Irreversible shear-induced vitrification of droplets into elastic nanoemulsions by extreme rupturing

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Many materials weaken through fracturing when subjected to extreme stresses. By contrast, we show that breaking down repulsive bits of matter dispersed in a viscous liquid can cause a dramatic and irreversible increase in the dispersion’s elasticity. Anionically stabilized microscale emulsions subjected to a history of high-pressure microfluidic flow can develop an unusually large elastic modulus as droplets are ruptured to the nanoscale, yielding “nanonaise.” As the droplet size approaches the Debye screening length, the nanoemulsion vitrifies. Consequently, the onset of elasticity for disordered uniform nanoemulsions can occur at droplet volume fractions far below maximal random jamming of spheres.

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Colloidal dispersions can behave in interesting and unusual ways when subjected to high shear stresses that alter their structures away from thermal equilibrium [1]. For instance, shearing a polymer entanglement solution can cause the polymers to stretch and even disentangle, leading to non-Newtonian shear-thinning behavior; the solution’s viscosity $\eta$ decreases at higher shear rates $\dot{\gamma}$ [2]. Other dispersions, such as concentrated hard spheres in a simple liquid, can exhibit a shear-thickening viscosity [3]: $\eta$ rises with $\dot{\gamma}$. Attractive hydrodynamic interactions between the hard spheres can lead to the formation of clusters of spheres that jam and can even percolate, effectively causing $\eta$ to diverge [4]. This increase in $\eta$ is reversible; thermal forces redistribute the spheres and the equilibrium particle structure returns. Clay-polymer “shake gels” can become temporarily elastic due to changes in the structure of interacting components after $\dot{\gamma}$ is raised [5–7]. All of these shear-induced rheological changes do not persist after the shear is removed and the shear-induced structures relax. In this work, we use the term ‘shear’ to refer generally to both simple shear and pure shear, also known as extensional flow [2].

Although it is relatively easy to cause a variety of complex dispersions in viscous liquids to become permanently elastic by changing their compositions, in general, it is quite difficult to transform a viscous dispersion of repulsive objects irreversibly into an elastic solid by subjecting it to a history of extreme shear without changing its composition. When making mayonnaise, an emulsion of oil droplets in an aqueous solution stabilized against coalescence by amphiphilic lipids and proteins from egg yolk, the elasticity is typically achieved by slowly adding more oil while vigorously stirring. The stirring causes the oil to be shear-ruptured from the macroscopic scale down to microscale droplets through the capillary instability [8], which is driven by the surface tension $\sigma$. As the droplet volume fraction $\phi$ increases and oil droplets begin to jam together and deform, the mayonnaise develops a shear elastic modulus $G'$, that is strong enough to overcome gravity, and the emulsion “sets”—it appears to become solid. The elasticity arises from work that must be done against surface tension to further deform droplets that are packed into a disordered foamlike structure [9]. This simple example shows that it is possible to transform a liquidlike dispersion into an elastic one by raising $\phi$ while shearing. Concentrated emulsions have been made somewhat more elastic through moderate shear introduced by sinusoidal amplitude variation rheometry [10]. However, a pathway to dramatically and irreversibly transform an emulsion that resembles a simple viscous liquid into one that resembles an elastic solid through shear without altering its composition has not yet been found.

The elasticity of glassy microscale emulsions of repulsive uniform droplets arises from the deformation of jammed disordered droplets [9,11]. At low $\phi < \phi_{\text{MRJ}}$, where the droplets are not jammed, the emulsion resembles a viscous liquid; whereas at large $\phi > \phi_{\text{MRJ}}$, the droplets repulsively jam and deform, and the emulsion resembles a solid. Here, $\phi_{\text{MRJ}} \approx 0.64$ is associated with maximally random jamming (MRJ) of spheres [12], formerly referred to less precisely as random close packing (RCP) [13,14]. The linear elasticity of concentrated emulsions arises from the additional deformation of the jammed droplets induced by the applied perturbative shear, and the Laplace pressure scale of the undeformed droplets sets the scale of the elastic storage modulus, $G' \sim \sigma / a$, where $a$ is the droplet radius. This fundamental understanding of the elasticity of disordered deformable objects as a function of $\phi$ also explains $G'$ for foams of gas bubbles [15].

