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Mesoporous Silica Colloids: Wetting, Surface Diffusion and Cationic Surfactant Adsorption

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ABSTRACT

We have investigated the wetting and surface diffusion of mesoporous colloidal silica particles at the water surface; and the adsorption of cationic cetyltrimethylammonium (CTA+) surfactant on these particles. Porous silica colloids diffuse at the surface of water and in the volume, interacting with cationic surfactants that can adsorb inside the pores of the particles. We observed that surfactant adsorption on mesoporous silica depends dramatically not only on the particle pore size but also on specific counterion effects. We measured striking differences both on a macroscopic property of the interface, i.e. surface tension, and also at a single particle level by evaluating the translational diffusion of partially wetted particles at the fluid interface. We varied the pore size from 2 to 7 nm and explored the effects of ions owing different hydration number and kosmotropic/chaotropic character. At concentrations lower than the critical micellar concentration, we evidence that cationic surfactants adsorb on silica as surface micelles and surfactant adsorption inside the pores occurs only if the pore diameter is larger than the size of surface micelles. With a view to understand the surprising different adsorption behavior of CTA+OH− and CTA+Br− on porous silica particles, we investigated the effect of counterions on the surfactant adsorption on porous silica colloids by tuning the pH and the counterion properties.

INTRODUCTION

Cationic surfactants are widely used in many industrial and fundamental research fields ranging from cosmetics, fabrics, nanoparticles synthesis, DNA folding, self-assembly and mesoporous material fabrication.1,2 Besides these applications, the fate of cationic surfactants and the resulting waste in water is an important issue, since cationic surfactant contamination of water represents a real environmental problem.3 Current methods to remove surfactants from water involve the use of organophilic clays, flocculants or carbon beds.4–8 In many separation methods, surfactant removal by adsorption onto solid substrates is required. Hence, both large surface areas and an effective attraction between the surfactant and the solid are necessary to drive soluble surfactants adsorption on the solid boundaries.

Adsorption of cationic surfactants on planar solid surfaces has been investigated for several surfactant-substrate systems. For cetyltrimethylammonium (CTA+) surfactants on silica surfaces, different adsorption regimes are found as a function of the surfactant concentration, surface charge and
hydrophobic interaction. At low concentrations, surfactant may adsorb onto the silica surface as a monolayer, as defective bilayers or as surface micelles. These micellar structures are flattened by the surface and possess a thickness of 3.5 nm and a lateral size of 9 nm. Upon increasing the concentration, bilayers may form due to the interaction between the hydrophobic chains. It is important to notice that the interfacial self-assembled structures usually appear at a concentration lower than the usual critical micellar concentration (CMC) in the volume. For CTA\textsuperscript{+} surfactants at concentrations C lower but close to CMC, surface excess concentrations \( \Gamma \) between \( \approx 1 \) and \( \approx 2 \, \text{mg/m}^2 \) have been reported.

As pointed out before, solid substrates with large specific areas are required for adsorption in order to separate surfactants from clean water. In this context, porous particles are ideal candidates given the large surface/volume ratio, which can be tuned by the pore size and volume fraction. However, adsorption inside a pore strongly differs from the adsorption on a planar interface. Pore size and morphology affect the adsorption process, which depends on differences in pressure and physicochemical properties inside and outside the pores.

Non-ionic surfactants adsorption onto mesoporous silica particles has been widely investigated by Findenegg and coworkers. Bilayers formation and other structures inside the pores has been reported. Very few experimental investigations can be found in the literature for ionic surfactants onto mesoporous silica. A small angle neutron scattering investigation reported some preliminary experiments for 8 nm pore size mesoporous silica at pH=9 where an increase of cationic surfactant adsorption occurs if the ionic strength is increased. Cationic surfactant interacting with silica nanoparticles have been also recently investigated.

Here, we have investigated the wetting and diffusion of porous silica colloids, and the adsorption of cationic surfactants on porous colloidal particles. Porous colloids of different pore sizes are deposited both in the bulk and at the interface. By varying pore size, surfactant concentration and investigating counterion effects we aim at controlling CTA\textsuperscript{+} cationic surfactant adsorption. These results could be used in order to develop a strategy to clean the surface and the volume of water with micron sized particles, which are small enough to perform Brownian motion and rapidly diffuse at the interface (and in the volume) but also large enough to slowly sediment. These porous colloids may be ideal candidates for an efficient removal of surfactants at low concentrations and present the advantage to be easily removed after decontamination by sedimentation or flotation. Note that larger particles are difficult to adsorb at the interface and may sediment very fast; whilst nanoparticles may remain dispersed in water due to the strong Brownian motion and cannot be separated by gravity or adsorption on bubbles as in flotation.

