

Boundary Mobility Controls Glassiness in Confined Colloidal Liquids

Gary L. Hunter,^{1,*†} Kazem V. Edmond,^{1,†} and Eric R. Weeks¹
¹*Department of Physics, Emory University, Atlanta, Georgia 30322, USA*
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We use colloidal suspensions encapsulated in emulsion droplets to model confined glass-forming liquids with tunable boundary mobility. We show that the dynamics in these idealized systems are governed by physical interactions with the boundary. Gradients in dynamics are present for more mobile boundaries, whereas for less mobile boundaries, gradients are almost entirely suppressed. The motions in a system are not isotropic but have a strong directional dependence with respect to the boundary. These findings bring into question the ability of conventional quantities to adequately describe confined glasses.

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When cooled quickly, some liquids avoid crystallization and vitrify, becoming mechanically solidlike glasses [1]. The glass transition temperature T_g marks the point where molecular motions all but cease. For polymers and molecular liquids, one typically quantifies a material's glassiness in terms of relaxation times τ , which behave roughly as the inverse of mobility and increase dramatically as the material is cooled toward T_g . For bulk glasses, these statements are sufficiently general and apply universally. However, for small systems, T_g and $\tau(T)$ exhibit a dependence on system size, and surprisingly, these quantities may either increase or decrease relative to their bulk values [2–21]. These size-dependent variations in material properties are often collectively referred to as “confinement effects,” and the understanding of such effects is essential to the development of functional nanoscale materials.

Research from the polymer and molecular liquid communities shows that the manner in which confinement affects mobility is dependent on the material in which the glass is confined [3–17]. In particular, the interaction (or lack of interaction) between the sample and the boundary is important. Solid boundaries that chemically bond to the confined material suppress molecular mobility within the sample, whereas mobile or chemically repulsive boundaries enhance mobility [4–12]. The experimental evidence supports models in which these types of boundary effects propagate into a sample and give rise to gradients in dynamics [4–12]. The idea is that confining a sample to a smaller space results in a large proportion of the sample being close to the boundary, and, therefore, as the system is made smaller, T_g and $\tau(T)$ are more statistically influenced by material near the boundary. Therefore, depending on the nature of the interactions, these quantities may increase or decrease. Despite the evidence for dynamical gradients, it remains unclear how boundary mobility affects motions in different directions, i.e., in directions tangential or perpendicular to the boundary [22–24].

Colloidal suspensions have served as a valuable model of supercooled liquids and glasses and have elucidated much

of the fundamental physics underlying the glass transition in molecular systems (see Ref. [1] and references therein). In hard-sphere colloids, phase behavior is controlled by volume fraction ϕ rather than temperature (qualitatively, $\phi \sim 1/T$). When confined within rigid, immobile boundaries, otherwise, liquidlike colloidal suspensions transition to glassy dynamics at lower ϕ than in bulk samples where $\phi_g \approx 58\%$, comparable to an increase in T_g for molecular glasses [18–20,25]. These reports also show that the length scales at which samples become glassy increase with increasing ϕ . The dynamics in confined colloids also depend on wall roughness [19,25], a feature found in molecular dynamics simulations of Lennard-Jones liquids [16,17]. To date, however, no experiments with confined colloidal glasses have discussed the effect of directly modifying the mobility of the confining boundary. Hence, it is unknown in what manner in which confined molecular liquids respond to boundary conditions is universal and also applies to colloids or if the response depends on specific details of the system.

To address these issues directly, we use bidisperse colloidal suspensions confined in emulsion droplets as model glass formers with tunable boundary conditions. We observe these systems with fast confocal microscopy and use particle tracking methods to determine the motions of many particles [26,27]. We limit our attention to systems at volume fractions $\phi = 33\% \pm 1.5\%$ or $\phi = 46\% \pm 1.5\%$ (hereafter referred to as 33% and 46%, respectively; details of the uncertainties in ϕ are given in Ref. [28]).

The colloids are fluorescent poly(methyl methacrylate) spheres [29] with small and large radii $a_S = 0.532 \mu\text{m}$ and $a_L = 1.08 \mu\text{m}$. The spheres are dispersed in a density- and index-matched solvent of cyclohexyl bromide and decalin and are stabilized against aggregation by an $\approx 15 \text{ nm}$ layer of poly(12-hydroxy-stearic acid) (PHSA). Electrostatic repulsion between the particles is screened by saturating the solvent with tetrabutylammonium bromide salt [30].

