Activated drying in hydrophobic nanopores and the line tension of water

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Edited by Pablo Gaston Debenedetti, Princeton University, Princeton, NJ, and approved October 4, 2012 (received for review May 6, 2012)

We study the slow dynamics of water evaporation out of hydrophobic cavities by using model porous silica materials grafted with octylsilanes. The cylindrical pores are monodisperse, with a radius in the range of 1-2 nm. Liquid water penetrates in the nanopores at high pressure and empties the pores when the pressure is lowered. The drying pressure exhibits a logarithmic growth as a function of the driving rate over more than three decades, showing the thermally activated nucleation of vapor bubbles. We find that the slow dynamics and the critical volume of the vapor nucleus are quantitatively described by the classical theory of capillarity without adjustable parameter. However, classical capillarity utterly overestimates the critical bubble energy. We discuss the possible influence of surface heterogeneities, long-range interactions, and high-curvature effects, and we show that a classical theory can describe vapor nucleation provided that a negative line tension is taken into account. The drying pressure then provides a determination of this line tension with much higher precision than currently available methods. We find consistent values of the order of -30 pN in a variety of hydrophobic materials.

drying transition | hydrophobicity | kinetics | nanobubbles | mesoporous silica

A remarkable property of water is its ability to form nanosize bubbles, or cavities, on hydrophobic bodies (1). Since their first direct observations through atomic force microscopy about a decade ago (2, 3), surface nanobubbles on hydrophobic surfaces have raised considerable interest, and they are believed to play a major role in surface-driven phenomena, such as boundary slippage of water flows, heat transfer at walls, vaporization and boiling, surface cleaning, etc. (4, 5). In a different context, the evaporation of water in the vicinity of hydrophobic bodies has been studied as a core mechanism for the hydrophobic interaction mediated by water (6–8), which plays a central role in biological matter. The formation of cavities able to bridge hydrophobic units provides a driving force for protein folding and supermolecular aggregation (9). Simulation examples of such drying-induced phenomena include the collapse of a polymer chain, multidomain proteins, and hydrophobic particles (9–13).

Despite their direct observation, the easy formation and the high stability of nanobubbles on hydrophobic bodies still raise fundamental questions (5, 14, 15). Because of significant theoretical work, it is now established that, at the scale of the nanometer, macroscopic concepts apply: hydrophobicity is described by interfacial energies, and the drying transition in hydrophobic confinement is a first-order transition triggered by the nucleation of a critical vapor bubble (1). The energy barrier limiting the kinetics of this transition is a strong signature of nanobubbles properties. Evaporation kinetics has also been pointed out as the most direct measure of the importance of hydrophobic collapse in protein folding (9). However, rate effects in the drying transition have not received much attention. A few numerical studies have addressed the rate of evaporation of liquid water confined between hydrophobic plates (16–19). The nucleation barrier has been measured with different methods, and it has been shown to increase strongly with the slit separation. The classical theory of capillarity has been shown to overestimate the numerical findings (17, 18). The classical capillarity is a key framework to understand nucleation phenomena, but it is based on macroscopic considerations and does not include specific features, like fluctuations or line energies that can affect interfaces at nanometric scales. There is, however, no consensus about the leading effect at these scales. Fluctuations are invoked in ref. 18 to explain the observed nucleation barrier reduction, whereas line tension is shown to account for the observed deviations in ref. 19. Experimental studies are scarce, because the rate or time variable is generally ignored in studies of adsorption and desorption of confined liquids.

Here, we use highly ordered nanoporous silicas to study the dynamics of water evaporation in hydrophobic confinement. Micelle-templated silicas (MTSs) have quasi-1D mesopores shaped in the form of cylinders of monodisperse radius adjustable from 1 to 5 nm. These model materials have been used as nanoscale laboratory to study the phase diagram of confined liquids (20, 21). In previous works (22, 23), we used silane-grafted MCM-41 (Mobil Crystalline Material 41) to study water confined by hydrophobic walls. Liquid water penetrates into the nanopores at high pressure, reaching 500 bars for nanometer-sized pores. The intrusion pressure of water in the cylindrical pores scales as the inverse of their radius down to radii of 1.3 nm, according to the Laplace law of capillarity (Eq. 1):

$$P_{\rm int} = -\frac{2\gamma_{\rm lv}\cos\theta}{R_{\rm p}} = 2\frac{\gamma_{\rm sl} - \gamma_{\rm sv}}{R_{\rm p}},$$
[1]