**MATERIALS AND METHODS**

*Materials.* All chemicals, except for the in-house synthetized colloids, were purchased from Sigma Aldrich and used without further purification. Two surfactants were investigated: cetyltrimethylammonium bromide CTA\textsuperscript{+}Br\textsuperscript{−} (CAS 57-09-0, molecular weight \( M_w = 364.45 \, \text{g/mol} \)) and cetyltrimethylammonium hydroxide CTA\textsuperscript{+}OH\textsuperscript{−} (10 wt.% in H\textsubscript{2}O, CAS 505-86-2, \( M_w = 301.55 \, \text{g/mol} \)). Three different batches of porous silica particles were also employed. The first two consist of spherical mesoporous silica colloids (CAS 7631-86-9) with an average radius \( \bar{R} = 1.23 \pm 0.10 \, \mu \text{m} \) (measured by scanning electron microscopy, SEM) and nominal pore diameter \( d = 2 \, \text{nm} \) and \( d = 4 \, \text{nm} \) and a specific area \( S_A = 1000 \pm 100 \, \text{m}^2/\text{g} \) for \( d = 2 \, \text{nm} \) and \( S_A = 350 \pm 50 \, \text{m}^2/\text{g} \) for \( d = 4 \, \text{nm} \). A third batch of non-spherical mesoporous silica particles was synthetized in the laboratory. These last particles are currently
called Santa Barbara amorphous-15 silica (SBA-15). The average pore diameter is larger than the 
previous ones: \( d = 7 \text{ nm} \) measured by volumetric nitrogen sorption isotherms. Stable particle dispersions 
were obtained by sonication of 1.2 mg/mL dispersions in an ultrasound bath for one hour. We measured 
the Feret diameter\(^2\) of these particles by scanning electron microscopy, which leads to an equivalent 
radius \( \bar{R} = 1.96 \pm 0.97 \mu \text{m} \) (\( i.e. \) half of the Feret diameter). The pores are about a micron long and 
connected via mesoporous walls. A total specific area of \( 750 \pm 50 \text{ m}^2/\text{g} \) was evaluated by Brunauer-
Emmett-Teller (BET) method. Pore size distributions were determined by the (Barrett, Joyner, Halend) 
BJH method.\(^2\) Mesoporous walls contribute significantly to the total specific area in terms of 
microporosity (size < 1-2 nm). We evaluate a total specific area \( S_A = 225 \pm 25 \text{ m}^2/\text{g} \) for the \( d = 7 \text{ nm} \) pore 
size (See Figure S1 in Supporting Information, SI).

NaOH, NaBr, NaCl salts were used as received. Fresh Milli-Q water was used throughout this work. 
Experiments were carried out at room temperature \( T = 22 \degree \text{C} \).

**Surface tension and sessile drop contact angle.** Surface tension was used to measure surfactant adsorption 
on porous particles. All measurements were performed in a 9 cm diameter polystyrene Petri dish 
containing a volume \( V = 35 \text{ mL} \) of surfactant solutions at different concentrations \( C \) and a fixed mass (1.2 
mg) of porous particles. For planar aqueous surfaces contained in a Petri dish, surface tension was 
measured by the Wilhelmy plate method (KSV NIMA, Biolin Scientific) using a filer paper plate. In order 
to evaluate the interfacial tensions of the silica-surfactant solutions-gas systems, we also measured the 
surface tension of aqueous solutions and the advancing contact angle \( \theta \) of sessile drops by a profile 
analysis tensiometer (PAT, Sinterface, Germany). A typical volume of the aqueous drop in PAT 
experiments is 15 mm\(^3\). Series of experiments were repeated at least three times and show a good 
reproducibility. For interfacial tension measurements, the standard deviations of the data obtained in 
different experimental campaigns lay between 1 and 3 mN/m, which can be related to the experimental 
protocols: particle deposition, aging of particle dispersions and surfactant solutions (\( i.e. \) maximum 5 
days).

**Particle dispersion deposition.** The final chosen method adopted to deposit porous particles at the 
interface is to drop 1 mL of a 1.2 mg/mL particle suspension on a tilted glass slide placed at the air-water 
interface and let it slip.\(^2\) Particle density at the interface can be calculated by counting the particle 
number using optical microscopy. For all porous particle systems, we evaluated an average surface 
coverage of 6.5±2 particles per mm\(^2\) (if 1.2 mg of silica porous particles are added in the Petri dish). From 
the latter value, one can estimate the ratio between the total number of particles \( N_{\text{tot}} \) and the ones trapped 
at the surface \( N_s \): \( N_{\text{tot}}/N_s \approx 10^6 \), which is very high, meaning that only few particles stay at the interface. 
We have put many efforts to find protocols to increase the number of particles remaining at the air-
aqueous solution interface and try several methods. However, protocols involving spraying or the use of 
additional volatile solvents were discarded since they do not allow the control of the particles numbers or 
introduce additional sources of contamination and uncertainties. It is important to note that even if \( N_{\text{tot}}/N_s \) 
\( \approx 10^6 \), the amount of particles at the interface is non-negligible for the adsorption of soluble surfactants, 
which accumulate preferentially at the interface but they are also present in the bulk, see Figure S2 in SI.

**Particle contact angle by gel trapping.** We measured the particle contact angle of porous silica colloids 
using a gel trapping technique.\(^3\) Few microliters of a diluted solution of particles were added to a gelled 
solution of Phytageal (CAS 71010-52-1, Sigma Aldrich) in a small container. After heating up the gel to 
the liquid state, the particles get trapped at the interface. Going back to the room temperature, a UV-glue 
was used to transfer the particles from the gel to the glue substrate. The contact angle was measured using
scanning electron microscopy images of the transferred particles on the solidified glue at their complementary contact angle positions.

