Colloidal dynamics and elasticity of dense wax particle suspensions over a wide range of volume fractions when tuning the softness by temperature

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1. Introduction

Emulsions are ubiquitous in our daily life and appear in various fields of research and applications such as cosmetics, pharmaceuticals, or food. Due to the large variety of industrial applications where emulsions are employed \cite{1}, a plethora of research studies have been carried out to probe the rheological properties of emulsion such as its behavior under shear flow or characterization of the yield stress at high volume fractions \cite{2–4}. Compared to suspensions of hard particles dispersed in a continuum phase, an emulsion is a soft particle system where the liquid-liquid interface is stabilized by surfactant or nanosized particles. Owing to this microscopic difference of the dispersed phase, the macroscopic properties are different in many ways as well. Utilizing the knowledge on the suspensions, it will be useful to compare the characteristics of emulsions that can be physically differentiated from those of suspensions. Thus, the existing models based on suspension systems can be used to describe the micro/macroscale dynamics of the emulsions as well, such as for example the Krieger–Dougherty equation \cite{5}. While such a comparison is possible for a liquid dispersion it will fail close and above the random close packing or jamming...
transition $\phi = \phi_{\text{rcp}}$ when soft particles or droplets are in contact. Emulsions droplets can be compressed and one can access a higher volume fraction regime, well above $\phi_{\text{rcp}}$. Emulsion droplets possess soft and adjustable interfaces that deform under high shear or normal stress conditions. The volume fraction regime accessible for hard particle suspensions is normally limited to concentrations below $\phi_{\text{rcp}}$ due to its hardness or the vanishing compressibility of the material [6,2]. Higher volume fractions can only be reached by breaking or deforming the hard particles irreversibly. Therefore, previous studies at high volume fractions above $\phi > \phi_{\text{rcp}}$ addressed intrinsically soft particles such as such as emulsions, polymeric microgels or star polymers and only a few studies compare hard sphere and soft sphere suspensions systems directly [6,2,7-9]. To overcome these limitations, and to compare hard and soft particle features using the same material, we study 1-eicosene wax dispersed in water system which has a melting point at $T_m = 24.6°C$. This system changes its characteristics from a hard particle suspension to an emulsion or vice versa by changing the temperature. Above $T_m$, wax melts and the particles become soft, while below the melting point, wax solidifies and becomes hard [10]. By changing the temperature, we can dial in softness or hardness while other physical properties of the system such as the solvent composition remain unchanged. We study the rheological properties of the suspensions over a wide range of volume fractions and compare the hard particle suspension and soft particle emulsion system in the liquid state. Moreover, we consider the systems dynamics above the jamming volume fraction at $\phi_{\text{jam}}$ using Diffusing Wave Spectroscopy (DWS) and rheometry. To this end we can access the physical properties of the high volume fraction suspension composed of micron-sized colloids, and can gain insight into the system’s internal structure and dynamics. In the experiments we study two different particle or droplet sizes and can thus also test whether the particle size plays an important role.

As explained later in detail, we find, that the system’s dynamics and rheology displays distinctly different properties depending on the volume fraction range considered, which we thus separate in three parts. First, at concentrations below 35 vol.%, denoted as the low volume fraction regime, microstructural changes lead to a characteristic slowing down of the particle or droplet diffusion as described in our earlier work [10]. Macroscopic influences of crowding are manifested by the increase in viscosity. No significant differences between emulsion droplets or hard wax particles were observed. Second, in the intermediate volume fraction regime ranging from 35 vol.% to $\phi_{\text{rcp}}$, we find microstructural differences between hard and soft particles, which affects the bulk rheology of the system such as the viscosity or storage modulus of the system. Finally, the third volume fraction regime above $\phi_{\text{rcp}}$ is denoted as the high volume fraction regime. In our work, the system can be prepared such that we can reach $\phi > \phi_{\text{rcp}}$ not only for the emulsion but even for the suspension, despite the non-deformability of hard particle systems. This can be realized by using the melting and solidifying process of wax particles, but it also makes the system properties depending on the history of the sample. In this case a high volume fraction emulsion system $\phi > \phi_{\text{rcp}}$ is first prepared and the droplets are subsequently solidified by a temperature drop. As a consequence the system becomes a densely packed solid particle suspension (of deformed wax particles).

