Van der Waals Emulsions: Emulsions Stabilized by Surface-Inactive, Hydrophilic Particles via van der Waals Attraction

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Abstract: Surface-inactive, highly hydrophilic particles are utilized to effectively and reversibly stabilize oil-in-water emulsions. This is a result of attractive van der Waals forces between particles and oil droplets in water, which are sufficient to trap the particles in close proximity to oil-water interfaces when the repulsive forces between particles and oil droplets are suppressed. The emulsifying efficiency of the highly hydrophilic particles is determined by the van der Waals attraction between the particle monolayer shells and the oil droplets enclosed therein and is inversely proportional to the particle size, while their stabilizing efficiency is determined by the van der Waals attraction between single particles and oil droplets which is proportional to the particle size. This differentiation in mechanism between emulsification and stabilization will significantly advance our knowledge of emulsions, thus enabling better control and design of emulsion-based technologies in practice.

Myriads of natural and industrial products are in the form of emulsions, in which fine oil droplets are well dispersed in water or vice versa by mechanical means with the aid of stabilizing agents. In almost all cases, emulsions need to be stabilized by surface-active agents: surfactants, amphiphilic polymers, solid particles or their combinations. By “surface-active”, it is meant that the agents are able not only to adsorb onto oil-water interfaces but also straddle the interface by projecting themselves partially into oil and partially into water. Up to date, the partial partitioning of surface-active agents at oil-water interfaces is universally accepted as the key to emulsion stabilization, which enables them to self-assemble into topologically ordered films and thereby stabilize the dispersed droplets by, for instance, electrical double layer (EDL) repulsion against coalescence. Counter-intuitively, here we show effective stabilization of oil-in-water (o/w) emulsions with surface-inactive agents—solid particles with exceedingly hydrophilic surfaces hardly wetted by oil at oil-water interfaces. Our results reveal that provided the EDL repulsion between the particles and oil droplets is suppressed, the van der Waals (VW) attraction between them is sufficiently strong to confine the particles in close proximity to oil droplet surfaces to form densely packed shells which efficaciously envelop the oil droplets for emulsion stabilization. In this new process, intriguingly, the emulsifying power of the highly hydrophilic particles is governed preferentially by the VW attraction between the densely packed particle shell as a whole and the oil droplets enclosed therein, while their stabilizing power is governed by the VW attraction between individual particles and oil droplets. To highlight the contribution of VW attraction, here the present o/w emulsions are referred to as “VW emulsions”.

Figure 1. (a) Formation of VW emulsions by homogenization of toluene and aqueous dispersions of 15 nm H-SiO₂ particles at pH 2.5, evidenced by milky upper phase (emulsion) and transparent lower phase (water) in the glass vial and negligible recovery of free oil after emulsification. No emulsions can be formed by 40 and 80 nm H-SiO₂ particles, evidenced by well-separated oil and water phases in the glass vials and fast and complete recovery of free oil after emulsification. (b) Dependence of the VW emulsion stabilization on the concentration of 15 nm H-SiO₂ particles in the aqueous phase at pH 2.5. The particle concentrations are marked on the photos. (c) Reversible stabilization of VW emulsions in response to pH. The inset indicates that the VW emulsions, stabilized by 15 nm H-SiO₂ particle at a concentration of 5.4×10¹⁵ particles/mL at pH 2.5, significantly destabilize upon increasing the pH to 4.0 and completely collapse at pH above 6.0. The pH adjustment is executed by dropwise addition of 0.1 M NaOH or 0.1 M HCl into the emulsions, followed by incubation for ca. 12 h under ambient condition. (d) Stabilization of VW emulsions (the upper phase in the photo) by citrate-stabilized gold particles of diameters of 16 nm at pH 2.5. The oil phase is tan in all experiments.