Particle tracking. Bright field optical microscopy was used for particle tracking. Experiments were performed in an upright Leica optical microscope mounted on an anti-vibration table. Images of isolated particles were recorded using a CMOS camera (Orca Flash 4.0, Hamamatsu) at rates of 80 or 100 frames per second using objectives of different magnifications (×63 and ×100). Tracking of the particle location was done by an IDL routine or by using an image correlation-based approach ("Stat Tracker St. Andrews") implemented in Labview (National Instruments).

Zeta potential and pH measurements. We measured the zeta potential ζ (Zetasizer Nano ZS, Malvern Instruments) and the pH of the particle dispersions in pure water or in the presence of surfactants. For porous silica colloids dispersions, a pH = 6.4 was always measured in pure water.

RESULTS AND DISCUSSION

Interfacial tensions of CTA⁺Br⁻ and CTA⁺OH⁻ at the liquid-gas and solid-liquid interfaces

We started characterizing the adsorption of CTA⁺ surfactants on non-porous silica by evaluating the interfacial tension between silica and the aqueous surfactant solutions, γSL. Young equation for a sessile drop of aqueous surfactant solutions on a silica substrate in air reads:

\[
γ_{SL} = γ_{SG} + γ_{LG} \cos θ,
\]

where the subscripts S, L and G represent solid, liquid and gas respectively. One can evaluate γSL measuring the interfacial tension γLG, the sessile drop static contact angle θ and assuming that γSG is a constant (and does not vary with the surfactant concentration). Surface tension at the air-aqueous solution interface γLG for CTA⁺Br⁻ and CTA⁺OH⁻ surfactant was measured by the Wilhelmy plate technique and pendant drop tensiometry. The two methods lead to very similar results as shown for a series of experiments in Figure 1A. The evolution of γLG as a function of the surfactant concentration C for both surfactants shows no significant difference. In addition, γLG values are stable and do not change in a typical experimental time range of 3000 s (inset of Figure 1A). Sessile drops made with surfactants solutions were deposited on clean glass slides and their contact angles θ were also evaluated by imaging the drop profile (and the cosθ is plotted in the inset of Figure 1B for a series of experiment). Hence, γSL can be calculated from eq. 1 with γSG = 68.5 mN/m as reported by Binks et al. for silica surfaces showing water contact angle similar to our experiments. Unexpectedly the silica-aqueous solution interfacial tension γSL shows a non-monotonic behavior if the surfactant concentration is increased. For C ≥ 0.1 mM, γSL increases if C increases, instead of decreasing as γLG. Hence, the Gibbs adsorption valid for ideal dilute solutions for C < CMC (≈ 0.8-0.9 mM) can be applied only for the air-liquid interface, where a surface concentration Γ ≈ 1 ng/m² at the CMC can be evaluated. For C < 0.1 mM at the solid-liquid interface, one could also observe a tiny decrease of the interfacial tension which could correspond to an increasing accumulation of non-interacting surfactants at the solid-liquid interface. However for C ≥ 0.1 mM, the increase of γSL points to the formation of self-assembled structures as bilayers or surface micelles, which form because of the hydrophobic interactions between surfactant tails. Note that the
concentration at which the interfacial tension $\gamma_{SL}$ increases is about 10 times lower than the CMC. An increase of the interfacial tension $\gamma_{SL}$ reveals that the surface is increasing in hydrophobicity due to the presence of hydrophobic surfactant groups. As already pointed out in the Introduction, the presence of bilayers or surface micelles on planar silica surfaces has been widely reported but to the best of our knowledge was never connected to a minimum in the interfacial tension $\gamma_{SL}$.

Figure 1. (A) Liquid-gas interfacial tension of CTA$^+$Br$^-$ (+, □) and CTA$^+$OH$^-$ (−, ▢) as a function of the concentration obtained by pendant drop tensiometry (+, −) and by Wilhelmy plate method (□, ▢); inset: interfacial tension as a function of time for pure water (>v) and CTA$^+$Br$^-$ (+) and CTA$^+$OH$^-$ (−) at $C = 0.5$ mM. (B) Solid-liquid
interfacial tension CTA\(^{-}\)Br\(^{+}\) and CTA\(^{-}\)OH\(^{-}\) evaluated by the Young equation and advancing contact angle measurement of sessile drops on silica (inset). Data for pure water (\(\forall\)) are also shown.

**Porous particles at the interface**

The contact angle of porous particles at the gas-liquid interface was measured by a gel trapping method (see Materials and Methods). Figure 2 shows images of particles at their complementary contact angle positions. We have also observed some crater shape footprints on the solidified substrate, which correspond to particles possessing very low contact angles. For spherical porous particle, we measured particle contact angle \(\alpha = 47° \pm 6°\) for \(d = 2\) nm and \(\alpha = 52° \pm 14°\) for \(d = 4\) nm (Figure 2A and B). Non-spherical porous particles show also a finite immersion at the air-water interface that is comparable to the contact angle observed for the spherical porous particles.

![Figure 2](image_url) SEM images of gel trapped \(R = 1.23\) \(\mu\)m spherical mesoporous silica colloids \(d = 2\) nm (A) and \(d = 4\) nm (B); and a non-spherical silica particle of equivalent radius \(R = 1.96\) \(\mu\)m (diameter = 3.92 \(\mu\)m) and \(d = 7\) nm (C).