We study the influence of the softness of particles by studying the bulk rheological properties of both suspensions and emulsion by applying oscillatory shear or a shear flow with a constant shear rate. Our experiments are designed to cover the widest range of volume fractions accessible. In particular we focus on the behavior of suspensions in the high volume fraction regime to clarify the influence of surface elasticity and particle softness by the comparison between emulsion and suspension. The structure of this paper is as follow: in experimental part, the preparation and characterization of the 1-eicosene system and also the experimental methods are described. In the results and discussion part, we first analyze the system for the low and intermediate volume fraction regime and then focus our attention on the high volume fraction regime. At high volume fractions we discuss the elastic properties of both the suspension and the emulsion system. A brief conclusion section will close the paper.

2. Experimental methods

We use 1-eicosene in water emulsion stabilized with 10 mM sodium dodecyl sulfate (SDS) as a model system. 1-eicosene wax has a melting point at $T_m = 24.6°C$ and the bulk viscosity in the liquid state at 30°C is approx. 4 mPa·s as measured in house with a rheometer. Samples were prepared using a custom-made couette cell: first, crude emulsions were made by rupturing oil–water–SDS mixture around 3000 rpm over 50°C and then the polydispersity of the sample was reduced using the depletion–fractionation method by Bibette [11,12]. Finally, two batches of samples with different size were obtained and the size distribution is plotted in Fig. 1, as derived from a particle sizing and counting analysis of bright-field microscopy images, see also Ref. [10]. Sample A has an mean diameter of 3.52 μm, polydispersity (standard deviation/mean diameter) of 25.6% and the volume fractions studied range from 10.7 vol.% to 90.2 vol%. Meanwhile, sample B has mean diameter of 1.75 μm, a polydispersity of 13.1% and the volume fractions studied range from 30.5 vol.% to 82.0 vol%.

The mass concentration of emulsion droplets is measured by drying and weighing and then converted to the volume fraction $\phi$ using known density of water and 1-eicosene. The emulsion samples are kept in an oven at 40°C and were found to be stable for up to a month but in some cases showed signs of coalescence after 8 weeks and longer. Experiments were done within one month after preparation to ensure that the droplet size distribution in the sample does not vary significantly. Below the melting point ($T_m = 24.6°C$), the system is expected to behave as a particle suspension, because wax particles solidify. However, the transition time to reach full solidification of the
droplets depends on the final temperature of the sample. At 20 °C, the transition of the emulsion to a suspension takes over 10 h which we considered too long to prevent drying or aging effects in our experiment. The reason for the relatively slow solidification of the droplets is probably due to the long-lived supercooled liquid state of the wax, which is favoured by the small size of the droplets. However, we have not studied this effect quantitatively. The time for solidification can be reduced to a more convenient time scale by lowering the temperature to 15 °C or less and at this temperature it takes less than 3 h for the system to reach a steady state. Melting takes place much faster already a few degree above T_m. Thus, above the melting point, the experiments were performed at 30 °C for the entire range of volume fractions. To make sure that there are no size changes of droplets, due to freezing and melting, we compared the viscosity and storage modulus of the sample over 10 temperature cycles and find no temperature history dependence.

The macroscopic mechanical properties of sample A and B were measured using a Physica MCR300 stress-controlled type rheometer (Anton Paar GmbH, Austria). A solvent trap was used to prevent drying. Before the measurement, for all experiments a preshear at 100/s for 60 s and rest for 300 s is applied to erase the effect of loading. We find that the emulsion (at high temperature) below ϕ_{tem} always shows a Newtonian behavior. For high volume fractions, for both the emulsions and the suspensions, we analyze the rheological behavior under small amplitude oscillatory shear. A strain sweep test is done at a frequency = 1 rad/s and the strain range of 0.001–100%. Frequency sweep tests are also performed in a range of strain amplitudes where linear viscoelasticity is guaranteed (0.01–1% in the volume fraction regime studied).