with γ_{sl} , γ_{sv} , and γ_{lv} being the solid/liquid, solid/vapor and liquid/ vapor surface tensions, respectively. This intrusion law shows the model character of hydrophobized MTS to provide a geometrically and energetically well-defined confinement. The drying transition is obtained by lowering the pressure. Liquid water becomes metastable and finally, empties the nanopores at a pressure P_{ext} lower than the intrusion pressure (24). The drying pressure P_{ext} is not described by the Young–Laplace law (1) and increases with temperature (23). Those features are in good qualitative agreement with a drying triggered by the nucleation of a critical vapor bubble on the pore walls.

Author contributions: G.V. and É.C. designed research; L.G. and T.B. performed research; T.B. and A.G. contributed new reagents/analytic tools; L.G., G.V., and É.C. analyzed data; and É.C. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10. 1073/pnas.1207658109/-/DCSupplemental.

We report here investigations of the drying kinetics of the hydrophobic nanopores. For this study, we have developed a device that allows us to perform intrusion–extrusion cycles at finite rates ranging from 0.1 to 100 s^{-1} (25). The dynamic study allows us to get quantitative access to the volume of the critical vapor bubbles initiating the drying and the energy barrier. Although the volume of the critical bubbles is remarkably well-predicted by the classical theory of capillarity, the latter overestimates considerably the energy barrier for their formation. We show that the low energy barrier observed is a strong support of a negative line tension of water on the silane monolayer. The drying pressure provides a measurement of this line tension, completely independent from other experimental methods and with a much higher precision.

Results

We use here three different MTSs: MCM-41 (26, 27), SBA-15 (Santa Barbara Amorphous 15) (28, 29), and HMS (Hexagonal Mesoporous Silica) (30, 31), which are all shaped in the form of cylindrical pores with a narrow size distribution. Their synthesis, silanization with octyldimethylsilane, and pore size determination with nitrogen sorption at 77 K are described in SI Text, section I. The materials exhibit some differences in their organization and internal pore surface. The pore radii of the grafted materials are 1.34 ± 0.1 nm (MCM-41), 1.54 ± 0.1 nm (HMS), and 2.15 ± 0.25 nm (SBA-15) (Table 1). An instrumented, deformable cell is filled under vacuum with the degassed material and pure water, and water is forced in the pores up to full saturation (Fig. 1). The cell volume V is then increased at a constant rate until water empties the nanopores, and the initial cell volume is recovered. The intrusion and drying transitions appear as quasiplateaus on the pressure-volume (P-V) curves (Fig. 1).

Fig. 2 shows the typical behavior of the drying pressure as a function of the time t_{ext} of the drying process. The latter is defined and measured as the time spent on the drying plateau (Fig. 1). A logarithmic growth is obtained for all of the MTSs at all of the temperatures investigated. In contrast, the intrusion pressure exhibits much smaller kinetic effect.

This logarithmic kinetics is a strong signature of the activated processes that govern the drying transition. We argue that the mechanism limiting the drying process is the nucleation in each pore of a vapor bubble extending across the section and forming



Fig. 1. Intrusion/extrusion of water in hydrophobic mesoporous silicas. A thermally regulated cell containing water and the material is placed in a traction machine (*Left*) to measure the pressure-volume isotherms (*Lower Right*). The volume change is driven at a constant velocity in the range of 0.08-80 mm/s. The intrusion and extrusion pressures, P_{int} and P_{ext} , respectively, are determined as the average pressure in the corresponding plateaus of the P-V isotherms (24).



Fig. 2. Variation of the extrusion pressure P_{ext} with the logarithm of the time t_{ext} during which extrusion occurs for the MCM-41 material at different temperatures. The other materials show similar logarithmic growth of P_{ext} with t_{ext} .

two disconnected menisci (Fig. 3). The drying time, in each independent pore of average length *L*, is then related to the rate of nucleation of a spanning bubble: $I = (\nu L/b)e^{-\Delta \Omega_c/k_B T}$, by $It_{ext} \cong 1$. Here, $\Delta \Omega_c$ is the energy of the critical vapor nucleus, *b* and ν are microscopic length scale and frequency, respectively, and $k_B T$ is the thermal energy. This rate leads to a classical nucleation law (Eq. 2):

$$\Delta \Omega_{\rm c} = k_{\rm B} T \ln(L\nu t_{\rm ext}/b).$$
 [2]