It is important to notice that the measured contact angle of a sessile water drop on silica, \(\theta \approx 26°\), is significantly lower than the contact angles of porous silica particles shown in Figure 2: \(\alpha > \theta\). In order to explain this result, we start discussing the partial wetting state of porous particles and compare it to the reference case of a bare solid spherical particle, see Figure 3. For a bare particle, the equilibrium of interfacial tensions leads to the Young equation (see equation 1). For smooth and homogenous surfaces, the bare solid micron-sized particle contact angle should be identical to the contact angle of a sessile liquid drop on a flat solid substrate: \(\alpha_B = \theta\).\(^{39}\) In the framework of the Cassie-Baxter model valid for composite interfaces, two different scenario can be considered for porous particles at the interface.

In the first case, we consider that the pores of the particles are completely filled by the liquid. In this case, the particle interfacial tensions \(\gamma_{SG}\) and \(\gamma_{SL}\) change if compared to the bare particle case. The interfacial tension of the particle exposed to the liquid will be reduced from \(\gamma_{SL}\) to \(f\gamma_{SL}\), where \(f\) is the solid area fraction on the particle surface, see Figure 3. The particle surface exposed to the gas phase can be regarded as a composite interface made of solid and liquid.\(^{39}\) The resulting equilibrium of interfacial tensions reads:

\[
f\gamma_{SG} + (1-f) \gamma_{LG} = f\gamma_{SL} + \gamma_{LG}\cos\alpha.
\]  

Except for \(f\), all the other variables in equation 2 are measured. In pure water, \(\gamma_{SG} = 68.5\) mN/m, \(\gamma_{LG} = 72.5\) mN/m and \(\gamma_{SL} = 4\) mN/m (Figure 1). According to equation 2, \(\alpha\) decreases if \(f\) decreases and the Young equation is recovered if \(f = 1\). Actually for porous particles with pores completely filled by the liquid, the particle contact angle \(\alpha\) is expected to be low, \(\alpha < \theta\), which explains the observation of crater footprints in SEM images and the large number of particles in the volume with respect to the particles present at the interface (see Particle dispersion deposition section). Note also that the interfacial energy...
gain associated to the adsorption of a bare particle at the interface $\gamma_{LG}A_0$ due to the removal of a bare fluid interface area ($A_0 = \pi R^2 \sin^2 \alpha$) is strongly reduced for a porous particle given the presence of liquid inside the particle. A second scenario can be considered to explain the results shown in Figure 2, where $\alpha > \theta = \alpha_B$ (Figure 3). In this case, the liquid is not present on the porous particle surface exposed to the gas phase. Hence, the external surface can be regarded as a composite surface made of solid and air, and the interfacial tension of this surface becomes $f\gamma_{SG}$. The resulting equilibrium of interfacial tensions reads:

$$f\gamma_{SG} = f\gamma_{SL} + \gamma_{LG} \cos \alpha.$$  \hspace{1cm} (3)

In this second case, the contact angle of a porous particle can be higher than the contact angle of an equivalent bare particles. Equation 3 can be rewritten in the form $\cos \alpha = f \frac{(\gamma_{SG} - \gamma_{SL})}{\gamma_{LG}} = f \cos \theta$ (see equation 1), which leads to solid area fraction $f = 0.76$ for 2 nm and $f = 0.68$ for 4 nm pore size porous particles.

![Sketch of Interfaces](image)

Figure 3. Sketches of a bare non-porous particle (left), a porous particle with pores completely filled by the liquid (center) and a porous particle with pores partially filled by the liquid (right) in mechanical equilibrium at the gas-liquid interface.

**Influence of particle pore size on surfactant adsorption**

In order to measure the surfactant adsorption on porous silica colloids we performed surface tension experiments using a Wilhelmy plate apparatus. Starting from a stable interfacial tension measurement at a given surfactant concentration $C$, porous colloids are deposited at the air-aqueous solution interface via a tilted glass slide. Note that the addition of few porous colloids (1.2 g in 35 mL water) in absence of surfactants does not affect the interfacial tension $\gamma_{LG}$. As a consequence of the addition of mesoporous particles, we record an increase of the liquid-gas interfacial tension connected to a surfactant removal from the aqueous solution due to the adsorption on porous silica colloids.