Colloid-filled droplets, such as those shown in Figs. 1(a) and 1(b), are created by depositing small amounts of

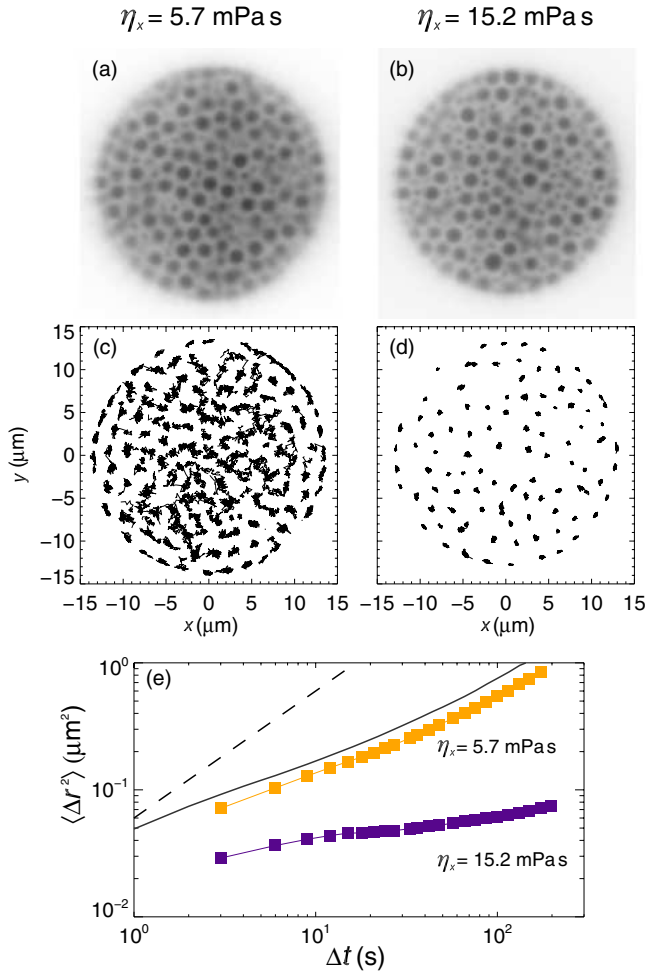


FIG. 1 (color online). Colloidal suspensions encapsulated within emulsion droplets. (a),(b) Raw confocal images of confined colloids at $\phi = 46\%$ with (a) $\eta_x = 5.7$ mPa s and $R = 14.4$ μm , and (b) $\eta_x = 15.2$ mPa s and $R = 14.1$ μm . Movies of these data are included in the Supplemental Material [28]. Note that these data are from separate experiments and that droplets are not physically near one another. (c),(d) Two-dimensional trajectories of the larger species of particles from the data immediately above. Motions are shown over a period of 600 s. Voids apparent in (c),(d) are due to untracked particles slightly out of the focal plane of the microscope. Bulk translational motions are subtracted with standard particle tracking routines [26], and bulk rotational motions are subtracted with a modified version of the procedures described in Ref. [27]. (e) Mean-square displacements for the large particles in the droplets shown. Light symbols (orange in color) correspond to data in (a),(c). Dark symbols (purple in color) correspond to data in (b),(d). The solid line is the MSD for an unconfined suspension at $\phi = 46\%$, and the dashed line has a slope of unity for comparison to Brownian diffusion.

suspension onto mixtures of glycerol and water and shaking gently by hand. A small amount of sodium dodecyl sulfate surfactant (3 mM) is present in the glycerol-water solution prior to shaking to stabilize the droplets against coalescence and prevent wetting of particles at the fluid-

fluid interface (we do not observe the Pickering effect under these conditions). The droplets are injected into glass chambers, allowed to sediment, and imaged via 2D or 3D confocal microscopy. The smaller particles move too quickly to be tracked reliably, so only trajectories for the large particles are computed [26,27]. However, the numbers of small n_S and large n_L particles are accurately identified in each droplet. Additionally, we find no evidence of particle size segregation over the course of several weeks. Droplet radii R are determined by measuring the well-defined radial coordinate of the outermost layer of large particles and adding a_L . Hence, the volume fraction is given by $\phi = (n_S a_S^3 + n_L a_L^3)/R^3$ [31]. The number ratio n_S/n_L varies slightly between droplets but is approximately $n_S/n_L \approx 0.9 \pm 0.1$ for $\phi = 33\%$ and 1.1 ± 0.2 for $\phi = 46\%$. Precise values are given in Ref. [28]. Only droplets with $R \leq 18$ μm could be successfully observed due to the limited field of view of the confocal microscope and the index of refraction mismatch between the internal and external phases. The droplet sizes are well below the capillary length for our solvents ($l_c \approx 6$ mm $\gg R$), so droplets do not significantly deform under gravity. Thermal fluctuations in the droplet surfaces are ≈ 1 nm; hence, these droplets function as smooth, nondeformable spherical confining cells.

