Title
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Permalink
https://escholarship.org/uc/item/4z8310pk

Journal
Nano letters, 18(4)

ISSN
1530-6984

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Publication Date
2018-04-01

DOI
10.1021/acs.nanolett.8b00208

Peer reviewed
The Interfacial Assembly of Polyoxometalate Nanoparticle Surfactants

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Supporting Information

ABSTRACT: Polyoxometalates (POMs) using \( \{\text{Mo}_72\text{V}_30\} \) as an example, dissolved in water, can interact with amine-terminated polydimethylsiloxane (PDMS-NH₂) dissolved in toluene at the water/toluene interface to form POM-surfactants that significantly lower the interfacial tension and can be used to stabilize liquids via interfacial elasticity. The jamming of the POM-surfactants at the water/oil interface with consequent wrinkling occurs with a decrease in the interfacial area. The packing density of the POM-surfactants at the interface can be tuned by varying the strength of screening with the addition of cations with differing hydrated radii.

KEYWORDS: POM-surfactant, liquid/liquid interfaces, structured liquids, self-assembly, wrinkling

Colloidal particles can assemble at oil−water interfaces, screening energetically costly interfacial interactions between immiscible liquids.1−5 Particles that are strongly bound to the interface can impart mechanical integrity to the assembly. These quasi-two-dimensional assemblies can support anisotropic stresses, giving rise to a wealth of complex, non-equilibrium, all liquid structures.3,6,7 The energy binding particles to the interface increases quadratically with particle radius,3 and so in the case of nanoparticles (NPs) the reduction in the interfacial energy per NP is small. Therefore, compressive forces exerted on the NPs as the interfacial area decreases are sufficient to eject the NPs from the interface, causing the liquid/liquid interface to ultimately relax to an equilibrium spherical shape. As such, it is difficult to structure liquids into nonequilibrium shapes with uniformly functionalized NPs. Janus-type NPs can be prepared where the reduction in the interfacial energy per NP is greater but, as of yet, they have not been used to structure liquids into complex shapes.8,9 Janus-type polymeric particles, specifically Janus-star shaped copolymers, as described by Müller and co-workers10,11 and by Jiang et al.12 can be used to shape liquids; however, the polymer chains can contract upon compression and the loading of the star-shaped polymers at the interface must be exceptionally high to effect tailored shaping of the liquids. Recently, a very simple method was developed where NP-surfactants were formed at liquid/liquid interfaces by dispersing functionalized NPs in one fluid and polymers end-capped with a complementary functionality in an immiscible second fluid.13 The NPs and polymers will interact at the interface, forming NP-surfactants having hydrophilic NP heads anchored by hydrophobic polymer tails. The NP-surfactants assemble spontaneously into a monolayer at the interface and the assemblies self-regulate by tuning the numbers of polymer chains interacting with the NPs. This strategy not only circumvents the detailed chemistries necessary to precisely functionalize the NPs but also affords a simple pathway to structure the liquids into highly nonequilibrium

Received: January 16, 2018
Revised: February 9, 2018
Published: March 20, 2018

DOI: 10.1021/acs.nanolett.8b00208
Nano Lett. 2018, 18, 2525−2529

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Polyoxometalates (POMs), well-defined molecular clusters comprised of early transition metal ions and oxo ligands having sizes ranging from 1 to 6 nm, provide an interesting candidate for use as NPs in the formation of NP-surfactants. In comparison to classic NPs, POMs are monodisperse, have well-defined structures, and have tunable surface properties, such as charge density. Diverse applications have emerged for POMs, for example as building blocks in supramolecular chemistry and as physical models for nanoscale clusters, and have found uses as multiphase microreactors, microfluidic devices, and all-liquid membranes, and multilength scale porous materials. Furthermore, the bonding between the NPs and end-functionalized polymers that result in the formation of the NP-surfactants is not covalent and the resultant assemblies are jammed; consequently, the NP-surfactant assemblies are responsive to external stimuli and the systems are reconfigurable.

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forms at the interface. In a typical POM-surfactant system, when the concentrations of PDMS-NH₂ solutions are varied and the aqueous solution of \{Mo₇₂V₃₀\} is fixed at 0.01 mg mL⁻¹, γ is found to decrease with increasing PDMS-NH₂ concentration. Above a concentration of 0.05 mg mL⁻¹, γ did not decrease further and remained constant at ∼22 mN m⁻¹. Here, the rate of reduction in γ increased with increasing PDMS-NH₂ concentration. In all instances, after the initial reduction in γ, there is a second slower process that leads to a continued slight decrease in γ with time. We attribute this slow process to the rearrangement of the POM-surfactant monolayer at the interface.

On the other hand, if the concentration of PDMS-NH₂ was fixed at 0.05 mg mL⁻¹ and the concentration of \{Mo₇₂V₃₀\} varied, the results in Figure 1b were obtained. For lower concentration of \{Mo₇₂V₃₀\} (<0.01 mg mL⁻¹), the reduction in γ is gradual. This is a result of the diffusion of POMs to the interface, the formation of POM-surfactants at the interface, and the reorganization necessary for subsequent POMs to be anchored to the interface. However, as the concentration of POMs is increased, the reduction in γ is much more rapid, and the equilibrium in γ is much lower ∼13 mN m⁻¹. Consequently, upon decreasing the volume of the droplet the POM-surfactant assemblies wrinkle, which is characteristic of a jammed assembly (Figure 1c and Movie S1). The droplets do not coalesce even when forced into contact (Figure 1d and Movie S2). During the compression, as the droplet’s interfacial area is reduced, the POM-surfactant assemblies jam causing either a wrinkling of the assembly or a permanent distortion of the droplet shape when the droplets are separated.