We also use DWS to analyze the system’s internal dynamics under multiple light scattering conditions, owing to high turbidity of the system [13,14]. Typical dynamic light scattering or static light scattering methods are not suitable to study such turbid media at concentrations exceeding 10 vol%. In this paper, we obtain a measure of the mean square displacement (MSD) from the commercial DWS Rheolab instrument (LSInstruments AG, Fribourg, Switzerland) in transmission geometry. The path length L of the DWS scattering cell is varied from 0.5 mm to 5 mm. The accessible range of the source’s optical transport mean free path l’ at a given concentration and the cell path length L. From the light scattering experiment, we capture changes of the particle movement as the temperature is varied.

3. Result and discussions

For the comparison of the physical properties of emulsion and suspension, we first analyze the behavior of the system at concentrations below ϕ_{tem} i.e. in the low and intermediate volume fraction regime and then above ϕ_{tem} (high volume fraction regime).

3.1. Low/intermediate volume fraction regime

We first consider the particle or droplet effective short time self diffusion coefficient D(ϕ), derived from the mean square displacements (MSD) probed using DWS: ⟨Δr^2(t)⟩ = 6D(ϕ)t, as shown in Fig. 2, to evaluate the effect of surface softness on the microscopic behavior of the system. All MSD curves are presented in Fig. S1 for both the suspension and the emulsion system. Diffusion coefficients for the low and intermediate volume fraction regime are calculated from the slope in Fig. S1 for t > 10^{-5} s. We find that the MSD curves vary linearly with time ranging from 10^{-3} to 10^{-2} s. To compensate the effect of temperature on the solvent viscosity we plot the dimensionless reduced diffusion coefficient D(ϕ)/D_0 using the diffusion coefficient at infinite dilution as a reference D_0 = kT/6πηR with R = 1.76 μm, Fig 1 (a) [6].

At low volume fractions, emulsion and suspension do not show any notable differences in D. The situation changes at around 35 vol%, where the data for the hard particle suspension clearly deviates from the soft particle emulsion data. The low/intermediate volume fraction regime is separated according to this criterion at around 35 vol%. In the intermediate volume fraction regime (35 vol% to ϕ_{tem}), the change ‘soft’ to ‘hard’ plays a significant role defining the particles dynamics, although the particles are not yet in contact with each other. For hard particle suspensions, D(ϕ) drops dramatically at around 35 vol%, which means the diffusive motion of a single particle begins to be slowed down by the presence of other particles. Such a drop in D(ϕ) of the suspension at around 35 vol% implies the confined diffusive motion of hard particles. This could be a consequence of weak van-der-Waals attractions [15] leading to the formation of aggregation clusters. Strong macroscopic creaming or gravitational settling is not observed on the time scale of the measurements indicating that either the clusters formed are short lived or that some kind of loose particle gel is formed [16].

We obtain information about the motion of individual particles from DWS and we also measure the bulk rheological properties. In Fig 3, the viscosity of the emulsions is shown for the low to the intermediate volume fraction regime. The emulsion behaves like a Newtonian fluid and we observe no dramatic changes in viscosity at around 35 vol% which coincides well with the data in Fig. 2 that shows similar trend of
Fig. 4. Reduced viscosity of the emulsion system at 30°C as a function of the droplet volume fraction (red triangles) using the water viscosity at 30°C as a reference. Solid line: prediction by the semi-empirical expression by Lionberger and Russel [17].

a gradual and continuous decrease for $D/D_0(\phi)$ [2,3]. In Fig 4, the reduced viscosity of the emulsion $\eta(\phi)/\eta_0$, extracted from the data shown in Fig 5 is plotted. The viscosity of emulsion below $\phi_{crp}$ fits well with the Lionberger and Russel equation, Eq. (1) [17]. The good agreement between the experimental data and the semi-empirical equation for hard spheres means that the liquid wax emulsion system at high temperatures mimics a hard sphere system up to volume fractions $\phi \leq 50\%$.

