors.¹¹ Investigations show that freezing under confinement can take place by both first-order and continuous phase transitions,^{3,12} while bulk liquids freeze only through first-order transitions. Liquid/vapor coexistence of confined fluids has received significantly less attention. It has been found for *n*-alkanes that the critical point exists at lower temperature and higher density compared to bulk.¹³ This result can be explained by competition between fluid–fluid and fluid–surface interactions in combination with the entropic effect.

ABSTRACT



Vapor pressure grows rapidly above the boiling temperature, and past the critical point liquid droplets disintegrate. Our atomistic simulations show that this sequence of events is reversed inside carbon nanotubes (CNT). Droplets disintegrate first and at low temperature, while pressure remains low. The droplet disintegration temperature is independent of the CNT diameter. In contrast, depending on CNT diameter, a temperature that is much higher than the bulk boiling temperature is required to raise the internal pressure. The control over pressure by CNT size can be useful for therapeutic drug delivery.

KEYWORDS: phase transition · carbon nanotube · water · atomistic simulation

CNTs are promising candidates for novel therapeutic technologies.^{14–16} Their unique mechanical, electrical, and optical properties can be utilized simultaneously to create drug delivery vehicles.⁵ Hydrophobic CNTs penetrate through cell membranes and reach locations that are generally inaccessible to polar drug molecules.^{17,18} Hence, one can encapsulate drugs inside CNTs and deliver them to desired parts of the living body. Unlike living tissues, CNTs absorb near-infrared light, which allows for the photoactivation of *in vivo* CNTs without damaging side effects.^{19,20} Upon activation, the photon energy rapidly transfers from electrons to phonons²¹ and, subsequently, is released as heat to the encapsulated solution.²² Heating destroys CNT capping agents, freeing the drug. Increasing the

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vapor pressure of the encapsulated solvent greatly contributes to the drug release process.²³

The current paper establishes the effects of spatial confinement on the evaporation and resulting vapor pressure of water droplets encapsulated within CNTs. Using classical molecular dynamics (MD), we show that vaporization of water inside CNTs occurs by an entirely different mechanism than in the bulk. In the bulk phase, boiling is followed by droplet disintegration, as the temperature rises above the critical point. In contrast, water droplets inside CNTs disintegrate at a significantly lower temperature and lower pressure than free droplets. In order to achieve the rapid pressure growth associated with boiling, confined water must be heated to a temperature far beyond the droplet disintegration temperature. While the temperature required for the pressure growth depends on the CNT diameter, the droplet disintegration temperature is diameter independent. The latter fact is quite remarkable, since most properties of confined liquids do depend on the CNT size. We show that the mechanism of water droplet evaporation inside CNTs is controlled by the strength of the CNT–water interaction.

RESULTS AND DISCUSSION

Bulk Liquid/Vapor Interface. Liquid vaporization can be simulated by introducing a liquid/vapor interface in the simulation cell, Figure 1. This can be achieved starting from conventional bulk liquid and extending the box along one of its three Cartesian directions. The resulting system contains an endless film of liquid surrounded by several nanometers of vacuum. To simulate the interface, it is critical that both liquid and gas phases are present simultaneously in the simulation cell. Since, generally, vapor is 3 orders of magnitude less dense than liquid, proper representation of the gas phase requires a significantly larger simulation volume compared to the liquid phase.

In order to select an optimal box size for the simulation of the liquid/vapor interface, we tested the thermodynamic properties of three systems with different simulation volume and the same amount of liquid in the NVT ensemble, Table 1. The liquid film was centered in boxes with dimensions of $3.74 \times 3.74 \times 10$, $3.74 \times 3.74 \times 20$, and $3.74 \times 3.74 \times 30$ nm. The internal pressure in these systems is generated mostly by the gas phase, and hence, the pressure component normal to the film surface is dominant. Water boils when its vapor pressure exceeds one atmosphere (for standard conditions). For TIP4P water, this happens at 368 K,²³ which is close to the experimental value of 373 K. The simulated boiling temperature is independent of the simulation box volume, as should be expected, Figure 2.