In Figure 4 we show a series of surface tension experiments of CTA$^+$Br$^-$ and CTA$^+$OH$^-$ solutions at different surfactant concentrations in the presence of a fixed number of porous silica particles of 4 and 7 nm pore diameters $d$. For CTA$^+$OH$^-$ in the presence of the $d = 4$ nm (Figure 4A) and 7 nm (Figure 4B) porous silica particles, we measured a significant increase in the surface tension, which is a clear evidence.
of surfactant removal from the aqueous solution. When adding \( d = 4 \) nm porous silica colloids, the interfacial tension increases by 8 mN/m, 5 mN/m and 4 mN/m for 0.25 mM, 0.5 mM and 0.7 mM respectively. As for the \( d = 7 \) nm porous silica colloids, the surface tension increases by 12 mN/m, 9 mN/m and 7 mN/m for 0.25 mM, 0.5 mM and 0.7 mM respectively. Surprisingly, no apparent increase in the surface tension with CTA\(^{+}\)Br\(^{-}\) was observed, which will be further analyzed and discussed in the following sections. No adsorption for both CTA\(^{+}\)Br\(^{-}\) and CTA\(^{+}\)OH\(^{-}\) was also observed for \( d = 2 \) nm porous silica colloids, see Figure 5. Surface tension measurements showed no variation before and after adding the \( d = 2 \) nm porous particles (Figure 5), leading to the conclusion that a pore size of 2 nm is too small for CTA\(^{+}\) cationic surfactant adsorption. This result agrees with the increase of \( \gamma_{SL} \) shown in Figure 1B and connected to the formation of surface micellar structures at concentrations much lower than the CMC. Surfactants may adsorb inside the pores as micellar structures and not as single molecules. The size of micelles in the bulk\(^{10}\) (for \( C \approx \text{CMC} \)) is between 2.6 and 4.7 nm; and for \( C < \text{CMC} \) the thickness of surface micelles is about 3.5 nm for CTA\(^{+}\)Br\(^{-}\). Hence, these dimensions are always larger than the size of 2 nm and explain why adsorption is not observed in \( d = 2 \) nm porous particles.
Figure 4. Liquid-gas interfacial tension as a function of time after the deposition of \( d = 4 \) nm porous silica particles (A) and \( d = 7 \) nm particles (B) at different CTA\(^{+}\)OH\(^{-}\) surfactant concentrations: \( C = 0.25 \) mM (.), \( 0.5 \) mM (!), \( 0.7 \) mM (?); and CTA\(^{+}\)Br\(^{-}\) concentration \( C = 0.7 \) mM (–). 1.2 g of particles were added in 35 mL solution.
Figure 5. Liquid-gas interfacial tension as a function of time for CTA^\text{+}OH^- at C = 0.5 mM in the presence of $d = 7$ nm (.), $d = 4$ nm (!) and $d = 2$ nm (7) porous silica particles (1.2 g of particles in 35 mL).

Knowing the initial CTA^\text{+}OH^- surfactant concentration $C$ and the interfacial tension $\gamma_{LG}$ after adsorption (at time $t = 3000$ s in Figure 4 and 5), we can extract the final surfactant concentration $C_f$ in the solution using the $\gamma_{LG}$ vs $C$ data (eventually interpolated) shown in Figure 1A. Hence, we can calculate the total mass of surfactant removed in the $V = 35$ mL solution as $(C - C_f)V$. In Figure 6A, the adsorption amount (which is the mass of surfactants divided by the mass of porous particles) is plotted as a function of the initial surfactant concentration. For $d = 7$ nm porous silica particles, the adsorption amount increases up to 2 mg/g at intermediate concentrations. For $d = 4$ nm porous silica particles instead a maximum in the adsorption amount is clearly observed at $C = 0.25$ mM, above which the adsorption amount decreases.

Knowing the mass $m$ (=1.2 mg) of the porous particles and their specific areas $S_A$ (see Materials and Methods), we can also calculate the surfactant surface concentration $\Gamma_S$ on the total silica area $(= mS_A)$:

$$\Gamma_S = \frac{(C - C_f)V}{mS_A} \quad (4)$$

Figure 6B shows $\Gamma_S$ as a function of CTA^\text{+}OH^- concentration for $d = 4$ and 7 nm porous silica particles. For $C = 0.1$ mM, we measured a weak adsorption of surfactants on porous silica particles $\Gamma_S = 1$-3 mg/m$^2$, which is equivalent to the surface excess of CTA^\text{+} surfactant forming dense monolayers at the interface. For non-spherical silica particles with $d = 7$ nm, $\Gamma_S$ increases up to 10 mg/m$^2$ at intermediate concentrations and slightly decreases if $C$ approaches the CMC ($\approx 0.9$ mM). For spherical silica particles with $d = 4$ nm, the surface concentration increases up to $\Gamma_S \approx 4$ mg/m$^2$ and significantly decreases if $C$ approaches the CMC. It is interesting to note that at low concentrations, $\Gamma_S$ is proportional to the pore size, meaning that pores are filled with surfactants. In this pore size range and for surfactant concentrations far from the CMC, the larger the pore diameter the larger the surfactant adsorption. Close to the CMC however surfactants start to self-assemble in the form of micelles in the volume. As a consequence, a competition between the self-assembly of surfactants in the volume and the adsorption inside the pores of the silica particles is expected. This competition may also depend on the confinement...
dictated by the pore size. If the pore confinement effect is very strong, micelles in the volume could hinder surfactant adsorption inside the particles’ pores. For \( d = 4 \) nm porous particles, the decrease of \( \Gamma_S \) as a function of concentration seems in agreement with this scenario. For pore size \( d = 7 \) nm, the pore confinement effect is weaker than for \( d = 4 \) nm given that the micellar dimensions are about 3.5 nm, i.e. the half of the pore size, which explains the weak decrease of \( \Gamma_S \) as a function of concentration close to the CMC, see Figure 6B.
Figure 6. Adsorption amount (A) and surfactant excess concentration on the total silica Area (B) for \(d = 7\) nm (−), and \(d = 4\) nm (I) solid particles as a function of CTA^+OH^- initial concentration.

**Porous particle diffusion at the gas-liquid interface**

At the single colloidal particle level, we have also noticed striking differences in the translational diffusion of spherical \(d = 4\) nm porous particle at the gas-liquid interface in the absence and the presence of CTA^+ surfactants. Tracking the particle center of mass at the air-water interface, we observed Brownian trajectories (see inset Figure 7A) and measured the mean squared displacement (MSD), which is plotted as a function of the lag time \(\Delta t\) in Figure 7A. Translational diffusion coefficients at the surface, \(D_s\), can be calculated by \(MSD = 4D_s\Delta t\).