To vary mobility at the confining boundary, we change the viscosity of the external aqueous phase. When a solitary particle of radius a is suspended in an unbounded Newtonian fluid, it experiences a constant drag coefficient, $6\pi\eta a$. Near a flat, mobile fluid-fluid interface, however, the drag coefficient is a nontrivial function of distance from the interface and the viscosities of both the suspending and external fluids [32,33]. The qualitative behavior in such a situation is intuitive: increasing (decreasing) the viscosity of the external fluid increases (decreases) the drag coefficient, provided that the particle is within a distance of $\approx 10a$ of the interface. Thus, by varying the viscosity of the external continuous phase, η_x , one modifies the viscous hydrodynamic coupling across the fluid-fluid interface and directly affects particle diffusivity. While the prior theory [32,33] was developed for an isolated particle near a flat fluid-fluid interface, one expects the qualitative picture to remain the same for denser suspensions and curved interfaces.

This is, indeed, what we observe, as shown in Figs. 1(c) and 1(d) comparing the motions of particles in droplets of similar size and ϕ , but where η_x differs by a factor of 2.7. Motion near the droplet interface is faster when η_x is smaller [Fig. 1(c)]; more surprising, however, is the dramatic difference in particle motion far from the boundary, where any direct hydrodynamic influence of the wall is screened [34]. Particle mean-square displacements (MSDs) given in Fig. 1(e) show the striking difference in the magnitude of particle motions in these two droplets and demonstrate that the ease with which particles

move at the boundary influences the dynamics in the whole system.

The effect of η_x as well as ϕ and R are more clearly represented by examining the MSDs at a single lag time Δt . MSDs as a function of Δt are given for all data sets in Ref. [28]. As shown there, MSDs vary quantitatively with Δt , but the behavior as a function of R or ϕ remains qualitatively the same for any Δt . Shown in Fig. 2 are particle MSDs at $\Delta t = 30$ s as a function of the droplet radius. For $\phi = 33\%$ and droplet radii $R \gtrsim 9 \mu\text{m}$, the dynamics are indistinguishable from those in an unconfined bulk sample. Therefore, no obvious effect of confinement or η_x is present at these length scales. Below this size, we observe decreased particle mobility and the onset of confinement effects.

With decreasing droplet size, we find further reduction in particle mobility; however, most significantly, the rate at which the dynamics decrease with R differs between the two η_x , with motions decreasing more strongly for larger η_x (dark symbols in Fig. 2). For the samples at $\phi = 46\%$, the mobilities are less than those in bulk for all droplets observed; therefore, the onset of a confinement effect occurs at droplet sizes $R \gtrsim 15 \mu\text{m}$. Hence, as ϕ increases, so do the length scales associated with the onset of confinement effects. This observation relates to growing length scales near the glass transition [1] and is consistent with previous research on colloidal suspensions confined in rigid chambers with immobile boundaries [18,21,25]. We also find that for $\phi > 50\%$ (data not shown), the particle

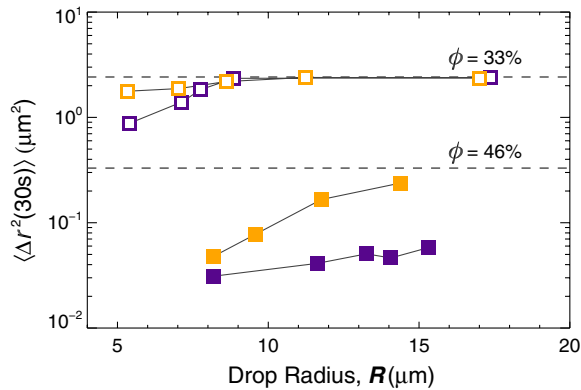


FIG. 2 (color online). MSDs at $\Delta t = 30$ s of particles in droplets of different ϕ and different η_x as a function of drop radius. Open symbols correspond to $\phi = 33\%$, and solid symbols are for $\phi = 46\%$. Light symbols (orange in color) are for $\eta_x = 5.7$ mPa.s, and dark symbols (purple in color) are for $\eta_x = 15.2$ mPa.s. Dashed horizontal lines are $\langle \Delta r^2(30 \text{ s}) \rangle$ in unconfined samples at the same ϕ . Data with $\phi = 46\%$ are from two-dimensional confocal images at the equatorial plane of the droplets, whereas data with $\phi = 33\%$ are from three-dimensional confocal images of the entire droplets. For meaningful comparison, we scale data for $\phi = 33\%$, such that $\langle \Delta r^2 \rangle = (2/3)\langle \Delta x^2 + \Delta y^2 + \Delta z^2 \rangle$, whereas for data at higher ϕ , $\langle \Delta r^2 \rangle = \langle \Delta x^2 + \Delta y^2 \rangle$.