The three curves shown in Figure 2 diverge at the critical point of TIP4P water. At this temperature the

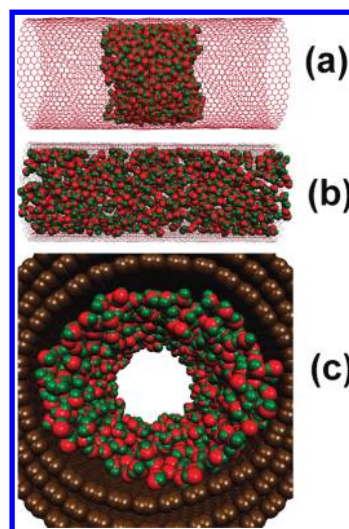


Figure 1. (31,31) CNT containing TIP4P water. (a) At a low temperature, $T = 300$ K, water exists as a droplet. (b) At a high temperature, $T = 650$ K, evaporated water is distributed uniformly throughout the tube. (c) At an intermediate temperature, $T = 500$ K, water adsorbs onto CNT sidewalls.

TABLE 1. Simulated Systems Consisting of a Water Droplet inside a CNT^a

system	CNT	d_{CNT} , nm	N_{waters}	T_b , K	T_c , K	T_w , K
1			1728	368	575	
2	(11,11)	1.49	89	897		490
3	(21,21)	2.85	462	702		500
4	(31,31)	4.20	1112	618		490

^a See Figure 1a. The CNTs are 10 nm long. System 1 represents bulk TIP4P water and includes a water layer in a rectangular box, as detailed in the text. T_b and T_c/T_w correspond to the steep rise of the vapor pressure (boiling) and droplet disintegration. For bulk, liquid phase disappears at the critical temperature, T_c . Confined water droplets disintegrate due to the wetting transition, T_w . The order of the boiling and disintegration events is reversed for confined droplets relative to bulk.

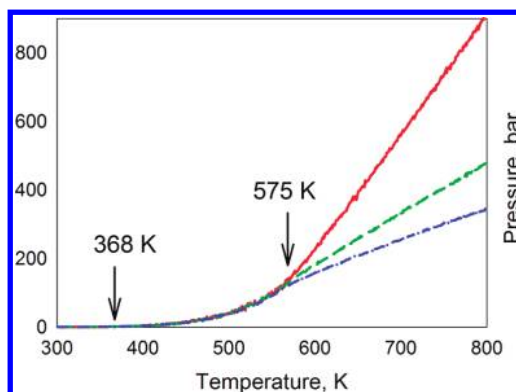


Figure 2. Pressure vs temperature for bulk water obtained for system 1, Table 1, placed in boxes with the same cross-section and different lengths: 10 nm (red solid), 20 nm (green dashed), and 30 nm (blue dash-dotted). Each system contains 1728 water molecules. The arrows show the boiling and critical temperatures, 368 and 575 K, respectively. Above its critical point, water cannot be condensed at any temperature; therefore, the three lines corresponding to different system volumes diverge.

liquid layer of water disappears and pressure becomes dependent on the simulation box volume. The TIP4P model of water,²⁴ Figure 2, underestimates the experimental values of the critical temperature and pressure, 647 K and 228.5 bar, respectively. The improved TIP4P/2005 model agrees better with experiment in this case.²⁴ Since both boiling and critical points coincide for the three MD simulation cells, we concluded that all of them provide proper representation of the liquid/vapor interface. In order to save computer resources, we chose the smallest box for the production simulations.

Disintegration of Confined Water Droplets and Anomalous Pressure Changes. Next, we investigated pressure changes inside the (11,11), (21,21), and (31,31) CNTs, Table 1, induced by gradual heating in the constant temperature, constant volume ensemble. The resulting curves shown in Figure 3 are qualitatively different from the data for bulk water. First, the temperature required for a rapid pressure growth depends on the CNT diameter, and second, the initial pressure growth is significantly sharper for confined water. Moreover, the pressure of the confined liquid remains very close to zero at low temperatures, while the pressure of bulk water is finite and grows continuously even below the boiling point. It should be noted, now, that the molecular processes underlying the pressure growth inside CNTs are qualitatively different from the boiling of bulk water, as discussed in detail below. The T_b values listed in Table 1, corresponding to the inflection points in Figure 3, can be called boiling temperatures only nominally. Both the T_b values and the $P(T)$ slopes correlate strongly with the CNT diameter. In particular, T_b is inversely proportional to the diameter.