In absence of surfactants, \(D_s\) for \(d = 2\) and \(4\) nm pore particles agrees with the hydrodynamic prediction for partially immersed particles at clean and flat fluid interface, see Figure 7B:\(^{40}\)

\[
D_s = \frac{k_B T}{k_B \eta R},
\]

where \(k_B T\) is the thermal agitation energy, \(\eta\) is the liquid viscosity and \(k_B\) is the drag factor accounting for the particle immersion or contact angle at the interface:\(^{40}\)

\[
k_t = 3\pi \left[1 + \frac{9}{16} \cos \alpha + O(\cos^2 \alpha)\right] \text{ (for } 0 < \alpha < 90^\circ).\]

For the particle diffusing in the bulk our experimental results agree with the Stokes-Brownian diffusion

\[
D_0 = \frac{k_B T}{6\pi \eta R} = 0.174 \mu m^2/s.\text{ In absence of surfactants, the good agreement between the } 1.4 < D_s/D_0 < 1.5\text{ experiments and the prediction shows that no additional dissipation due to contact line fluctuations (as observed for bare silica colloids with } 1.2 < D_s/D_0 < 1.3)\text{ occur on partially wetted porous particles. As pointed out before the external porous particle surface can be regarded as a composite surface made of hydrophilic silica and fluids. Hence, this surface presents less defects than the surface of a bare particle, which may explain the negligible line friction due to surface defects.}\(^{41}\) As a perspective, one may wonder if these porous particles could be good probes for passive surface microrheology since the only sources of dissipations seem to be the viscosity.

Surface diffusion coefficients of porous particles change significantly in the presence of CTA^+OH^- (\(C = 0.5\) mM): \(1.1 < D_s/D_0 < 1.2\), and in CTA^+Br^- (\(C = 0.5\) mM) aqueous solutions: \(D_s/D_0 \approx 0.3\), see Figure 7. The experiments are performed at times \(\approx 3000\) s, where an increase of the gas-liquid interfacial tension of 5 mN/m for CTA^+OH^-; and no change for CTA^+Br^- were measured, see Figure 4. Hence the presence of surfactants affect strongly the porous particle translational diffusion at the surface. For porous particles in the presence of CTA^+OH^-, adsorption of surfactants inside the pores occurs during the porous particle Brownian motion, whilst in the presence of CTA^+Br^- no adsorption dynamics occurs. In order to quantify the decrease of the surface translational diffusion \(D_s\), we use a model describing the drag felt by a spherical particle straddling a viscous and incompressible interface possessing a surface viscosity \(\eta_s\) at low Boussinesq numbers \(B_0 = \frac{\eta_s}{\eta R} < 1\). The drag factor in equation 5 in this case reads:\(^{42}\)

\[
k_t = k_t^{(0)} + B_0 k_t^{(1)} + O(B_0^2),
\]

\[
k_t^{(0)} \approx 6\pi \sqrt{\tanh[32(1 + \cos \alpha)/(9\pi^2)]}
\]

\[
k_t^{(1)} \approx -4\ln\left[\frac{2}{\pi} \arctan\left(\frac{(1 + \cos \alpha)}{3}\right)\right]
\]
For CTA\textsuperscript{+}Br\textsuperscript{−} (C = 0.5 mM) aqueous solutions, a surface viscosity $\eta_s \approx 10^{-8}$ Pa.s.m fits well the experimental result. In the literature, a large range of values of $\eta_s$, from $10^{-8}$ to $10^{-6}$ Pa.s.m, are reported for soluble monolayers by surface macrorheology.\textsuperscript{43} Usually a large difference is also measured between surface macro- and micro-rheology ($\eta_s = 10^{-10} \ldots 10^{-9}$ Pa.s.m),\textsuperscript{44} which could be due to additional dissipations or some boundary conditions not included in the analysis of the results.\textsuperscript{45,46} Recent experiments using noncontact microrheology, with completely wetted particles as probes, report $\eta_s \approx 10^{-8} \ldots 10^{-7}$ Pa.s.m,\textsuperscript{45} which agree with our results for CTA\textsuperscript{+}Br\textsuperscript{−}. For CTA\textsuperscript{+}OH\textsuperscript{−} instead our results $\eta_s \approx 5.10^{-10}$ Pa.s.m are comparable to passive surface microrheology using partially wetted particles as probe particles.\textsuperscript{44} Given that the liquid-gas interfacial tensions are similar for CTA\textsuperscript{+}Br\textsuperscript{−} and CTA\textsuperscript{+}OH\textsuperscript{−}, one possible explanation of the difference in $D_s$ between the two surfactant solutions is related to the surfactant dynamics occurring at the porous particle surface. As already pointed out before, the adsorption of CTA\textsuperscript{+}OH\textsuperscript{−} surfactants inside the pores could lead to a local depletion of the surfactant concentration close to the external surface of the porous particle and therefore in a smaller surface viscosity felt by the particle.