The variation in the packing density of the POM-surfactants was investigated by the addition of salt to the aqueous phase to tune the surface properties of \{Mo₇₂V₃₀\}. When the volume of the droplet coated with the POM-surfactants is decreased to the point where wrinkling occurs, this onset-of-wrinkling volume can be used as a measure of the initial packing density of the POM-surfactants at the interface; \(V_w/V_i\) is the amount that the initial assembly of the POM-surfactants must be compressed to solidify the assemblies, whereas \(V_w\) is the volume of the droplet when wrinkling is observed and \(V_i\) is the initial volume of a droplet. Figure 2a shows the value of \(V_w/V_i\) as a function of concentration of different salts after aging for 1 h. The snapshots were captured when the aqueous phase was withdrawn from the droplets. For each of the snapshots, the image was on the left-hand side (lhs) taken after 1 h aging, whereas the image on the right-hand side (rhs) shows the initial visible wrinkling state (Figure 2a and Movie S3–S6). It is evident that \(V_w/V_i\) depends on the type of salt added to the aqueous phase. Without the salt, \(V_w/V_i\) was ∼0.37, whereas it gradually increased to ∼0.52 with increasing concentration of NaCl. This saturates at a NaCl concentration of 20 mM. These results show that NaCl promotes the assembly of POM-surfactants at the interface. On the other hand, adding KCl or RbCl to the aqueous phase leads to much higher initial packing density of POM-surfactants. \(V_w/V_i\) rapidly increased to near unity, especially for the RbCl, \(V_w/V_i\) increased to 0.98 with the addition of only 2 mM salt, indicating a near complete coverage of the interface with POM-surfactants initially. When the concentration of RbCl is increased to 10 mM, the droplet self-wrinkles after ∼55 s, as shown in the third to fourth images in Figure 2b. The self-wrinkled droplet eventually fell from the needle (Figure 2b and Movie S7). This self-wrinkling behavior demonstrates the ability of the salt to promote the formation and assembly of the POM-surfactants, the significant reduction in the interfacial tension (Figure S4 and S5) that allows the droplet shape to extend (significantly increasing the interfacial area), and the budding-off of the droplet. Gravitational forces underpin the extension of the droplet.

\{Mo₇₂V₃₀\} carries 31 negative charges on its surface, theoretically balanced by 8 Na⁺, 14 K⁺, 2 VO⁵⁺, and 5 H₂O⁺ counterions. A very interesting and reasonable explanation for the above results is that the affinity of cations to the \{Mo₇₂V₃₀\} macroions relies on the size of hydrated cations. The smaller the size of the hydrated cations, the stronger the affinity. Here, we chose the monovalent cations Na⁺, K⁺, and Rb⁺, thus...
the ionic strength is the same at a fixed concentration. Rb$^+$ has the smallest hydrated size such that it can replace the original counterions with a stronger affinity to the $\{\text{Mo}_{72}\text{V}_{30}\}$ macroions. This stronger affinity reduces the repulsion between the $\{\text{Mo}_{72}\text{V}_{30}\}$ macroions, promoting the formation of POM-surfactants and the generation of high area density POM-surfactant assemblies at the interface. This is essentially a screening effect (Figure 3). Na$^+$, however, is less likely to exchange with the counterions, resulting in a smaller increase in the packing density of POM-surfactants at the interface. The observed self-wrinkling, extension and breaking off of the droplet (Figure 2b), can be attributed to the high packing density of the POM-surfactants, including the counter Rb$^+$ and the continuous decrease in $\gamma$ with time (Figure S4, even though the initial $\gamma$ is not that low, Figure S5). If more RbCl, for example, 20 mM, was added to the aqueous phase, the $\{\text{Mo}_{72}\text{V}_{30}\}$ precipitated due to the excess screening effects (Figure S7), which is evidenced by the lack of wrinkling after aging 1 h.

We have demonstrated the interfacial activity of $\{\text{Mo}_{72}\text{V}_{30}\}$ to interact with PDMS-NH$_2$ at a water/toluene interface to form POM-surfactants that considerably reduce interfacial tension. The jamming and wrinkling of the interfacial POM-surfactant assemblies were observed when the interfacial area was decreased. The packing density of $\{\text{Mo}_{72}\text{V}_{30}\}$ at the water/toluene interface was tuned by adding different salts where the size of the hydrated cations appeared to be of importance, with the screening of repulsive interaction between the POMs promoted a denser packing of the POM-surfactants at the interface.

# ASSOCIATED CONTENT

## Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.nanolett.8b00208, Materials and methods, $\gamma$ of water/toluene and POM-surfactant monolayer assembly (Figure S1–S6), the DLS data of POM aqueous (Figure S7) (PDF).

Movie S1 of droplet (AVI)  
Movie S2 of droplet (AVI)  
Movie S3 of droplet (AVI)  
Movie S4 of droplet (AVI)  
Movie S5 of droplet (AVI)  
Movie S6 of droplet (AVI)  
Movie S7 of droplet (AVI)

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Figure 3. Schematic diagram of the POM-surfactants assembled at the water/toluene interface at different screening effect conditions.