Further understanding of the effect of the CNT–water interaction on the evaporation of confined water droplets is provided by the data of Figure 4. Here, we use the Lennard-Jones epsilon parameter of the carbon atoms as a tool to control the strength of water attraction to the hydrophobic CNT sidewalls. Changing epsilon affects the repulsive part of the potential as well; however, this change is less important for the present study, since the infinite repulsive wall remains even for small epsilon values, for which the attraction disappears. Focusing on the (31,31) system, we obtained pressure as a function of temperature for a series of diminishing values of epsilon. As should be expected, decreasing the attraction of water to the CNT sidewalls increases pressure. When epsilon approaches zero, the $P(T)$ curve for the confined droplet starts to resemble the bulk data. Nevertheless, even in the absence of the CNT–water attraction, the confined droplet and bulk lines never become identical. The difference in the T_b values is equal to 16 K; $T_b = 384$ K for confined water in the absence of CNT–water attraction, and $T_b = 368$ K for bulk TIP4P water.^{23,24} The difference arises solely due to confinement and is

smaller for larger tubes. In the limit of small epsilon, the transition from the low-pressure region to the region with rapidly rising pressure is smooth, and pressure remains finite even at low temperatures. In contrast, for larger epsilon, pressure remains essentially zero at low temperatures, suggesting that water molecules evaporating from the droplet adsorb on the CNT walls.

The fact that water pressure inside CNTs is significantly affected by changes in the van der Waals parameters of CNT sidewalls indicates that pressure can be tuned by CNT functionalization.^{25,26} Chemical changes made to the outer surface of single-walled CNTs will influence the internal vapor pressure. Relative to the attractive CNT–water interaction, the effect of spatial confinement *per se* on the internal vapor pressure is quite small. Overall, both confinement and the attractive interactions can be used to control quantitatively the evaporation process inside CNTs. Decreasing epsilon by a factor of 4 is comparable to the change in the CNT diameter from 2.85 to 4.20 nm; compare Figures 3 and 4. The decrease of epsilon by 4 times results in the decrease of boiling temperature from 618 K to 555 K, Figure 4, while the increase of the CNT diameter from 2.85 nm to 4.20 nm results in the decrease from 702 K to 618 K. The changes of temperature in these cases are comparable. Decrease of the CNT diameter from 2.85 nm to 1.49 nm in the (11,11) CNT creates a substantial growth in the T_b value, up by 195 K. At the same time, pressure generated in the (11,11) CNT at temperatures above T_b is relatively small, indicating that substantial pressure of water vapor inside CNTs with diameters below 2.5 nm cannot be attained. The large surface-to-volume ratio ensures that adsorption of water on the CNT surface greatly influences thermodynamic properties of liquids confined in small CNTs.

The data presented in Figures 3 and 4 suggest the following scenario for evaporation of water droplets inside CNTs. At temperatures well below T_b for a given tube water molecules with high kinetic energy escape the droplet but are immediately adsorbed on CNT walls. Vapor pressure is not generated. This process can occur at temperatures significantly above the boiling temperature of bulk water. Raising the temperature further increases the kinetic energy of water, which works to overcome the van der Waals attraction, releasing water from the CNT walls. As a result, pressure grows rapidly inside CNT.

The Wetting Transition. The transport of water molecules from the confined droplet to the CNT walls merits further investigation. Figure 5 shows the water density distribution function calculated along the axial direction of the three CNTs at a series of temperatures. The corresponding bulk data are computed along the direction perpendicular to the water layer. The data for 300 K show stable liquid droplets in equilibrium with