At present, no theory accurately predicts the linear shear modulus of emulsions by self-consistently including energy contributions from droplet deformation, entropy, and stabilizing repulsive interactions between droplet interfaces. Simulations of disordered uniform spherical droplets determined the repulsive jamming point to be $\phi \approx 0.64$ [11,16–18], in good agreement with experiments. The first of these simulations modeled the energy of deformation between two droplets, including the effects of the average local
coordination number, using Surface Evolver [19]. More recent simulations of random monodisperse foam have provided a much more accurate picture of the structure [20,21], but all simulations have neglected entropy and the electrostatic repulsions, instead treating interactions between the deformable surfaces as being “hard.” This is a reasonable assumption for most microscale emulsions and even larger foam bubbles, since ionic surfactants strongly inhibit droplet coalescence through short-range Debye-screened repulsions in the pair interaction potential \( U \). In this case, an effective volume fraction, \( \phi_{\text{eff}} = \phi h(1 + h/(2a)) \), where \( h \) is the separation between droplet surfaces, effectively accounts for small corrections of at most a few percent introduced by the short-range repulsion [21].

The glaring weakness in the existing explanation of the measured elasticity of uniform disordered emulsions is the \textit{ad hoc} assumption of a model for the film thickness \( h(\phi) \) that has been chosen to create a universal scaling curve of \( G'(\phi_{\text{eff}}) \). Although the model for \( h(\phi) \), which consists of a linear decrease from 17.5 nm at \( \phi_{\text{MB}} \) to 5 nm at \( \phi=1 \), is consistent with a measured value for the chosen stabilizer [22], it is very unlikely that this \textit{ad hoc} model for \( h(\phi) \) would be appropriate as the droplet radii approach the nanoscale. Moreover, this model is not based on any fundamental theory of interaction between surfactant-coated droplet interfaces.

Here, we reveal a striking effect: a liquidlike viscous material can be transformed into a solidlike elastic material through a history of extreme shear without altering its composition. Thus, a physical process that causes an irreversible breakdown of the structures within the material can be used to dramatically transform the material’s rheological behavior from that of a liquid to that of a solid. This is highly unusual, because many materials actually weaken irreversibly through fracture or relax back after being subjected to such high shear conditions. We demonstrate “shear-induced vitrification” using an ionically stabilized model emulsion system: we subject a microscale silicone oil-in-water “premix” emulsion at fixed \( \phi < \phi_{\text{MB}} \) to enormous strain rates near \( 10^8 \text{ s}^{-1} \). This extreme shear ruptures droplets down to nanoscale sizes, and the resulting disordered “glassy” nanoemulsion [23] can be quite elastic even though \( \phi \) itself has not changed. By analogy to “mayonnaise,” which commonly refers to elastic emulsions of microscale droplets, we refer to elastic nanoemulsions as “nanonaise.” As the rupturing occurs, \( h \) decreases towards the Debye-screening length, \( \lambda_D \), and the droplets repulsively jam into a Debye glass. We attribute the large elasticity of the nanoemulsions at low \( \phi \) to a combination of the increased influence of the Debye screened repulsions as well as an overall increase in the Laplace pressure, \( \Pi_L = 2\sigma/a_c \), of the undeformed nanodroplets. Using a simple model for disordered networks of repulsive elements, we extract the average interaction potential \( U(h) \) from \( G'(\phi) \) and this potential is in satisfying agreement with a Debye-screening law. Thus screened electrostatic repulsions between relatively undeformed nanodroplets play a key role in the elasticity of ionically stabilized nanoemulsions.

To make the premix emulsion, we disperse polydimethylsiloxane (PDMS or “silicone oil”) droplets at the desired \( \phi \) into an aqueous solution of sodium dodecylsulfate (SDS) at a concentration \( C_{\text{SDS}} \), typically above the critical micelle concentration (CMC) of 8 mM, using a mechanical mixer. The resulting microscale premix emulsion is polydisperse, having a broad size distribution centered at approximately \( \langle a \rangle \approx 5 \text{ \mu m} \). The premixed emulsion provides a feed to a high-pressure “hard” stainless steel–ceramic microfluidic flow device. Roughly 3 mL of emulsion is pulsed through microfluidic channels of 75 \( \mu m \) in a converging extensional flow geometry every second [24]. The microfluidic device mechanically amplifies the input air pressure \( p \), ranging up to 10 atm, to create liquid pressures up to about 2400 atm, yielding enormous peak strain rates of about \( 10^8 \text{ s}^{-1} \). To mitigate heating by viscous dissipation, the temperature of the output emulsion is controlled using a heat exchanger. At \( \phi=0 \), the extreme shear does not effect the viscosity of the surfactant solution.