$$\eta = \frac{\eta_0}{1 - \phi(1 + \phi - 0.184\phi^2)}$$

(1)

Meanwhile for the suspension, unlike the emulsion, it behaves like a viscoelastic material in which an elastic modulus can be measured for both the low and intermediate volume fraction regime, indicating a system spanning weak network of force chains [18]. Fig. 5 shows the result of the strain sweep test of a 50.05 vol% sample as an example. Here the storage modulus and loss modulus have plateau values in low strain regime followed by a decrease as strain increases. No local maximum is observed and both $G'$ and $G''$ decay steadily. The details of the suspension system will be further discussed in next chapter.

3.2 High volume fraction regime

Hard particle suspensions cannot be prepared at high volume fractions above $\phi_{crp}$ due to the non-deformability of particle itself. Here, we use the 1-eicosene wax system to achieve a highly concentrated particle systems. Due to the deformability of the soft particle, the system easily reaches the high volume fraction regime. When particles are solidified after the sample composition has been fixed at $\phi/\phi_{crp} > 1$ the hard particle suspension retains a memory of it's prepared state. In this high volume fraction regime, the particles must have a deformed shape with facets at the interface. Images of such facets are clearly visible also for the case of pickering emulsions as shown Ref. [19]. In the limit of very high volume fractions approaching $\phi \approx 1$, the system shows a foam-like structure [3,2] which consists of a network of thin films and plateau borders composed of the continuum phase. The mechanical stability of emulsion system is then determined entirely by the strength of the thin film at the plateau borders [3].

Next, we will have a closer look at the dynamics and the stability of emulsions and suspension in this high concentration regime. For the particulate system, any given particle is at least temporarily surrounded or caged by nearest neighbors. If the concentration is not high enough (i.e. well below $\phi_{crp}$), the movement of particles can be slowed down for a short period of time but the particle can finally escape from its cage by diffusive motion, also called long time diffusion [20]. Meanwhile, as the concentration increases above $\phi_{crp}$, not only the short time scale diffusive motion but the long time scale diffusive motion (escaping from the cage) of the particles essentially vanishes and particles become permanently trapped in their cages. Initially the motion of particles or droplets in concentrated system is limited to the short time self-diffusion within the cages and long time scale dynamics shows highly correlated dynamics which may be entirely arrested eventually [21,22]. When particles begin to touch the systems jams at $\phi_{crp}$ and except for a few rattlers the motion of the particles or droplets is determined by their intrinsic softness [23,24].

This microscopic behavior of the particles can be analyzed using the DWS technique which can detect and measure the local positional fluctuations of particles on nanometer length scales, even inside a highly opaque medium. The measured intensity fluctuations of multiply scattered light can be converted to the autocorrelation function and the latter is then used as a measure of the dynamics of the sample. A detailed description of the technique is given for example in ref. [25-27]. One can follow the average particle dynamics over wide range of length and time scales. Here we convert the intensity autocorrelation function $g_2(t) = \langle I(t)I(t + \tau)\rangle/\langle I(t)^2\rangle$, of scattered light to the time dependence of the mean square displacement $\langle \Delta^2(r) \rangle$, neglecting collective scattering effects [26,28,29].

We quantify the dynamics of the system in different ways depending on the volume fraction regime considered. The viscous properties are defined by the diffusion coefficient obtained from the slope of the MSD, data shown in Fig. 2, for the low and intermediate volume fraction regime. Meanwhile, for the high volume fraction regime, the elastic properties can be quantified by the plateau value of the MSD as shown in Fig. 6.