![Figure 7](image)

Figure 7. (A) Mean squared displacement as a function of the lag time for $d = 4$ nm porous particles at the gas-liquid interface in pure water ($\Psi$), in $C = 0.5$ mM CTA\textsuperscript{+}OH\textsuperscript{−} aqueous solution (--) and in $C = 0.5$ mM CTA\textsuperscript{+}Br\textsuperscript{−} aqueous solution (8). The inset shows a typical particle Brownian trajectory at the surface. (B) Ratio between the surface and the bulk diffusion coefficients as a function of the particle contact angle for the same systems shown in (A) and for $d = 2$ nm porous particles at the gas-liquid interface in pure water (!). Solid line is the hydrodynamic prediction for a bare and clean interface. Dashed and pointed lines are drawn from a model valid for viscous and incompressible interfacial layers with a given surface viscosity $\eta_s$.

### Influence of the counterion on surfactant adsorption

We start this section by analysing the remarkable difference in surfactant adsorption on porous particles observed between CTA\textsuperscript{+}OH\textsuperscript{−} and CTA\textsuperscript{+}Br\textsuperscript{−}. For planar non-porous interfaces, both interfacial tensions at the liquid-solid and liquid-gas point to a very similar behaviour for CTA\textsuperscript{+} surfactants, independently from the counterions. However, CTA\textsuperscript{+}Br\textsuperscript{−} surfactants do not adsorb on $d = 4$ and 7 nm porous particles, while...
CTA$^+\text{OH}^-$ surfactants adsorb with very high surface excess concentrations $\approx 4$-10 mg/m$^2$ for $C$ around 0.5 mM.

To explain this striking different behavior we started to look at the pH of the solution. Adding CTA$^+\text{OH}^-$ surfactants in the solution in fact alters the pH of water, which changes from 6.4 to 10.5 for $C = 0.5$ mM. It is also well known that for planar silica surfaces, cationic surfactant adsorption usually increases if the pH increases because of the increase negative charge of the silica surface.$^{35}$ In order to test this hypothesis, we have added Na$^+\text{OH}^-$ to a CTA$^+\text{Br}^-$ solution ($C = 0.5$ mM) in the presence of porous particles. The final pH measured is very similar (pH = 10.7) to the pH of CTA$^+\text{OH}^-$ surfactants where a strong adsorption was observed. However also in this case, CTA$^+\text{Br}^-$ surfactants do not significantly adsorb on porous silica particles, see Figure 8. It is also important to notice that for planar interfaces, CTA$^+\text{Br}^-$ adsorb on silica even if the pH remains around 6.$^{35}$

If OH$^-$ ions are not sufficient to trigger surfactant adsorption on porous particles, we consider the opposite scenario for which the ion Br$^-$ is able to hinder CTA$^+$ adsorption on silica. Therefore, in the presence of porous particles, we measure the surface tension of CTA$^+\text{OH}^-$ solution adding 1 mM of Na$^+\text{Br}^-$. In this case, we observe that the surface tension of CTA$^+\text{OH}^-$ (in the presence of porous particles) remains constant instead of increasing, see Figure 8. Thus, it confirms that CTA$^+$ do not adsorb inside the pores of $d = 4$ and 7 nm silica particles if Na$^+\text{Br}^-$ is present in the aqueous solution.
Figure 8. Salt effect on the liquid-gas interfacial tension as a function of time after porous particle deposition for $d = 4$ nm particles (A) and $d = 7$ nm particles (B). The surfactant concentration is fixed, $C = 0.5$ mM, CTA$^+$OH$^-$ data (○) are the same as in Figure 2. Data for CTA$^+$Br$^- +$ NaOH (●), CTA$^+$OH$^- + [NaBr] = 1$ mM (▲), CTA$^+$OH$^- + [NaCl] = 1$ mM (▲) and CTA$^+$Br$^- + [NaCl] = 1$ mM (●) as a function of time are also shown.

In order to verify if this effect is specific to Na$^+$Br$^-$, we have also tested another salt: Na$^+$Cl$^-$. The surface tension of CTA$^+$OH$^-$ (in the presence of porous particles) in this case increases only by 1 to 3 mN/m, which confirms a salt hindrance effect on adsorption (see Figure 8).

Attempting to rationalize these finding, we consider some specific ionic phenomena related to adsorption phenomena. Some properties of OH$^-$, Br$^-$, Cl$^-$ and Na$^+$ are listed in table 1. The main differences between the three negative ions are their hydration numbers (3 for OH$^-$, 2 for Cl$^-$ and just 1 for Br$^-$) and their
Jones-Dole viscosity $B$ coefficients connected to their ability to weaken or strengthen the interfacial water structure close to hydrophobic molecules such as the surfactant tails. $Cl^-$ is at the border ($B \approx 0 \text{ M}^{-1}$) between chaotropic $Br^- (B = -0.04 \text{ M}^{-1})$ and kosmotropic $OH^- (B = +0.18 \text{ M}^{-1})$ behaviors.\textsuperscript{47} Hence $OH^-$ may favor hydrophobic interactions between CTA$^+$ hydrophobic groups, which supports the adsorption of surface micelles; whereas $Br^-$ leads to a weakening of these hydrophobic interactions. We have also measured the zeta potential $\zeta$ on bare porous particles,\textsuperscript{48,49} which shows always negative values: $\zeta = -28.5$ mV for $d = 4$ nm and $\zeta = -7.8$ mV for $d = 7$ nm; as expected for hydrophilic silica.\textsuperscript{50} For CTA$^+$OH$^-$ surfactants, $\zeta$ remains negative for $C = 0.25$ mM: $\zeta = -19.5$ mV for $d = 4$ nm and $\zeta = -7.8$ mV for $d = 7$ nm particles. On the contrary, in the presence of CTA$^+$Br$^-$, positive zeta potentials were measured at the same concentration ($C = 0.25$ mM): $\zeta = +5.3$ mV for $d = 4$ nm and $\zeta = +5.9$ mV for $d = 7$ nm particles, which points to the accumulation of positive charges on the external surface of the porous silica colloids. Hence, CTA$^+$ from CTA$^+$Br$^-$ (or H$_2$O$^+$) may accumulate on the silica portion of the porous particle outer surface. This outer silica surface is very small with respect to the total porous particle surface, which may explain the negligible change of the interfacial tension (or pH) observed in the experiments. Both the chaotropic effect of Br$^-$ and the repulsion due to the positive $\zeta$ lead to a hindrance of CTA$^+$ adsorption inside the pores, which could explain the weak adsorption of CTA$^+$Br$^-$ surfactants on porous silica colloids.