Dimensionally, we expect that $\Delta\Omega_c$ depends on the pressure only through a term PV_c involving the volume of the critical nucleus V_c . Hence, the drying pressure should express as (Eq. 3):

1

$$P_{\text{ext}} = \frac{k_{\text{B}}T}{V_{\text{c}}} \ln \frac{t_{\text{ext}}}{t_{\text{o}}} + P_{\text{ext}}^{\text{o}}(T), \qquad [3]$$

with $P_{\text{ext}}^{\text{o}}(T)$ being the extrusion pressure measured at some reference extrusion time t_{o} . If the volume of the vapor nucleus does not depend on the liquid pressure, P_{ext} is expected to grow logarithmically with t_{ext} . This result describes our data very well. We get values of V_{c} from the inverse of the slope of the experimental P_{ext} vs. $\log(t_{\text{ext}})$ plots for each material and temperature. We obtain $V_{\text{c}} = 10.2 \pm 1.5 \text{ nm}^3$ for the MCM-41, $V_{\text{c}} = 17.8 \pm 2.7 \text{ nm}^3$ for the HMS, and $V_{\text{c}} = 51 \pm 17 \text{ nm}^3$ for the SBA-15 (Fig. 3). The drying of SBA-15 shows a weaker dynamical behavior than the two others, and therefore, the uncertainty on V_{c} is much larger.

Because classical capillarity describes successfully the intrusion pressure, we compare V_c to the macroscopic calculation in the work by Lefevre et al. (22) for the nucleation of a bubble in a cylinder. The energy barrier is given within 5% by the approximate expression (Eq. 4):

$$\Delta\Omega_{\rm c} \simeq P_{\rm L}K_1(\theta)R_{\rm p}^3 + K_2(\theta)\gamma_{\rm lv}R_{\rm p}^2, \qquad [4]$$

where $P_{\rm L}$ is the liquid pressure, and K_1 and K_2 are functions of the Young's contact angle θ (detailed in *SI Text*, section I). Note here that, in contrast to bulk nucleation, the critical volume $V_{\rm c} = R_{\rm p}^3 K_1(\theta)$ does not depend on the applied pressure. The reason for this lack of dependence is that the formation of two disconnected menisci from a bubble growing at the wall of a cylinder occurs through a capillary instability, which is explained in Fig. 4. The theoretical volume $R_{\rm p}^3 K_1(\theta)$ of the critical nucleus can be calculated by taking the contact angle obtained from the



Fig. 3. (*A*) Schematic representation of the nucleation process: the critical vapor bubble is able to form two disconnected menisci. (*B*) Nucleus volume V_c measured from the slope of the logarithmic growth of P_{ext} as a function of t_{ext} using Eq. **3** in the three materials at various temperatures. The SBA-15 has a lower extrusion pressure with a smaller slope, leading to a larger uncertainty. The colored rectangles are the theoretical values $K_1(\theta)R_p^3$ calculated using nitrogen sorption pore radii and the contact angle derived from the intrusion pressure (Table 1 and *SI Text*).

intrusion pressure (Eq. 1) as the value of the Young's contact angle θ . Fig. 3 shows its remarkable agreement with the experimental value for the three materials and the different temperatures studied. This agreement is obtained without adjustable parameter: pore radii and their uncertainty are used as derived from the nitrogen sorption isotherms (*Materials and Methods*).

Discussion

The ability of classical capillarity to describe quantitatively the volume of the critical vapor bubble is remarkable. We should emphasize that we observe here a cavitation process occurring at positive pressures of order of 80–150 bars, whereas usual cavitation in water occurs at strongly negative pressures. This effect is the result of hydrophobic confinement.

However, if classical capillarity predicts, with great precision, the critical bubble volume and the slow dynamics of the drying pressure, it fails by orders of magnitude in predicting the reference pressure (Eq. 5):

$$P_{\text{ext}}^{o} = \frac{k_B T}{V_{\text{c}}} \ln \frac{L\nu t_{\text{o}}}{b} - \frac{K_2(\theta)}{K_1(\theta)} \frac{\gamma_{lv}}{R_{\text{p}}}.$$
[5]

Assuming a microscopic scale $b \sim 1-10$ Å and a microscopic frequency $\nu \sim 10^{12}$ to 10^{13} s⁻¹, Eq. 5 gives a negative value of P_{ext}^o (as low as -200 bars), which is in strong contrast with experimental observation giving P_{ext}^o above 50 bars for all materials and temperatures studied (we have chosen $t_0 = 1$ s as the reference time). Put into energy units, the classical model overestimates the nucleation barrier at least by $150 k_{\text{B}}T$, and is utterly unable to account for the occurrence of cavitation in the hydrophobic mesopores.