Because the flow is inhomogeneous, it is desirable to recirculate, or “pass,” the emulsion through the microfluidic device more than once to ensure that all droplets experience the peak shearing conditions. After each pass \( N \) we recover a small volume of the emulsion and perform standard small-strain linear oscillatory viscoelastic rheometry using cone-and-plate and small Couette geometries to determine the frequency-dependent storage modulus, \( G'(\omega) \) and loss modulus, \( G''(\omega) \). Dynamic light scattering (DLS) of highly diluted emulsions provides the average radius \( \langle a \rangle \) and the standard deviation \( \delta a \). All measurements are conducted at room temperature, \( T = 23 \text{ °C} \).

Although shear-induced elastic vitrification can be achieved in only one pass at the highest air pressure \( p \approx 10 \text{ atm} \), we use a lower \( p \approx 3.4 \text{ atm} \) to show the hallmarks of vitrification over a larger range of \( N \) (Fig. 1). For fixed \( C_{\text{SDS}} = 116 \text{ mM} \) and \( \phi = 0.40 \), as \( N \) increases, a viscous re-

![FIG. 1. (Color online) Frequency dependence of the storage, \( G'(\omega) \) (solid symbols), and loss \( G''(\omega) \) (open symbols) moduli, of an oil-in-water emulsion with \( \phi = 0.40 \) and \( C_{\text{SDS}} = 116 \text{ mM} \) subjected to \( N = 2 \) (triangles), 3 (squares), and 6 (circles) passes of extreme microfluidic shear at an air pressure \( p = 3.4 \text{ atm} \). As \( N \) increases, the nanoemulsion becomes a highly elastic glass with \( G' > G'' \) down to low \( \omega \).](image-url)
polydispersity $\delta a/(a) = 0.15$ while fixing $C_{SDS} = 10 \text{ mM}$ [23]. This yields three different uniform concentrated nanoemulsions having radii: $\langle a \rangle = 75 \text{ nm}, 47 \text{ nm},$ and $28 \text{ nm}$. For each $\langle a \rangle$, we set the largest $\phi$ by ultracentrifuging at $20,000 \text{ rpm}$, and then diluting each stock nanoemulsion with surfactant solution. Strikingly, the rise in $G'_p(\phi)$ (Fig. 3) for nanoemulsions can be found as low as $\phi = 0.23$, much lower than $\phi_{MR}$. The sharp rise in $G'_p(\phi)$ is followed by a more gradual increase toward large $\phi$. This behavior is similar to $G'_p(\phi)$ for microscale emulsions, yet “nanonaise” is strongly elastic at much lower $\phi$ than has been observed for repulsive microscale emulsions.

Using a simple model of repulsive disordered objects, we obtain the droplet interaction potential $U(h)$ from $G'_p(\phi)$. Assuming $z=6$ nearest neighbors per droplet, the osmotic pressure is: $\Pi(\phi) = 3U(\phi)/V_{uc}$, where the unit-cell volume $V_{uc} = V_d/\phi$, and $V_d$ is the volume of a droplet. For dispersed repulsive networks under osmotic pressure, both experiments and simulations support the conjecture that $G'_p(\phi) = \Pi(\phi)$ [9], so we find $U(\phi) = G'_p(\phi)V_d/3\phi$ as the interaction energy per droplet-droplet “contact.” To determine $h$, we shift the measured $G'_p(\phi)/(\sigma/a)$ towards higher $\phi$ so that it overlaps with the prediction for deformable droplets with “hard” interactions [11]: $G'_p(\phi_{sat}) = 1.74(\sigma/a)\phi_{sat}(\phi_{sat} - \phi_{MR})$ (Fig. 4, inset). This shift provides $\phi_{eff}$, and we calculate $h = 2a/(\phi_{sat}(\phi_{sat})^{1/3} - 1)$, assuming the droplets are spherical. Since the Debye-screened repulsive potential is proportional to the square of the charge, we normalize $U(h)$ by $a^2$, assuming a constant surface charge density $\rho_s$ for all $\langle a \rangle$. This rescaling collapses all of the potentials onto a single master curve (Fig. 4), which we fit to $B^2P^2\exp(-h/\lambda_D)/(h e_0)$, where $B$ is a constant and $e_0 = 80$ is the relative dielectric permittivity of water. For $\rho_s = 3.2 \times 10^3 \text{ esu/cm}^2$ and $C_{SDS} = 10 \text{ mM}$ [26], the fit yields $B = 5.9 \pm 0.4$ and $\lambda_D = 3.8 \pm 0.5 \text{ nm}$, in good accord with the reported $\lambda_D = 3.5 \text{ nm}$ [27]. The excellent col-