According to thermodynamics, the depth (∆E) of the potential well that traps solid particles at an oil-water interface with tension (γOW) is correlated with the contact angle (θ) of the particles at the
interface and their radius ($r$), which is symmetrically expressed with respect to $\theta=90^\circ$ as $^{[8]}$:

$$\Delta E = \pi \sigma_0 r^2 (1 \pm \cos \theta)^2$$  \hspace{1cm} (1)

The magnitude of $\Delta E$ can easily be $> 10^3$ kT ($k$ is the Boltzmann constant and $T$ is the absolute temperature) for solid particles with $\theta$ in the range of $20^\circ$ - $90^\circ$, even when $r = 5$ nm. Once adsorbed at the oil-water interface, therefore, the particles with such intermediate surface wettability are almost irreversibly trapped resulting in the outstanding long-term stability of particle-stabilized emulsions, known as Pickering emulsions. When $\theta < 20^\circ$, $\Delta E$ quickly diminishes so highly hydrophilic particles can hardly be held at the interface to stabilize a Pickering emulsion. In stark contrast, here we demonstrate that highly hydrophilic particles with $\theta=0^\circ$ can effectively and reversibly stabilize an o/w emulsion. The VW emulsions are stabilized by small particles more effectively than by large ones, while the interchange of differently sized particles readily occurs between the surfaces of the dispersed oil droplets and the continuous water phase. These unique features make the VW emulsions reminiscent of surfactant-stabilized ones. In contrast, Pickering emulsions are more effectively stabilized by large particles rather than by small ones and thus the preferential displacement of small particles by large particles on dispersed droplets was observed when $r < 2.5$ nm according to equation 1 $^{[8]}$.

Here uniform, spherical, silica particles with diameters of either 15, 40, 80, 150, and 200 nm were synthesized in water (Figure S1 and Table S1) and thoroughly cleaned to ensure the particle surfaces as well as the particle dispersions were completely free of the organic additives (Figure S2). As-prepared naked, hydrophilic silica particles, denoted as H-SiO$_2$, were dispersed in water (pH 5.5). When as-prepared particle dispersions were mixed with apolar organic solvents such as toluene and hexadecane, no stable emulsions were formed even under intense homogenization (Figure S3). When the pH of the as-prepared particle dispersions is adjusted to 2.5, intriguingly, 15 nm H-SiO$_2$ particles can effectively stabilize an o/w emulsion simply by hand shaking but larger particles cannot even by intense homogenization (Figures 1a and S4). Nevertheless, H-SiO$_2$ particles with diameter $> 15$ nm become effective in emulsion stabilization at pH 2.5 when the particle concentration in water is noticeably increased. For instance, 150 nm H-SiO$_2$ particles can effectively stabilize o/w emulsions at pH 2.5 after the as-prepared particle dispersions are concentrated by $> 5$ times (Figure 2b). The resulting o/w emulsions destabilize with an increase in pH and are completely broken into well-separated oil and water phases at pH $> 6.7$, while they are effectively re-stabilized upon lowering the pH back to 2.5 (Figure 1c). Since the isoelectric points of both H-SiO$_2$ particles and oil droplets in water are close to pH 2.5 $^{[10]}$ (Figure S5), the pH-response of the resulting o/w emulsion stabilization reflects the presence of pH-dependent EDL repulsion between the H-SiO$_2$ particles and oil droplets. Similar to 15 nm H-SiO$_2$ particles, citrate-stabilized gold particles with diameter of 16 nm can also effectively stabilize oil droplets in water at pH 2.5 (Figure 1d), at which the EDL repulsion between the citrate-laden surfaces of gold particles and oil droplet surfaces is minimized, implying the irrelevance of the surface chemical nature of highly hydrophilic particles in stabilizing o/w emulsions.