Fig. 4. Normalized potential energy $\Delta\Omega/4\pi\gamma_{l\nu}R_p^2$ of a vapor bubble resting on the wall of a cylinder as a function of its normalized volume V/R_p^3 . The Young's contact angle is $\theta = 120^\circ$. The different plots correspond to different values of the liquid pressure P_L . The shape of the bubble surface is represented for different volumes. At a critical volume V_c , the bubble becomes unstable and spreads over the pore section, forming two disconnected menisci. The maximum value $\Delta\Omega_c$ is equal, or very close, to $\Delta\Omega$ just before instability. (*Inset*) The reduced critical volume V_c/R_p^3 as a function of the reduced liquid pressure $P_L R_p/2\gamma_{l\nu}$.

Why does the classical theory account so well for the critical bubble volume and so badly for its energy? A major breakdown of classical capillarity at the nucleus scale cannot be invoked. A reduced value of the water surface tension as low as $\gamma_{\rm lv} \sim 40$ mN/m could indeed provide the adequate extrusion pressures. However, such a large deviation from the macroscopic value is consistent with neither the $1/R_{\rm p}$ scaling of the intrusion pressure found in the MCM-41 nor the state-of-the-art numerical models (32, 33) or capillary forces investigations (34, 35). Actually, the smallest radius of curvature of water menisci reached in our experiments, obtained for MCM-41 intrusion, is of 3.2 nm, not really a molecular value.

Next, one should consider the role of defects able to reduce the energy barrier. Water-repellent defects are needed to favor the vapor phase. However, a close analysis shows inconsistencies with the intrusion process. Chemical defects should have a very large area: taking a maximum value of $\gamma_{\rm lv}$ for the defect dewetting energy $(\gamma_{\rm sl} - \gamma_{\rm sv})_{\rm def}$, which is unrealistically high, the minimum area *A* providing the needed energy $A\gamma_{\rm lv}(1 + \cos \theta)$, is of the order of 20 nm² in the MCM-41 and more in the SBA-15 (values of θ in Table 1). This result corresponds to extended defects covering a significant cylinder portion. Eq. **4** can then be



Fig. 5. Water repellent defects. (Upper) Local defects. (Lower) Extended defects.



Fig. 6. Comparison of two different methods for calculating the normalized energy barrier $\Delta\Omega_c/4\pi\gamma_{lv}R_p^2$ for vapor nucleation in a cylinder. The different colors correspond to different values of the line tension τ . The Young's contact angle is $\theta = 120^\circ$, and the *x* axis is the normalized liquid pressure. The continuous lines correspond to the first method used in ref. 22, leading to Eq. 6 in the text. The × symbol corresponds to the proper method developed here and described in *SI Text*, section III: the critical nuclei are directly calculated taking into account the finite line tension. The case of MCM41 at 50 °C corresponds to $\tau/\gamma_{lv} = -0.2SR_p$ and has been plotted for illustration. The value of the energy barrier at the extruding pressure $P_L = 178$ bars is 48 k_BT . (*Inset*) A nucleus shape for $\tau/\gamma_{lv} = -0.3R_p$.

used to estimate self-consistently the energy barrier on the extended defect, with a local contact angle θ_{def} . Taking the MCM-41 for instance, $[P_{ext}^{o}(50^{\circ}\text{C}) = 178 \text{ bars})]$, we need a local contact angle $\theta_{def} \sim 135^{\circ}$ to provide the adequate barrier. However, the calculated intrusion pressure on such extended defect should be 750 bars, much higher than the maximum pressure around 500 bars reached in the experiment. Such strong defects should not be wetted, and no nucleation should be needed for emptying the pores hosting them (Fig. 5). A similar inconsistency is obtained for topographic defects, such as bumps on the pore walls and pores constrictions (*SI Text*, section II). Thus, the low nucleation barrier shown by the drying pressure cannot be easily attributed to wall defects in the framework of classical capillarity.