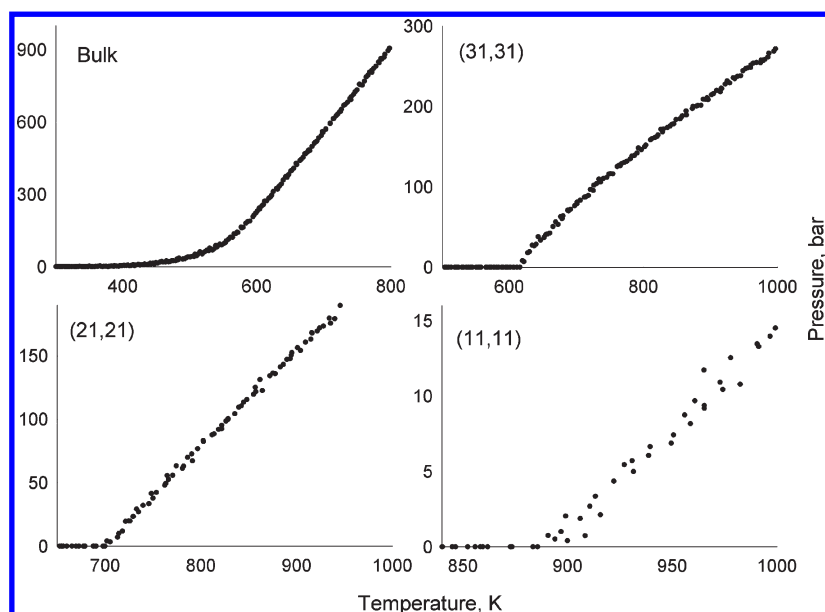


Figure 3. Pressure vs temperature for bulk and confined water droplets. Pressure rises rapidly above a certain threshold temperature, nominally boiling temperature, which is different for each system depending on the strength of the confinement effect.

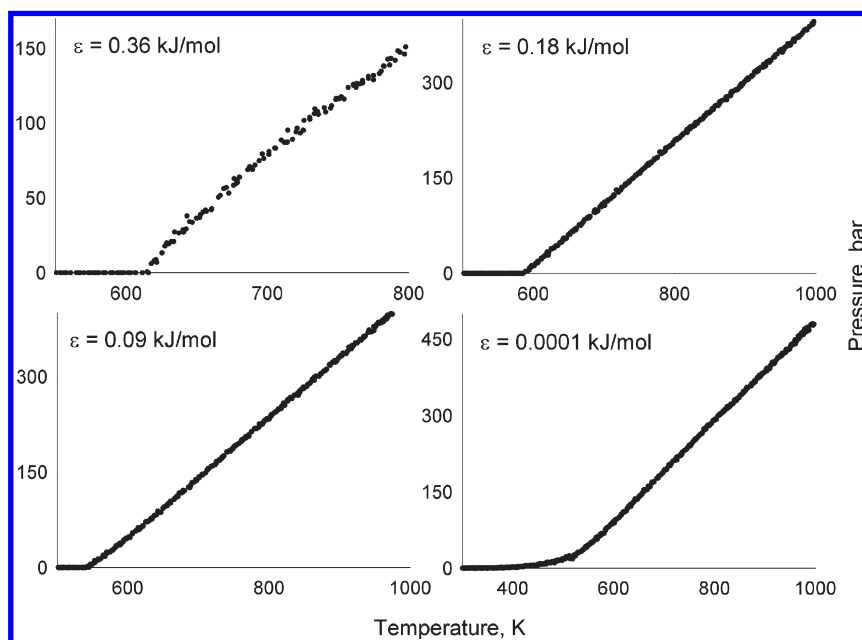


Figure 4. Pressure vs temperature in the (31,31) CNT system with different degrees of attraction of water molecules to CNT carbon atoms. The attraction strength was varied by changing the epsilon value in the Lennard-Jones equation. As epsilon decreases, the data approach those for bulk water, Figure 3.

low-density vapor. As the temperature is increased, the droplets disintegrate. The liquid layer representing bulk water disappears around 575 K, corresponding to the critical temperature of the TIP4P water model, Figure 2. The confined droplets destruct at a significantly lower temperature, T_w , around 500 K, due to the wetting transition.²⁸ Unlike most properties of confined media, T_w is independent of the CNT diameter.

It is interesting to note that the water droplet confined inside the smallest (11,11) CNT is very mobile.

It migrates by about 2 nm to the left within 40 ns of MD simulation. Such behavior can be explained by smoothness of the CNT hydrophobic sidewalls²⁷ and a relatively large contact area of the (11,11) CNT and the droplet. The hydrophobic interaction between water and the (11,11) CNT walls disrupts strong intermolecular interactions between water molecules. As a result, water molecules behave relatively independently, and the overall droplet is mobile. In contrast in larger CNTs, water–water interactions are disrupted