As the concentration increases, the movement of a single particle over long time scales changes: the diffusion slows down which is reflected as a decrease in slope for the short time MSD, while the MSD reaches a plateau and the plateau value or height is a measure of the solid-like character of the sample due to the restricted motion by other surrounding particles in the long time scale limit. This plateau value reflects the elasticity of the system: as the plateau is lower, the network of force chains becomes tighter and the material is stronger [30]. The elasticity index can be defined, as in denoted in Eq. (2), to quantify the systems’ elasticity in a qualitative passive microrheology approach [31,27,32,33].

El(Elaticity index) = 1/(plateau height of MSD)

(2)

Compared to the elastic modulus in bulk rheology, which is observed over the whole range of volume fractions studied, the El can only be probed in the high volume fraction regime because the particle movement is caged, over the time scales accessible to DWS, only for the volume fraction above $\phi_{crp}$ in Fig. 7 we show that the volume dependence of the El is different for the suspension compared to the emulsion:

![Fig. 5. Strain sweep test at $\omega = 1/s$ for a 50.1 vol% suspension, sample A (G’: solid symbols, G”: open symbols).](http://doc.rero.ch)
for the emulsion the EI continues to increase steadily while for the suspension the EI rapidly jumps to a high value at \( \phi \sim \phi_{rcp} \) and then appears increase further only very slowly at higher concentrations. In general the EI of the suspension is substantially higher than that of emulsion over the whole range of volume fractions considered here.

The origin of elasticity of highly concentrated emulsion is different from that of suspension: the elasticity comes from the compressed repulsive droplets permitting the storage of interfacial elastic energy. [34,35] This means the droplets are more and more deformed as the volume fraction increases, and the surface area of the droplets increase, which creates additional storage energy by imposing shear deformation at the interface. The increase of EI for the emulsion shown in Fig 7 can be interpreted in the same way: as the volume fraction increases, more flat facets and contacts between neighboring particles are created, thus the elasticity index increases by the distortion of particle itself. This is due to the deformable characteristics of the soft particle.

However, for the suspension, the situation is very different, even though the particles are also deformed at high volume fraction regime and thus have flat facets at the interfaces. Preformed facets of the hard particles cannot be deformed anymore once the particles are solidified. This implies that the distorted surface areas for single hard particle do not change and therefore the EI of the suspension increases more gently as the shape of the single particle is adjusted to the geometrical constraints at volume fractions above \( \phi_{rcp} \), blue symbols in Fig. 7.

To compare the EI to the bulk elastic moduli of the system, we also assessed the macro-rheological properties in high volume fraction regime. Strain sweep and frequency sweep test results are presented as an example for the 87 vol% of sample B sample (smaller size droplets). In Fig. 8(a), both the emulsion and the suspension show a linear regime until a certain critical strain is reach, known as the yield strain [36]. For the suspension the linear regime is much narrower and we observe no local maximum in loss modulus \( G'' \), while \( G'' \) of the emulsion shows a clear maximum, in agreement with previous studies on model emulsion systems [3]. Interestingly, for the suspension of 87 vol%, the form of the graph is almost the same compared to the data taken in the intermediate volume fraction regime (Fig. 5). The increase in volume fraction merely shifts the modulus upwards by more than two orders of magnitude.

With respect to the frequency dependence of the elastic moduli of the emulsions and the suspension, shown in Fig. 8b, we find that the storage and loss modulus are both are much higher for the suspension than for the emulsion, by a factor of 60 for \( G' \) and 1800 for \( G'' \) at \( \omega = 10 \text{ rad/s} \). Moreover the frequency dependency also show some difference: \( G' \) and \( G'' \) are almost independent of frequency for the emulsion, while for the suspension, \( G' \) slightly increases with frequency while \( G'' \) remains constant indicating that the system becomes more elastic.

To clearly reveal the difference in the frequency dependence, the storage modulus at frequency = 0.1, 1, 10, 100 rad/s are plotted versus volume fraction in Fig 9. Both the suspension and the emulsion show a
solid-like response in the high volume fraction regime but follow a rather different frequency dependence. For the emulsions, except for the low volume fraction regime (equal and below 35 vol%), little dependence on frequency is observed for \( \omega > 10 \text{ rad/s} \). Meanwhile the suspension, shows a clear frequency dependence of all volume fractions.