Finally, the effect of long-range interactions can be estimated from the value of the disjoining pressure $A_{slv}/6\pi D^3$, where A_{slv} is the wall-vapor-liquid Hamaker constant, and D is the distance of a meniscus portion to the wall. In the heart of the pore, with a typical Hamaker constant of 10^{-20} J, the disjoining pressure is of the order of 5 bars, which is not relevant. It is more important close to the contact line. This effect is, indeed, described by the thermodynamic concept of line tension introduced by Gibbs more than a century ago to account for the excess energy caused by long-range interactions close to a three-phases contact line (36, 37). With an expected magnitude of the order $\tau \sim \gamma_{\rm lv} a$ (a) being the molecular size, and $\tau \sim 20$ pN for water), line tension plays a significant role only for liquid objects of nanometric size in the three dimensions of space. In contrast to surface tension, it can be negative and thus, reduce the energy of a sessile nanobubble. Experimental determinations of line tension are, however, notoriously difficult, and values reported for water on different substrates vary greatly in amplitude (from 10^{-11} to 10^{-6} N) and sign and tend to depend on the method used (37, 38). The most direct methods, based on the size dependence of the contact angle of sessile drops/bubbles (39), are limited by the difficulty of exact contact angle measurements at the required scale (1-100 nm) (40) and the bias induced by surface heterogeneities (41). In a previous work, Lefevre et al. (22) attributed the low energy barrier for the drying of silanized MCM-41 to a negative line

tension and estimated an amplitude of some 10^{-11} N. In their recent numerical study of the evaporation kinetics of water confined between hydrophobic plates, Sharma and Debenedetti (19) also found nucleation energy governed by the line tension but with a positive value. The solid phases are, however, very different in the two cases: the surfaces in ref. 19 are nonsupported 2D solid phases (a single layer of atoms), which should behave quite differently from a 3D solid phase for long-range interactions. In their systematic numerical study of 3D solid and fluid phases interacting with Lennard–Jones potentials, Weijs et al. (42) found a negative line tension for contact angle values between 70° to 130° (42).

We have studied (with a finite element method) the effect of a line tension τ on the critical nucleus energy. The surprising result is that, although the line tension changes the shape of the critical nucleus, the critical energy is simply given by the expression (22) (Eq. 6)

$$\Delta\Omega_{\rm c} \simeq P_{\rm L}K_1(\theta)R_{\rm p}^3 + \gamma_{\rm lv}K_2(\theta)R_{\rm p}^2 + \tau K_3(\theta)R_{\rm p}, \qquad [6]$$

where $K_1(\theta)$ and $K_2(\theta)$ are the very same functions as in Eq. 4 (case without line tension) and $K_3(\theta)R_p$ is equal to the contact line perimeter of the critical nucleus computed without line tension. This result reflects the almost exact compensation between the line energy gained in changing the shape of the nucleus and the associated losses in volume and surface energies (Figs. 4 and 6).

In contrast to the previous effects, the impact of line tension on the energy barrier is huge: a value of τ order $\gamma_{1v}a \sim 20$ pN changes the energy barrier by hundreds of $k_{\rm B}T$ and the drying pressure by hundreds of bars. Thus, we interpret the high value of the drying pressure in the hydrophobic mesopores as a strong support, if not a direct proof, of a negative line tension of water on the C8-silanized silica. The value of the line tension can then be calculated from the experimental values of $P_{\rm ext}^{\rm o}$ using Eq. **6** and the tabulated values of K_1 , K_2 , and K_3 given in *Materials and Methods* (Eq. 7):

$$P_{\text{ext}}^{o} = \frac{k_B T}{V_{\text{c}}} \ln \frac{L \nu t_{\text{o}}}{b} - \frac{K_2(\theta)}{K_1(\theta)} \frac{\gamma_{lv}}{R_{\text{p}}} - \frac{\tau}{R_{\text{p}}^2} \frac{K_3(\theta)}{K_1(\theta)}.$$
 [7]

The result is shown in Fig. 7 and Table 1. The precision of ± 2 bars on the drying pressure gives a relative precision better than 10^{-2} on τ . This is a much higher resolution than the one provided by currently available experimental methods. The values