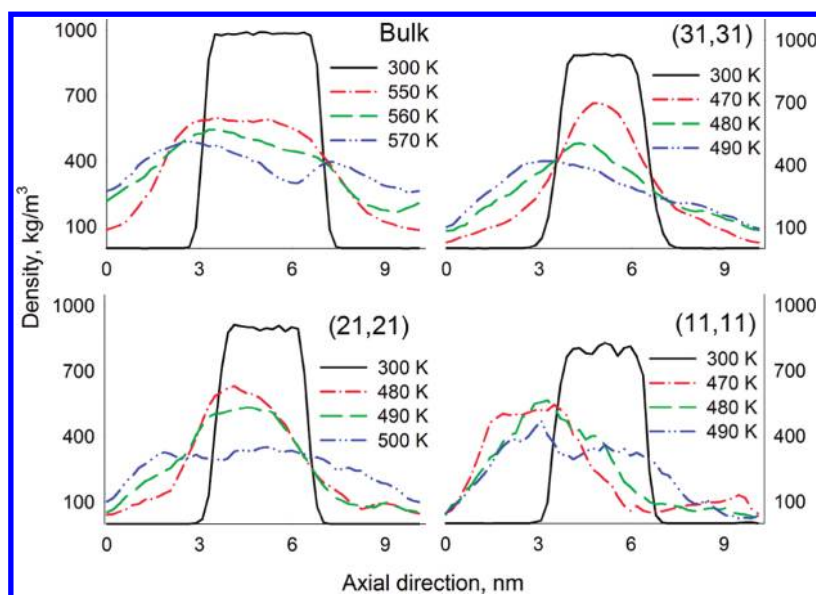


Figure 5. Water density distribution for free liquid film and inside the (31,31), (21,21), and (11,11) CNTs computed along the tube axis at several temperatures. In bulk, the density becomes uniform above the critical temperature of 575 K. Under confinement, the droplet disintegrates at a lower temperature of about 500 K, independent of the CNT diameter.

less by the CNT walls, water molecules remain strongly bound to each other, and the droplets are less mobile.

The temperature-driven transport of water from the droplet, Figure 1a, to the CNT walls, Figure 1c, is an example of a wetting transition: if a fluid does not spread across a surface at a low temperature, then it does so at a higher, wetting transition temperature, T_w , Table 1. Our results well agree with the Monte Carlo simulations of water droplets on a graphite surface, showing the wetting transition at 480 K for a different model of water.²⁸

The wetting transition inside CNTs partially filled with water entirely undermines the possibility to use the capillary theory in order to predict the rapid vapor growth of confined water, Figures 3 and 4. In principle, one can use the Kelvin equation in order to compute the change of the vapor pressure due to the presence of a curved meniscus of water in the CNT capillary. Then one can employ the Clausius–Clapeyron relation in order to estimate the boiling point elevation for the given vapor pressure, relative to the unconfined system. Such calculations provide good agreement with the atomistic simulation for sufficiently large CNTs that are completely filled with water.²³ However, the boiling point elevations on the order of tens of Kelvins predicted by the capillary theory are significantly smaller than the temperatures required for the rapid pressure growth in the current MD calculations, Figure 3. Such disagreement should be expected,²⁹ since the meniscus disappears at an elevated temperature, Figure 1c. The capillary theory neglects the interaction of molecules in the saturated vapor with the walls of the nanovessel.

Figure 6 depicts the radial density distribution functions calculated perpendicular to the CNT axes

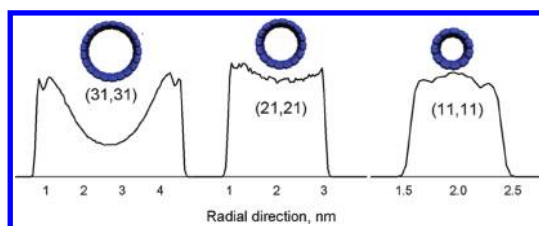


Figure 6. Radial density distributions of water confined inside the (31,31), (21,21), and (11,11) CNTs at 500 K. Water molecules tend to adsorb onto CNT walls, as illustrated in Figure 1c.

for water confined by the three CNTs at 500 K. At this temperature, the droplets have already disintegrated; see Figure 5. The radial distribution function for the (31,31) CNT shows two distinct density maxima near the opposite sidewalls. The widths of the maxima correspond to about two layers of adsorbed water. The slice across the (31,31) CNT system shown in Figure 1c gives an atomistic illustration of this situation: water molecules are adsorbed at the CNT walls, and a 1 nm wide cylinder of vacuum is created in the middle of the tube. The CNT–water attraction provides an additional force driving disintegration of the water droplets. It explains why the confined water droplets destruct at a temperature that is lower than the critical temperature of bulk water.