In order to evaluate the partitioning of H-SiO$_2$ particles at the oil-water interface at pH 2.5, the refractive index profile across the planar interface was accessed by means of ellipsometry according to our previous study $^{[11]}$. Figure 2a reveals significant changes in ellipsometric scans at the toluene-water interface after pure water is replaced by aqueous dispersions of 200 nm silica particles with partially hydrophobized surfaces. These particles straddle the toluene-water interface with a contact angle through water of 125 $\pm$ 2$^\circ$ and a surface coverage of 0.7 $\pm$ 0.2 based on a particle monolayer model at the interface $^{[11]}$. In contrast, no noticeable difference in ellipsometric scans is observed in the presence or absence of 200 nm H-SiO$_2$ particles at pH 2.5. This points to the absence of particles at the interface or to such weak adsorption which cannot be detected by ellipsometry. This is in good agreement with excellent resistance of naked silica surfaces against attachment of oil droplets in water at pH 2.5 (Figure S6) and little surface activity of as-prepared H-SiO$_2$ particles at pH 2.5 (Figure S7). Thus, we can conclude that H-SiO$_2$ particles are hardly (if at all) wetted by the oil phase at the toluene-water interface. Figure 2b shows that the o/w emulsions stabilized by 200 nm H-SiO$_2$ particles at pH 2.5 undergo complete destabilization after being frozen to -18 °C and thawed to room temperature, again implying non-wetting of the particles by the oil phase at the interface. In contrast, Pickering emulsions stabilized by 200 nm silica particles with partially hydrophobized surfaces are sufficiently stable against the freeze-thaw action, since $\Delta E$ arising from the partial wetting of 200 nm particles ($r=110$ nm) with reasonably hydrophilic surfaces ($\theta=30^\circ$) at the oil-water interface is over 7000 kT (Figure 2b).

![Figure 2](https://example.com/image2.png)

**Figure 2.** (a) Ellipsometric scans at the oil-water interface before (open squares) and after (open circles) addition of 200 nm H-SiO$_2$ particles (left panels) and 200 nm SiO$_2$ particles with partially hydrophobized surfaces (right panels), pH = 2.5. Lines are the fits to the data of amplitude tanθ (upper panels) and phase shift (lower panels) as a function of the incident angle. The particle concentration is ca. $1 \times 10^{12}$ particles/mL. (b) Destabilization of VW emulsions, stabilized by 200 nm H-SiO$_2$ particles at pH 2.5, by freeze-thaw action; the free oil phase is completely recovered after 20 min thawing at room temperature. The particle concentration is ca. $5 \times 10^{11}$ particles/mL. In contrast, Pickering emulsions stabilized by 200 nm partially hydrophobized SiO$_2$ particles are stable against freeze-thaw action. The oil phase is toluene in all experiments.

To effectively stabilize o/w emulsions, obviously, not only minimal EDL repulsion but also sufficient strong attraction between particles and dispersed oil droplets are necessitated in order to confine the H-SiO$_2$ particles in close proximity to the surfaces of the oil droplets in water at pH 2.5. Hydrophobic attraction can be ruled out taking into account non-wetting of H-
SiO₂ particles by the oil phase at the oil-water interface (Figure 2a) and excellent oil-repellence of naked silica surfaces in water at pH 2.5 (Figure S6). The VW attraction is therefore the only interaction trapping the H-SiO₂ particles in close proximity to the surfaces of dispersed oil droplets to form densely packed shells that envelop the oil droplets. To test our hypothesis, apolar organic solvents with different refractive indices were utilized as the oil phase (Table S2). Figure 3 shows effective stabilization of o/w emulsions by 15 nm H-SiO₂ particles at pH 2.5 when the refractive index of the oil phase, e.g. hexadecane, toluene and xylene, is significantly larger than that of water to create positive Hamaker constant (Table S2) and thus VW attraction between particles and dispersed oil droplets in water. The stabilization efficiency noticeably decreases as the refractive index decreases, e.g. octane and hexane. In contrast, no emulsion is formed when organic solvents of refractive index less than that of water are used, e.g. perfluorohexane, which creates VW repulsion between particles and oil droplets as a result of negative Hamaker constant (Table S2). The revealed refractive index effect supports the notion that the formation and stabilization of o/w emulsions is driven by the VW attraction between H-SiO₂ particles and oil droplets in water.