Fig. 7. The reference extrusion pressure P_{ext}^{o} as a function of temperature in the three materials. The dashed line is the best fit obtained by fitting the line tension in Eq. 6. The pore radii, contact angles, water surface tension, and expressions of K_1 , K_2 , and K_3 are listed in *SI Text*, section I. In each material, we allow a small linear variation $\tau(T) = \tau_o(1 + \alpha(T - T_o))$. The thermal coefficient α is of the order -10^{-3} K⁻¹ (Table 1). For comparison, the thermal coefficient of water surface tension in the same range is $-2.4 \ 10^{-3}$ K⁻¹.

quoted in Table 1 are calculated using the microscopic quantities b = 1 Å, $\nu = 10^{12} \text{ s}^{-1}$, and $L = 10 \,\mu\text{m}$. Changing the ratio $b/\nu L$ by a factor 10 or 0.1 changes τ by less than $\pm 0.6 \text{ pN}$ (that is, 1 to 2.5%). The line tension found has consistent values for the three materials ranging from -23 pN in MCM-41 to -35 pN in SBA-15. To account fully for the temperature variation of P_{ext}^{0} , we have allowed a small thermal variation $\tau(T) = \tau_0[1 + \alpha(T - T_0)]$ (Fig. 7 and Table 1). The absolute value $|\alpha|$ of the thermal coefficient is less than 10^{-3} K^{-1} , to be compared with the thermal coefficient of water surface tension in the same range, $-2.4 \, 10^{-3} \text{ K}^{-1}$.

The values that we find here are of the same amplitude but opposite sign to the line tension of water measured on hydrophilic surfaces, such as quartz (43) and silica (39). On strongly hydrophobic surfaces, measurements have focused on the shape of nanobubbles. State-of-the-art investigations have not evidenced a variation of the contact angle with the bubble size (40, 44, 45), corresponding to an upper amplitude of about 100 pN for the line tension of water. Therefore, we compare our results with the systematic study by Weijs et al. (42) performed for Lennard-Jones fluids, although the water/silane/silica system investigated here is chemically different. The (negative) line tension is characterized by the ratio of the tension length $l = -\tau/\gamma_{\rm lv}$ to the molecular size a. The ratios found here (with a = 2.7 Å) are from 1.2 to 1.9, which is in good agreement with thermodynamic expectations and in qualitative agreement with the work by Weijs et al. (42) $(l/a = 0.8\hat{2}$ at contact angle 117°).

Finally, we find a difference of about -20% (respectively, +20%) for the line tension in MCM-41 (respectively, SBA-15) with respect to HMS. This difference could be because of the topography of the pore walls (SBA-15 is known to have a rougher surface than MCM-41) or a systematic trend with the solid surface curvature, such as described in ref. 46.

Conclusion

In summary, we have shown here that the drying of water brought into metastable equilibrium inside hydrophobic cavities is a dynamical process with slow logarithmic dynamics. This finding illustrates the importance of carrying out dynamical ratedependent study of adsorption and desorption phenomena. Such studies are a sensitive probe of the specific mechanisms that control the formation and dynamics of nanobubbles on hydrophobic surfaces. Here, we find that the mechanism that allows one to interpret quantitatively the rate dependence of the extrusion pressure is the thermally activated appearance of a critical vapor nucleus that can be well-described by the macroscopic theory of capillarity, if a negative line tension of water is taken into account. Our approach provides an accurate independent estimate of this line tension, which is consistent with what can be

Table 1. Nucleation volume and line tension in the various materials

	MCM-41	SBA-15	HMS
R _p *	1.34 ± 0.1	2.16 ± 0.25	1.54 ± 0.1
θ _{50°C} [†]	114.8°	119.1°	115°
<i>δθ</i> (°/°C ⁻¹)	0.036	0.077	0.059
V_c^{\dagger}	10.2 ± 1.5	51 ± 17	17.8 ± 2.7
P_{ext}^{o}(50^{o}C)^{§}	178	60.2*	109.4
τ _{50 °C} ¶	-23.3	-35.5	-30.1
α(°C ⁻¹)	-1.0 10 ⁻³	-0.8 10 ⁻³	-0.7 10 ⁻³

*Pore size (nm).

[†]Intruding contact angle at 50 °C; $\theta(T) = \theta_{50 \text{ °C}} + \delta\theta(T - 50)$. [‡]Nucleation volume (nm³).

Nucleation volume (nm).

[§]Reference extrusion pressure at 50 °C (bar; $t_o = 1$ s). [¶]Line tension at 50 °C (10⁻¹² N); $\tau(T)/\tau_{50} \circ_{C} = 1 + \alpha(T - 50)$.