motions for our range of observable R are very small and are on the scale of our uncertainty in tracking. As with data at lower ϕ , the rates at which the dynamics slow with droplet size depend on η_x , indicating that particles within the droplet are, indeed, affected by properties outside or at the fluid-fluid interface. In general, after the onset of confinement effects, the particles in larger droplets move faster than those in the smaller droplets, and a higher viscosity external phase results in lower particle mobility. The change in mobility with η_x is reminiscent of observations in confined polymers and small molecule glasses, where T_g and τ are strongly dependent on properties of the confining interface [3–7,11–17].

Changes in T_g and $\tau(T)$ (increases or decreases) relative to the bulk have been ascribed to mobility gradients originating from interactions at the boundary [4,5,7–11]. Visualizing the confined particles with confocal microscopy allows such gradients to be observed directly. Individual particle MSDs are resolved into radial and angular components and the mobility computed as $\Delta r = \sqrt{\langle \Delta r_r^2 + \Delta r_\theta^2 \rangle}$ as a function of a particle's distance s from the droplet interface (see Ref. [28] for further details of the calculation). We limit our discussion of mobility gradients to experiments where $\phi = 46\%$ but results from the lower ϕ case are similar and are given in Ref. [28].

Shown in Figs. 3(a) and 3(b) are particle mobilities Δr as a function of interfacial distance s for droplets of different sizes and different η_x . The dynamical gradients are apparent in droplets with the lower viscosity outer phase, and the slope of the gradient decreases with decreasing droplet size, perhaps even becoming negative for the smallest droplet studied. For the larger droplets, the particles are generally slower at the boundary than in the center of the droplet. With decreasing droplet size, the mobility curves shift to lower values, as would be expected from Fig. 2. In the case of the higher η_x , there are no obvious indications of a mobility gradient, but the decrease in mobility with decreasing droplet size is present. Small oscillations in mobility are the result of density fluctuations due to particle layering and are a generic feature of confined particles [18,21,25,35]. Layering has been shown to have a significantly smaller effect on mobility than confinement (see Refs. [18,19,25] for detailed discussions of this effect). The average structural properties of comparable droplets is independent of η_x (see Ref. [28]), and so it does not explain differences in mobility.

Decomposing mobility into radial and tangential components reveals further similarities between systems with different η_x . In Figs. 3(c) and 3(d), the radial motions near the interface are quite small for all droplets in a given η_x , but the mobilities of the particles nearest the interface do exhibit a slight dependence on droplet size. Approaching the center of the droplet, the radial mobilities increase, though this observation is much more prominent in the

lower η_x case. The increase of radial mobility far from the interface is similar to previous findings for the component of mobility perpendicular to a smooth or rough rigid boundary [15–18,21,25,35] but variations of this type in response to boundary mobility have not been reported.

In contrast, the tangential mobilities shown in Figs. 3(e) and 3(f) appear highest for the layer of particles immediately adjacent to the droplet interface ($s \leq 2a_L \approx 2 \mu\text{m}$). At $s \approx 2 \mu\text{m}$, the mobility falls in a stepwise fashion and remains essentially constant in the remainder of the droplet, with some larger fluctuations near the droplet center where statistics are poorer. As with the radial component, a decrease in droplet size results in decreased average mobility and is a trend present for droplets in either external phase, but more pronounced when the external viscosity is lower. Given the observations in Figs. 3(c)–3(f) for systems with different η_x , the qualitative similarity combined with the stark quantitative difference points to the strong effect that boundary mobility can have on the dynamics of the entire system.

The functional form of the mobility gradient in our case is unclear. Several models whose dynamics vary continuously with interfacial distance have been substantiated [5–7,15,17] but do not adequately describe our data.