Some additional specific ion effects occur on the surface of silica, where silanol groups are known to undergo ion exchange reaction:\textsuperscript{50,51} $X^+ + SiOH = X(SiO) + H^+$ for pH $< 8$, where $X^+$ is a cation (H$_3$O$^+$, Na$^+$ or CTA$^+$), and for pH $> 8$, SiOH + OH$^-$ = SiO$^-$. Hence a competition between H$_3$O$^+$, Na$^+$ or CTA$^+$ cations is also expected. Na$^+$ from Na$^+$Br$^-$ and Na$^+$Cl$^-$ in fact competes with CTA$^+$ in the adsorption on silica,\textsuperscript{52} which can explain the weakening of CTA$^+$OH$^-$ adsorption in the presence of a sodium salt (see Figure 8). Note that similarly to OH$^-$, Na$^+$ possesses an elevated hydration number and a positive $B$ coefficient corresponding to a kosmotropic behavior, see Table 1.

<table>
<thead>
<tr>
<th></th>
<th>OH$^-$</th>
<th>Br$^-$</th>
<th>Cl$^-$</th>
<th>Na$^+$</th>
</tr>
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<tbody>
<tr>
<td>Bare ion radius (nm)$^{53}$</td>
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<td>0.195</td>
<td>0.181</td>
<td>0.095</td>
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<tr>
<td>Hydrated radius (nm)$^{53}$</td>
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<td>1</td>
<td>2</td>
<td>4.5</td>
</tr>
<tr>
<td>$B$ (M$^{-1}$)$^{57}$</td>
<td>+0.18</td>
<td>-0.04</td>
<td>-0.01</td>
<td>+0.06</td>
</tr>
</tbody>
</table>

Table 1

CONCLUSIONS

We have herein reported some fundamental aspects of the wetting and surface diffusion of porous particles at the liquid-gas interface and explored the mechanisms underlying cationic surfactant adsorption inside the particle’s pores. Contact angle of porous particles at the gas-liquid interface can be described in the framework of the Cassie model in partial wetting, and it depends strongly on the presence of the liquid inside the pores of the particle. For pores completely filled by the liquid, the particle contact angle is expected to be very low and porous particles may detach from the interface and diffuse in the bulk. Particle translational diffusion at the pure water interface agrees with the hydrodynamics prediction,
which makes these particles ideal probes for interfacial microrheology. Porous silica colloids are already able to adsorb efficiently CTA\(^+\)OH\(^-\) surfactants without any chemical surface treatments on silica. We determined a significant adsorption of CTA\(^+\)OH\(^-\) surfactant on \(d = 4\) and \(7\) nm pore size colloids. The lack of adsorption for \(d = 2\) nm porous particles agrees with the presence of surface micelles and not monolayers of surfactants inside the pores of silica. We quantified CTA\(^+\)OH\(^-\) surface concentration and showed that for \(C < CMC\), the adsorption increases with the pore size. Therefore, we investigated the effect of the counterion on surfactant removal in order to understand the opposite adsorption behaviour observed between CTA\(^+\)OH\(^-\) and CTA\(^+\)Br\(^-\) on porous silica colloids. pH and surface charge effects are not able alone to explain our experimental findings, which points to the importance of kosmotropic/chaotropic counterion character. Surfactant adsorption on the porous particle affects also the particle translational diffusion at the interface, which points to different boundary conditions on the particle external surface. Finally, these results can be used to implement an environmental friendly strategy to remediate the surface of water from soluble and insoluble pollutants. Yet for future investigations, we plan to functionalize porous silica particles with hydrophobic groups to target contaminants that accumulate preferentially only on the water surface. Moreover, based on our expertise on self-propelled Janus silica particles at the interface,\(^{54,55}\) we plan to investigate the role of enhanced active diffusion on the kinetic of surfactant removal.\(^{56}\) Attention will be paid to the effect of ionic species present in the aqueous medium on the self-propulsion particle behavior.\(^{57,58}\)

ASSOCIATED CONTENTS

Supporting information
Non-spherical mesoporous silica particles pore size distribution, Particle deposition protocols.

ACKNOWLEDGEMENTS

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(51) Wängnerud, P.; Olofsson, G. Adsorption Isotherms for Cationic Surfactants on Silica Determined