Water molecules inside the smaller (21,21) and (11,11) CNTs are also driven to CNT walls following droplet disintegration at 500 K. However, no vacuum is seen in the middle of the tubes, Figure 6. Only seven and three water molecules can be arranged perpendicular to the axes of the (21,21) and (11,11) CNTs, respectively. The inner radius of the (21,21) CNT is only 1.3 nm, and dipole–dipole interactions of water molecules adsorbed on the opposite sidewalls are strong.

This strong electrostatic interaction stabilizes a continuous distribution of molecules across the smaller tubes. The strong intermolecular interaction of water molecules across the whole width of the two smaller CNTs explains why vapor pressure remains nearly zero to significantly higher temperatures: compare 897 and 702 K for the boiling temperatures in the (11,11) and (21,21) CNTs with 618 K for the (31,31) CNT, Table 1.

Thermodynamic Considerations. The information presented above is now sufficient to rationalize the qualitative and quantitative differences in the evaporation of water droplets inside CNTs and in the free state. A free water droplet exists in equilibrium with its vapor pressure. If the volume of the system is fixed as in the current study, vapor pressure grows slowly with increasing temperature below the boiling point and rapidly above the boiling point. Finally, the water droplet disintegrates at the critical temperature. This sequence of events is reversed inside CNTs. The droplet disintegrates while pressure remains nearly zero, and only at a much higher temperature does the pressure start growing abruptly and at a rapid pace. These differences arise primarily due to the attractive interaction between the water droplet and CNT walls. The interaction prevents evaporated water molecules from diffusing freely inside the tube. Instead, they adsorb onto CNT sidewalls.

Thermodynamically, one can consider several phases of water in partially filled CNTs, including water droplets (3D liquid), water layers on CNT walls (2D liquid), individual water molecules physisorbed onto CNT walls (2D gas), and confined water vapor (3D gas). The energies and entropies of these phases vary oppositely of each other. In particular, the interaction energy decreases in the above series, since the water–water interaction is stronger than the water–CNT interaction. At the same time, the entropy grows in this series, as water molecules are allowed to explore a larger fraction of space inside CNTs. Raising the temperature increases the importance of the entropy term in the thermodynamic free energy of the system. As a result, the system undergoes a sequence of transitions from 3D liquid to 2D liquid (wetting transition), to 2D gas (not seen in our calculations due to a high concentration of water molecules), and then to 3D gas (evaporation).

The CNT–water interaction is responsible for lowering the droplet disintegration temperature from the critical temperature of 575 K for bulk TIP4P water down to 500 K. The value of 500 K is determined by the strength of the interaction and is largely independent of the CNT diameter. Once water transforms from a droplet into a film adsorbed on the CNT walls, vapor pressure can be increased only when the temperature becomes sufficiently high to overcome both the adsorption energy and the intermolecular interactions of water molecules within this layer. The desorption

temperature increases in small CNTs, because additional energy is required to overcome the electrostatic interaction between water layers adsorbed on the opposite walls.

Experimental Implications. The temperature dependence of the water droplet behavior predicted by our simulations can be detected experimentally, for instance, using transmission electron microscopy (TEM).²⁹ Our results indicate that it is particularly important in such experimental studies to achieve good control over CNT quality and functionalization, since the presence of defects and different functional groups on CNT walls can alter the strength of the CNT–water interaction and change the quantitative details of the considered phenomena. At the same time, the qualitative sequence of events predicted in our simulation should remain robust to variations in CNT samples.

The sharp growth of vapor pressure inside CNTs above a certain temperature, whose value can be controlled by CNT diameter and functionalization, can prove advantageous in CNT applications for drug delivery. The threshold temperature values may turn out too high for this purpose. In such a case, one can use other solvents with lower boiling temperatures and weaker intermolecular interactions, such as carbon dioxide or a carbon dioxide/water mixture, so-called carbonated water. The desired increase in the vapor pressure can be achieved at a significantly lower temperature for CNTs filled with water completely²³ compared to the CNT/droplet systems considered in the current work.