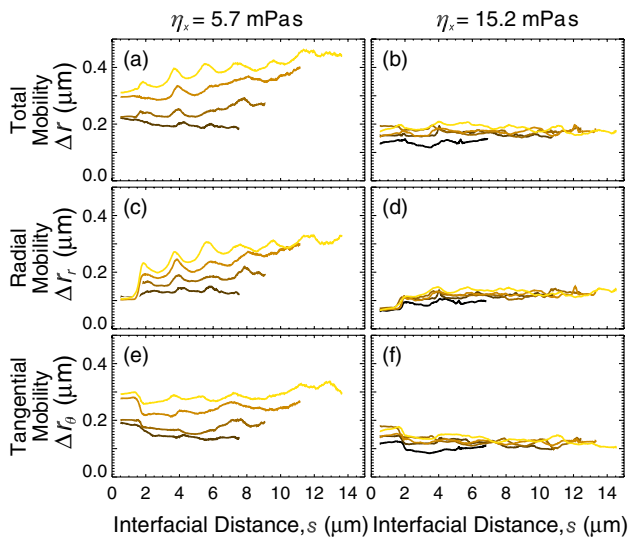


FIG. 3 (color online). Mobility as a function of interfacial distance. (a),(b) Particle mobility $\Delta r = \sqrt{\Delta r_r^2 + \Delta r_\theta^2}$ as a function of distance s from the fluid-fluid interface for systems with $\phi = 46\%$ and at $\Delta t = 30$ s. Data are from two-dimensional confocal images at the equatorial plane of the droplets and so include only one angular direction tangential to the interface. Data in the left panels (a),(c),(e) have $\eta_x = 5.7$ mPa s and right panels (b),(d),(f) have $\eta_x = 15.2$ mPa s. Droplets decrease in size from light to dark (orange to black in color). Where the data terminate at larger distances is approximately the droplet radius. See Tables II and III in Ref. [28] for precise radii. (c),(d) Radial components Δr_r , and (e),(f) tangential components Δr_θ of particle mobility.

Prior models of confined glassy materials assume that molecular mobility is a function of T , distance from the boundary, and the boundary conditions [5–8,11,15,17]. Hence, more confined samples have a larger fraction of their material close to a boundary and the influence of the boundary dominates the sample-averaged dynamics. The data in Fig. 3 show that mobility in our samples depends on these factors (a dependence on ϕ rather than T), but strikingly, it also depends on R . For example, the data in Fig. 3(a) for $s < 6 \mu\text{m}$ vary appreciably with the overall droplet size R . Thus, the slowed motion in more-confined droplets appears not just as the result of a stronger interface effect but is also partially due to a finite size effect, perhaps arising from the more pronounced curvature of the smaller droplets. Indeed, confining geometry has been found to be an important parameter in confined polymers [36]. While many differences exist between our idealized systems and polymers or supercooled molecular liquids, e.g., variations in particle geometries and particle-particle interactions, the response of a confined material to boundary conditions appears somewhat universal. Whether a general form incorporating appropriate variables can successfully describe the variety of confinement scenarios remains an open question.

Our data demonstrate that, as with polymers and molecular liquids, the properties of a medium external to a confined colloid can have a marked impact on the dynamics, even relatively far from the boundary. Thus, colloids continue to be a valuable model for glassy materials. Because particle mobilities depend on η_x , it is clear that confinement effects, in general, are not merely the result of a system's finite size; similar to polymers and small molecule glasses, the magnitude of motions occurring near the confining interface strongly influences motions in the remainder of the material.

An interesting implication of our observations relates to the interpretation of relaxation in confined glasses and how relaxation events and time scales in these scenarios can be appropriately quantified. For isotropic motions in the bulk of a glassy material, these quantities appear well defined. However, if relaxation is anisotropic, occurring on different time scales in different directions, then conventional definitions of relaxation may need revision. As found in prior experiments [18,21,25] and simulations [17,35], motions parallel and perpendicular to the boundary depend strongly on the shape and roughness of the confining boundary. Our results show that the fluidity of the boundary is an equally important parameter. The directional dependence of particle mobilities implies that relaxation also occurs anisotropically. Ignoring this anisotropy means that measurements of τ could be biased toward slower or faster motions (depending on the technique), obscuring or missing important physics entirely. While extremely sensitive, techniques used in polymers (e.g., fluorescent response of dyes and fluorescence recovery after photobleaching) and

molecular liquids (e.g., dielectric relaxation and solvation dynamics) are not yet able to simultaneously distinguish directionally dependent relaxations. In this respect, experiments with confined colloidal glasses may provide valuable insight for experiments with other confined glass-forming liquids and for understanding the glass transition in general [37].

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*GLHunter@gmail.com

†Present address: Center for Soft Matter Research, Department of Physics, New York University, New York, NY 10003, USA.

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