![Figure 3](image-url)

Figure 3. Dependence of the VW emulsion stabilization of 15 nm H-SiO₂ particles at pH 2.5 on the refractive index of the oil phase. The particle concentration is 1.4 × 10¹¹ particles/mL. The photos are taken in 6 h after homogenization of the oil and the particle dispersion.

To elucidate the formation and stabilization of VW emulsions, we start calculating the energy associated with VW attraction between a H-SiO₂ particle with radius r and a giant oil droplet through water (∆Epw/w) with r (Figure 4), which cannot account for 15 nm H-SiO₂ particles being superior to large ones in emulsion formation and stabilization (Figure 1a) or the particle concentration effect (Figure 1b). For effective stabilization of o/w emulsions, oil droplets must be armored by densely packed particle shells. Hence we consider the VW energy of a number of particles (N) participating in the stabilization of a single oil droplet and estimate the VW attraction between a monolayer shell of densely packed particles and an oil droplet with radius R enclosed therein (∆Epw/w) as follows:

\[ W_{pw/w}^{VW} \approx N V W_{pw}^{VW} = -R^2/\xi r \]  

where N = R²/π₃ as R ≫ r. In contrast to Wₑpw, Wₑpw increases with 1/r and rises progressively to several 10⁻⁴ ε when r < 50 nm (Figure 4). This accounts well for the size-dependence of the emulsification effectiveness of H-SiO₂ particles (Figure 1a). It may also provide clues to the highly effective stabilization by surfactants (r = 1 nm) in spite of the very small value of ∆Epw/w trapping them at an oil-water interface. For a given r, on the other hand, Equation 3 indicates that the Wₑpw increases with N, which accounts for the particle concentration effect on emulsion formation (Figure 1b).

![Figure 4](image-url)

Figure 4. Variation of the values of Wₑpw/w (open circles) and Wₑpw/w (open squares) as a function of particle diameters (2r) calculated based on Equation 2 and Equation 3, respectively. For simplicity of calculation, x is set to 1.5 nm to achieve sufficiently large Wₑpw and, at the same time, retain other possible repulsions such as hydration and osmotic interaction to avoid the near-surface contact between the particle and the oil droplet. Hexadecane is chosen as the oil phase for calculation.

To gain better understanding of the particle size effect on the formation of VW emulsions, apolar organic solvents were emulsified in aqueous dispersions of the binary mixtures of small and large H-SiO₂ particles at pH 2.5, in which the numbers of the small and large particles were adjusted to be sufficient for either of them to effectively create the VW emulsion. Intriguingly, the resulting o/w emulsions are stabilized solely by the small particles, while the large particles remain exclusively in the continuous water phase (Figure S5a). This stresses that small H-SiO₂ particles are superior to large ones in emulsification. From a viewpoint of diffusion kinetics (Table S1), small particles reach the proximity to the oil-water interface faster than large one and thus may be the kinetic favorable for emulsification. This particle diffusivity effect on emulsification kinetics, however, may be largely smeared out under intense homogenization. Thus, it is plausible to postulate that the emulsifying power of H-SiO₂ particles is governed by Wₑpw/w. To unravel the particle size effect on VW emulsion stabilization, the o/w emulsions, stabilized by small H-SiO₂ particles, were placed atop the aqueous dispersions of large H-SiO₂ particles after the free small particles were completely removed from the continuous water phase of the VW emulsions. The displacement of the small particles by the large ones
becomes noticeable after 1 month and the complete displacement is achieved after 3 months, yielding VW emulsions stabilized solely by the larger particles (Figure 5b). By contrast, the substitution of small H-SiO$_2$ particles dispersed in the continuous water phase for large ones stabilizing the dispersed oil droplets is much less effective and becomes visible after 3 months (Figure 5c). These two sets of results stress that the large particles is superior to small ones in VW emulsion stabilization, thus implying that the stabilizing power of H-SiO$_2$ particles is governed by $W_{\text{p/so}}$. It is worth noting that in as-prepared VW emulsions energetically favorable substitution of large H-SiO$_2$ particles for small ones on the dispersed oil droplets is surprisingly slow (ca. 3 months) taking into account that the particles remain non-wetted by the oil droplets and particle exchange occurs exclusively in the continuous water phase. This slow substitution may reflect the difficulty for small particles to escape from the densely packed particle shells enveloping the oil droplets, which requires the particles to overcome the VW attraction with the nearest neighboring particles. This inter-particle VW attraction may therefore offer an additional contribution to VW emulsion stability.