Table 2. Other characteristics of the materials

	MCM-41	SBA-15	HMS
R _{BJH} – R _{BDB} (nm)	1.25 – 1.43	1.91 – 2.40	1.44 – 1.64
R _p (nm)	1.34 ± 0.1	2.15 ± 0.25	1.54 ± 0.1
P _{int} (50 °C; bar)	432.1	291.1	325.5
θ _{50 °C}	114.8°	119.1°	115°
δθ(°/°C ^{−1})	0.036	0.077	0.059
K ₁ (θ _{50 °C}) − K ₂ (θ _{50 °C})	4.27 – 3.24	4.28 – 2.614	4.27 – 3.21
K ₃ (θ _{50 °C})	12.366	12.422	12.37

inferred from simulation data or atomic force measurements of droplet shapes. The existence of a negative line tension is an important ingredient for understanding the very high stability of nanobubbles on hydrophobic surfaces and the vaporization of water from repellent cavities, and it has wide implications for heterogeneous cavitation, hydrophobic interactions in biological matter, and more generally, properties of very small liquid objects. Additional work is under progress to study the compressibility of the water/silane hydrophobic interface in the mesopores (47).

Materials and Methods

The MCM-41 was synthesized from the structuring agent octadecyltrimethylammonium bromide (C18NMe3Br) as described in ref. 27. MCM-41 is a model material presenting independent cylindrical pores that are hexagonally ordered and have a smooth internal surface. SBA-15 was synthesized from the triblock copolymer poly(ethylene oxide)-poly(propylene oxide)-poly(ethylene oxide) EO20PO70EO20 as a structuring agent under acidic medium at 60 °C for 24 h (29). SBA-15 also has a hexagonal arrangement of cylindrical pores, but depending on the synthesis temperature, these mesopores can be connected by a secondary network of smaller micropores. We chose a temperature of 60 °C, which prevents the micropores from growing and creating interconnections (29). The HMS was prepared with $\mathsf{C}_{16}\mathsf{NH}_2$ as the structuring agent, with a ratio EtOH/H₂O = 0.19 at an ambient temperature for 24 h (31). The HMS has a smooth internal surface but is less ordered than MCM-41, and the pore network can be randomly connected in few locations. The three mesoporous silicas are silanized by grafting chlorodimethyloctylsilane as described in ref. 48. Before and after silanization, the MTS was characterized by nitrogen adsorption at 77 K (49). The pore size determination was done using two methods: the Barret–Joyner–Halenda method (50), which is classically used but has been shown to underestimate the pore size of hydrophilic silica (49), and the Broekhoff-de Boer method. The pore size of the hydrophobic materials is taken as the average of the two results $R_{p} = (R_{BdB} +$ R_{BJH} /2, and the uncertainty is taken as their difference $\Delta R_{\text{p}} = (R_{\text{BdB}} - R_{\text{BJH}})/2$. The values are gathered in Table 2.

The experimental device, cell preparation, and obtention of the P-V isotherms of water in the hydrophobic materials are detailed in ref. 25. The intrusion (respectively, drying) pressure is defined as the average pressure value on the corresponding plateau. The intrusion pressure depends weakly on the intrusion rate, and we use quasistatic values obtained at a rate of 0.1 s⁻¹. It also depends very weakly on temperature, because $P_{int} = P_{int}(T_o) + \Delta P_{int,T}$ $(T - T_o)$. The Young contact angle derived from Eq. 1 also changes with temperature, because $\theta = \theta(T_o) + \delta\theta(T - T_o)$. Values for the three materials are summarized in Table 2.

In the data analysis, we use the following expressions to interpolate the functions K_1 , K_2 , and K_3 with 90° $\leq \theta \leq 135$ °:

 $K_1(\theta) = 4.1661 + 0.11242 \times \sin(0.11819 \times \theta + 0.16478),$

 $K_2(\theta) = 20.32 - 0.14879 \times \theta$, and

 $K_3(\theta) = 3.355 + 0.14136 \times \theta - 0.00054762 \times \theta^2$.

The theoretical calculation of the shape and energy of a vapor bubble in a cylinder in the presence of a line tension is performed with a finite element method and a relaxation algorithm.

ACKNOWLEDGMENTS. We thank Thierry Abensur and the European Aeronautic Defence and Space-Astrium Space Transportation Companies for supporting this research.

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