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**FIG. 2.** (Color online) Shear-induced vitrification of an emulsion (see Fig. 1) is associated with droplet breakdown as the number of passes $N$ increases. (a) The average droplet radius $\langle a \rangle$ decreases and then saturates. Bars denote the standard deviation $\delta a$, not the error in the mean. An exponential decay with a constant saturation fits the data (line). (b) The storage modulus $G'$ at frequency $\omega = 10 \text{ rad/s}$ increases many decades and saturates; this fit by an exponential increase to a saturation (line). (c) The lower crossover frequency $\omega_c$ becomes very small for $N \geq 4$, signaling vitrification.
Debye-screening length of line is a fit to a Debye-screened surface repulsion, yielding a from all nanoemulsion data shown in Fig. 3 curve: $U(h)/a^2$, determined from all nanoemulsion data shown in Fig. 3 (same symbols). The line is a fit to a Debye-screened surface repulsion, yielding a Debye-screening length of $\lambda_D = 3.8 \pm 0.5$ nm. Inset: To determine $h$, $G'_p$ from Fig. 3 are scaled with $\sigma/a$ and shifted in $\phi$ onto a master curve: $G'_p(\phi_{\text{eff}})/(\sigma/a)$.

The lapse in Fig. 4 clearly demonstrates that a realistic model for $U$ must be used to accurately predict $G'_p$ of nanoemulsions at low $\phi$.

In addition to providing a satisfying explanation of $G'_p(\phi)$ without resorting to the ad hoc expression for $h(\phi)$, our interpretation of nanoemulsion rheology provides a macroscopic method for measuring $U(h)$ for soft, glassy repulsive colloidal suspensions of spheres. By introducing a repulsive contact disorder (RCD) interpretation of the mechanical shear elasticity of vitreous nanoemulsions, which assumes that $z=6$, jamming occurs at $\phi_{\text{MRO}}$, and $G'_p(\phi) = \Pi(\phi)$, we obtain the microscopic $U(h)$. In prior work on repulsive colloidal crystals, $G'_p(\phi)$ has been related to the microscopic $U(h)$ essentially by assuming $G'_p(\phi) = K_{\Pi}(\phi)$, where $K_{\Pi}(\phi)$ is the osmotic compressional modulus [28, 29], and packing occurs at $\phi = 0.74$ and $z = 12$. When this approach for crystals is applied to glassy colloidal systems, it fails to provide the correct scaling and it does not yield realistic $\lambda_D$ and $p$.

By contrast, $U(h)$ found using the RCD model is consistent with Bragg scattering experiments on magnetically manipulated ferrofluid emulsions at the same $C_{SDS}$ [26]. Although the assumption $G'_p(\phi) \approx \Pi(\phi)$ has been confirmed by simulations [17], it has received only minimal theoretical attention [30]. In principle, the RCD approach can be applied to obtain $U(h)$ when $G'_p(\phi)$ is known for any concentrated, soft, glassy repulsive colloidal system of spheres. By contrast, other techniques such as optical trapping [31], the surface forces apparatus [32], and ferrofluid emulsions [26], are typically performed as $\phi \to 0$. Nanoemulsions that are charge stabilized, whether by cationic or anionic surfactants, should exhibit similar $G'_p(\phi)$, whereas nonionic and polymer-stabilized nanoemulsions should exhibit different $G'_p(\phi)$ due to repulsions related to molecular compressibility.

In summary, shear-induced vitrification through irreversible structural breakdown provides an exciting route for making nanoemulsions that are highly elastic at surprisingly low $\phi$. This unusual and potentially useful property of nanoscale arises from the much greater relative importance of charge-screened repulsions between nanodroplets as $a$ approaches $\lambda_D$. Our understanding of nanoscale motivates a search for a broader range of dispersions that may elastically vitrify irreversibly when extreme shear causes the breakdown of repulsive elements. This also highlights a need for a self-consistent theory that accurately predicts $G'_p$ of emulsions and nanoemulsions, including repulsive interactions, droplet deformation, and entropy. Finally, we anticipate that careful microscopic rheology of disordered nanoemulsions can provide a window to the microscopic interaction potential of surfactants and other molecules.

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FIG. 4. (Color online) Scaled interaction potential as a function of separation between the droplet surfaces, $U(h)/a^2$, determined from all nanoemulsion data shown in Fig. 3 (same symbols). The line is a fit to a Debye-screened surface repulsion, yielding a Debye-screening length of $\lambda_D = 3.8 \pm 0.5$ nm. Inset: To determine $h$, $G'_p$ from Fig. 3 are scaled with $\sigma/a$ and shifted in $\phi$ onto a master curve: $G'_p(\phi_{\text{eff}})/(\sigma/a)$.

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