By analogy to the hard sphere system, the elasticity of the emulsion can be represented by a plateau elastic modulus, \( G_p \) [37], \( G_p \) is obtained from the analysis of the strain sweep test at frequency = 1 rad/s in the linear regime for samples A and B with different particle sizes, Fig. 10. Below \( \phi_{rcp} \), the droplets of the emulsion are not deformed so the microscopic movement of particles is similar for the emulsion and the suspension [38]. Above \( \phi_{rcp} \), the droplets are confined and start to be distorted by the presence of the neighboring particles and this additional deformation results in the elasticity of the emulsion [38]. However, the origin of elasticity of the suspension in this high volume fraction regime is not well understood, since the state can only be reached as long as the wax is liquid and it is not entirely clear microscopically how the particles adapt to this confinement once solidified.

As expected, for the emulsion in the intermediate volume fraction regime, \( G_p \) is higher when the size is smaller as shown in Fig. S2. However, for the suspension system \( G_p \) shows no clear size dependence in the high volume fraction regime. To quantitatively analyze this size dependency of \( G_p \), we introduce Laplace pressure \( \Delta \rho \) to normalize for the size effect in the plateau storage modulus as described in ref. [34]. In our study sample A and B both have the same interfacial tension since the system is based on the same wax(1-eicosene) and surfactant and only the size is different by a factor of about 2, Fig. 1, so that the Laplace pressure ratio between sample A and B has the value of approximately 2.

By using Laplace pressure ratio, one can take account of the size effect even without knowing the exact value of the surface tension. In Fig 10 we plot the scaled plateau storage modulus over the whole volume fraction range studied. The data collapse for emulsions and suspensions independently. Surprisingly, the master curve in Fig 10, compensating the droplet size effect just by simple scaling using the Laplace pressure ratio, can be achieved not only for the emulsion but also for the suspension system. The size range explored is very limited however and the experimental uncertainty is almost comparable in some cases, so a firm conclusion about the scaling of the elasticity of the suspension with particle size cannot be derived form our data.

For the emulsion, the size dependence is well adjusted using the Laplace pressure ratio over the whole volume fraction regime as shown in Fig. 10. If we take a closer look at the emulsion case, the \( G_p \) shows a different concentration dependence in the intermediate and high volume fraction regime compared to the emulsions. For the intermediate volume fraction regime\( (\phi < \phi_{rcp}) \), \( G_p \) follows the model equation \( G_p \sim \phi^{1/2} \) with a sharp rise at \( \phi_{rcp} \sim 0.5 \) (Fig 10 , red dotted line) [39]. The parameter \( \phi_{rcp} \), denotes a critical volume fraction we find to be approximately \( \phi_{rcp} \sim 0.5 \) which is somewhat lower compared to the effective close packing prediction of \( \phi_{rcp} \sim 0.6 \) in monodisperse model emulsions [40]. For the high volume fraction regime \( (\phi > \phi_{rcp}) \), \( G_p \) follows the network-spring model equation \( G_p \sim \phi/\phi_{rcp} \) with the same value of \( \phi_{rcp} \sim 0.5 \) (Fig 10, red dashed line) [40]. So for the whole volume fraction regime, the 1-eicosene emulsion acts as a fairly standard monodisperse model emulsion system.

For the suspension case, the size dependence and the concentration dependence of \( G_p \) changes as the volume fraction increases. For both intermediate and high volume fraction regime, \( G_p \) shows a power law behavior but the exponent we empirically derive, decreases from 3 to 2 as the concentration increases. Interestingly, we observe a sudden jump of the storage modulus at \( \phi < 55\% \) which, as we believe, is a signature of the hard particle nature of the frozen wax particles. Similarly to the emulsion this value is slightly lower than the nominal value of \( \phi_{rcp} \).

The finite elasticity at higher volume fractions can be understood in terms of the sample preparation protocol and the interaction between the wax particle subject to the forces imposed by the neighbouring particles in contact.