CONCLUSIONS

To recapitulate, MD simulations of water droplets inside the (11,11), (21,21), and (31,31) CNTs have been carried out over a wide range of temperatures. Under confinement, water droplets disintegrate at a lower temperature than in the bulk phase; however, little vapor pressure is generated. Sharp growth of vapor pressure inside CNTs is achieved by heating the system substantially above the droplet disintegration temperature. This situation is in contrast to the behavior of free water droplets, which produce high vapor pressure at the boiling temperature that is much smaller than the critical temperature required for droplet disintegration. The differences in the evaporation mechanisms of the free and confined water droplets are rationalized by the CNT–water interaction. First, the droplet transforms into a film of water molecules adsorbed at CNT walls. The temperature of this process, known as the wetting transition, is independent of the CNT diameter, which is quite unusual for properties of confined liquids. The droplet disintegration temperature is determined by the strength of the CNT–water interaction. Second, at a substantially higher temperature water abruptly generates strong vapor pressure. This temperature does depend on

the CNT diameter for small CNTs, since it is determined not only by the CNT–water interaction, but also by the interaction between water layers adsorbed on the opposite sides of the CNTs. The sharp increase of pressure

inside CNTs upon heating and the control over the temperature of this transition provided by the CNT diameter and sidewall functionalization can be utilized in therapeutic applications of CNTs.

METHODS

The simulations were performed with systems consisting of a water droplet encapsulated within a multiwalled CNT, Figure 1a. Three coaxial CNTs were used to prevent interaction of inner and outer water molecules. Guided by the 0.34 nm experimental inter-layer distance, the multiwalled CNTs were selected such that an (n,n) innermost CNT defined in Table 1 was encapsulated by the $(n+5, n+5)$ and $(n+10, n+10)$ outer CNTs. A water layer in equilibrium with its vapor pressure in a series of rectangular boxes was used to represent the properties of unconfined water, system 1 in Table 1.

The simulated CNTs were treated as flexible nonpolarizable objects. The potentials for bonds, angles, dihedrals, and pairwise 1–4 carbon–carbon Lennard-Jones interactions within the CNTs were represented by the AMBER force field (FF).³⁰ Water molecules were represented by the TIP4P four-site potential.³¹ Among the incredibly diverse set of water FFs, TIP4P is most successful in reproducing the liquid/vapor phase transition temperature, Figure 2. The electrostatic interaction energy for pairs of solvent molecules was calculated within the 1.3 nm cutoff radius using the reaction-field-zero technique.³² The Lennard-Jones (12,6) interactions were computed using the shifted force method,³² with the switch region from 1.1 to 1.2 nm. The Lorentz–Berthelot combination rules were used to derive the cross-term Lennard-Jones parameters. The selection of the interaction potentials for the CNT–water systems followed the well-tested practice.³³

The trajectories were propagated using GROMACS.³² The leap-frog integration algorithm with a 0.002 ps time-step was applied. The simulations started with empty, multiwalled CNTs surrounded with liquid water. A 5000 ps MD simulation in the NPT ensemble allowed water to enter CNTs and achieve relaxed configurations.³⁴ The thermostat of Bussi *et al.*,³⁵ often referred to as “V-rescale”, with the response time of 0.5 ps was used to maintain the temperature at 300 K. The Parrinello–Rahman barostat³⁶ with the time constant of 4.0 ps was used to maintain pressure at 1 bar. Following initial equilibration, all water molecules outside CNTs were deleted. Water droplets were created by removing 3.5 nm of confined water from each end of the CNTs, Figure 1a.

Simulated annealing was carried out by gradually heating the systems from 300 to 1000 K with at the rate of 1.0 degree per 200 ps. To ensure that the simulated water structure had sufficient time to adapt to the constantly increasing temperature, NVT runs were performed at several temperatures selected arbitrarily between 300 and 1000 K. The vapor pressure did not change during these test runs.

The pressure in the confined systems was defined as $P = 2(E_{\text{kin}} - G)/V$, where V is the system volume, and P , E_{kin} , and G are pressure, kinetic energy, and virial tensors, respectively. The virial tensor is given by $G = -(1/2)\sum_{i,j} \mathbf{r}_{ij} \otimes \mathbf{F}_{ij}$. The summations in the above equations involved pairs of atoms belonging only to water molecules, excluding CNT atoms. For isotropic systems, pressure is derived as $P = \text{trace}(P)/3$. For the confined droplets, pressure was defined by its component along the CNT axis.

Conflict of Interest: The authors declare no competing financial interest.

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