![Figure 5.](image)

Figure 5. (a) Comparison of the intensity-averaged hydrodynamic diameter profiles of aqueous dispersions of 15 nm (green curve) and 40 nm (red curve) H-SiO$_2$ particles and mixtures of 15 nm and 40 nm H-SiO$_2$ particles (blue curve) with that of the aqueous phase obtained after homogenization of the binary particle dispersion and toluene to form VW emulsions (black curve). (b) Substitution of large particles for small ones at toluene-water interfaces of VW emulsions via incubation of 40 nm H-SiO$_2$ particle-stabilized emulsion in aqueous dispersion of 150 nm H-SiO$_2$ particles for 1 (blue curve) and 3 months (black curve). (c) Substitution of small particles for large ones at the toluene-water interfaces of VW emulsions via incubation of 150 nm H-SiO$_2$ particle-stabilized emulsion in aqueous dispersion of 15 nm H-SiO$_2$ particles for 3 months (black curve). Figures b and c show the intensity-averaged hydrodynamic diameter profiles of the aqueous phase obtained by destabilizing the VW emulsion via free-fall action after particle exchange for a given time. The hydrodynamic diameter profiles of the corresponding small (green curve) and large particle dispersions (red curve) are shown for comparison. Note that in all experiments, the concentrations of small and large particles in water are adjusted to be sufficient for stabilization of VW emulsions by themselves alone.

Taken together, our results demonstrate that highly hydrophilic particles can effectively yet reversibly stabilize oil droplets in water as a result of the VW attraction between particles and oil droplets when the EDL repulsion between them is minimized. However, the resulting VW emulsions are vulnerable to external disturbance and subject to the dynamic exchange of different sized particles between the dispersed oil phase and the continuous water phase. These unique dynamic features make VW emulsions clearly distinct from Pickering emulsions, where solid particles are reversibly trapped at oil-water interfaces via partial surface wetting, and largely reminiscent of surfactant-stabilized emulsions. Our results underline that the formation of VW emulsions is controlled by $W_{\text{p/so}}$ but their stabilization is by $W_{\text{vw}}$. The revealed differentiation in mechanism between emulsification and stabilization should advance our knowledge of emulsions, especially those stabilized by binary surfactants or surfactants and solid particles, in which small surfactants appear more effective on emulsification [13].

The present work underlines that the attractive forces between stabilizing agents and dispersed droplets, such as VW attraction and image charge attraction[14], provide an important role in emulsion formation and stabilization, which must be sufficiently strong to ensure the stabilizing agents are effectively confined in close proximity to oil-water interfaces. It may offer a more generic model which enables us not only to articulate the pre-history of formation of emulsions before stabilizing agents adsorb to oil-water interfaces to form monolayer shells but also to rationalize the formation of unusual emulsions in the presence of ions such as hydroxide anions[15,16]. It may also unravel a subtle but important snapshot of how particles (as well surfactants) adsorb to oil-water interfaces from water, thus shedding light on many fundamental processes occurring at fluidic interfaces[16-18]. Taking into account the ubiquity of VW interactions, hopefully, our work will stimulate innovative designs of colloidal formulations in many disparate technical applications ranging from drug delivery for oral administration to bitumen emulsion for road construction.

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Keywords: Emulsion • Particles • Interfaces • Colloid • Interactions

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Surface-inactive, highly hydrophilic particles can effectively and reversibly stabilize oil-in-water emulsions thanks to van der Waals attraction between particles and oil droplets in water, thus referred to as van der Waals emulsion. In contrast to Pickering emulsions, the new emulsions are vulnerable to external disturbance and subject to the dynamic exchange of different sized particles between oil and water phases.