For the concentrated system above \( \phi_{rcp} \), the particles should have non-spherical shape for both the emulsion and the suspension [2] and have facets in their interfaces but the characteristics of the facets are
different. For the emulsion, the droplets are soft and the facets could be modified by the increased osmotic pressure caused by increase of volume fraction or by shear stress. The shape of the particles in the suspension cannot easily be distorted by particle external stresses. The generation of facets in the suspension system could be categorized in two different types: for the low and intermediate volume fraction regime, the facet is created by the crystallization process during the solidification expanding in radial direction from the center of particles, as shown in our previous work [10], meanwhile for high volume fraction regime, the facet is not only generated by crystallization but also induced by the presence of other neighboring particles in the liquid state before the temperature is reduced below T_m. Moreover during the cooling process reaching T < T_m stresses stored and force chains created in the emulsion can become frozen in and the elastic response, i.e. the shear modulus, in the solid wax will then be increased. We can also speculate that the number of contacts imposed by the jammed elastic emulsion [23,24] is retained after solidification. Although particles and droplets in both systems, in the high volume fraction ‘overpacking’ regime, have facets induced by the presence of other surrounding particles, contacts and facets contribute to the elasticity of the systems in a different way as clearly shown in Fig 7.

4. Conclusions

In this paper, we have studied the micro- and macroscopic behavior of 1-eicosene wax dispersed as micron scale droplets or particles in water over a wide range of volume fractions. Since 1-eicosene has a melting point around 24.6°C, changing the temperature makes the system alter from an emulsion to a suspension or vice versa without affecting any other physical properties such composition or particle/droplet size. In previous studies, for the hard particle system, concentrated suspensions above φ_{rcp} could not be prepared in a well-defined way due to the hard nature of the particles. However, in our work, we could overcome this problem and we could enter the high volume fraction regime for both particulate systems, hard and soft, by just changing the temperature of the wax suspended in system. This way we could broaden the scope of the comparison between emulsions and suspensions covering a wide range of volume fractions. We analyzed the rheological properties and the microscopic dynamic properties using DWS. The results of our experimental study are summarized schematically in the diagram shown in Fig. 11. For a better understanding of the system, we divide the volume fraction axis into 3 parts: the low volume fraction regime (up to 35 vol%), the intermediate volume fraction regime (35 vol% ~ φ_{rcp}), and finally the high volume fraction regime (above φ_{rcp}) based on the systems microscopic motion and macroscopic flow characteristics.

Around 35 vol%, which indicates change from the low to the intermediate volume fraction regime, the slope of diffusion coefficient suddenly drops but only for the suspension while it continues to evolve gradually for the emulsion. This implies that the change of the surface softness and the solidification of the particles play a significant role for suspension for the intermediate volume fraction regime likely leading to weak attractive interactions owing to the modified interfacial properties of the wax particles after solidification and a now dominant contribution due to van-der-Waals attraction. At concentrations higher than φ ~ φ_{rcp} characterizing the highest possible volume fraction where hard particles are maximally packed or jammed [41], both emulsions and suspensions act as highly elastic, particulate systems but the detailed rheological properties and the microscopic dynamics are very different. For the emulsions, as the volume fraction increases, the droplets deform more and more and the facet area of single particles also increases. This is reflected in the increase of elasticity index (EI) derived from single particle MSD’s, and we can identify the structural disorder and droplet deformation as the origin of sample elasticity. In this situation the modulus G_p is droplet size-dependent due to the decrease in Laplace pressure with increase in radius α/L. However, for the suspension, no additional facet area can be newly created after solidification. The generation of facets is only possible for liquid droplets and the configuration of wax particles is then frozen in during the solidifying process, likely retaining some memory of the number of contacts and force chains created in the emulsion state. The elasticity is then independent of the Laplace pressure of the liquid and only given by the structural conformation and residual elasticity of the solid wax.

Conflicts of interest

F.S. is a board member and shareholder of LS Instruments AG.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version

References


