Slow dynamics in cylindrically confined colloidal suspensions

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Abstract. We study bidisperse colloidal suspensions confined within glass microcapillary tubes to model the glass transition in confined cylindrical geometries. We use high speed three-dimensional confocal microscopy to observe particle motions for a wide range of volume fractions and tube radii. Holding volume fraction constant, we find that particles move slower in thinner tubes. The tube walls induce a gradient in particle mobility: particles move substantially slower near the walls. This suggests that the confinement-induced glassiness may be due to an interfacial effect.

Keywords: colloidal glass, confinement, glass transition
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1. INTRODUCTION

Understanding the glass transition is one of the enduring questions of solid-state physics [1–6]. The problem is simply stated: in some cases, when a hot viscous liquid is cooled, the viscosity rises dramatically but smoothly as a function of temperature. At some temperature the viscosity is so large that the sample appears like a solid; this identifies the glass transition temperature. The same phenomenon can likewise be induced by increasing the density (increasing the pressure) [7]. In contrast to a regular phase transition which occurs at well-defined temperatures and pressures, the glass transition can depend on details such as the cooling rate. Likewise, while phase transitions are signaled by abrupt changes in the sample properties or their derivatives, the properties of a glass-forming material such as viscosity and diffusivity change smoothly, and to an extent the definition of the transition temperature (or pressure) is a bit arbitrary [8, 9]. One of the key questions is what is changing microscopically that is responsible for the macroscopic changes in viscosity; no structural length scale has yet been found that would clearly explain the viscosity change [10, 11].

One clever way to probe length scales is to confine a sample: rather than studying a macroscopically large sample (a “bulk” sample), studying a microscopic-scale sample. Many experiments show that glasses change their properties when their size is sufficiently small, both for small-molecule glasses and polymer glasses [8, 12–14]. One of the key observations is that the glass transition temperature \( T_g \) changes for confined samples. In some cases \( T_g \) increases: confined samples are glassier. However, in other cases \( T_g \) decreases. The key difference explaining the increase or decrease seems to be the boundary conditions [14]. Samples with free surfaces, such as thin free-standing polymer films, are less glassy (lower \( T_g \)). Samples confined to pores or on substrates can be more glassy (larger \( T_g \)), in particular if the sample molecules form strong chemical bonds to the confining boundaries.

We wish to use colloidal samples as a glass-forming system which can be studied in confinement. Colloidal suspensions are composed of solid particles in a liquid. As the particle concentration is increased, the sample becomes more and more viscous [15–18]. Above a critical concentration, the sample behaves as a glass, and a large number of similarities have been observed between the colloidal glass transition and glass transitions of polymers and small molecules [9]. The most widely studied colloidal glass transition is that of hard-sphere-like colloids, and the control parameter is the volume fraction \( \phi \) [19]. The glass transition point has been identified as \( \phi_g \approx 0.58 \), with simulations demonstrating that this requires some polydispersity [20, 21].

Experimentally, the colloidal glass transition shifts to lower volume fractions in confined samples: confinement makes colloidal samples glassier [22–27]. This has been studied exclusively in parallel-plate geometries, where samples are confined between two glass walls that are closely spaced. Often these experiments use bidisperse samples (mixtures of two particle sizes) so that the flat walls do not induce crystallization [22, 24]. An alternate approach is to roughen the walls [25].

The geometry of these prior colloidal experiments most closely resembles thin films, which are used to study the glass transition of polymers or small molecule glass formers in thin slits. However, small molecule glass formers are more commonly studied by using nanoporous substrates; a variety of these nanoporous substrates are reviewed in Ref. [8]. Some of these substrates are quite disordered with pores of a variety of shapes and sizes. Others are ordered: for example, porous oxide ceramics have a regular lattice of cylindrical nanopores with well defined sizes [8, 28], as do anodized aluminum oxide membranes [29]. Experiments find that confinement in cylindrical pores can both enhance or di-
minish glassy behavior [28]. Simulations show that the boundaries play an important role in this: rough walls that frustrate layering of particles result in glassier dynamics, and smooth walls result in less glassy dynamics [30, 31].

In this paper, we present a study of colloidal samples confined in cylindrical glass tubes, to mimic the geometry of cylindrical nanopores. We use confocal microscopy to observe both the structure and dynamics of the samples. Similar to prior colloidal work, we find that confined colloidal samples are glassier. In particular, particle motion is dramatically slower at the capillary tube walls, demonstrating that we see an interfacial effect. We use a bidisperse sample to prevent confinement-induced crystallization or other ordering, known to occur for monodisperse cylindrically confined spheres [32–37]. Nonetheless, the particles layer against the walls, and these layers slightly influence the motion in ways similar to previous observations [22, 24]. The data add to the analogy with confined small-molecule glass-formers. Additionally, they are of interest for colloidal suspensions themselves: the implication is that for microfluidic applications, it will be more difficult than anticipated to flow dense colloidal suspensions, as they will be glassier in small tubes than an equivalent bulk sample. Colloids confined in cylinders or small channels have been studied before, but only in dilute concentrations [38] and/or in extremely thin channels that are only one particle diameter across [39, 40].

2. EXPERIMENTAL METHODS

Our goal is to use hard-sphere-like colloids as a model system. Colloids have proved to be effective models with similarities to hard-sphere computer simulations; see [9] for a discussion. A hard-sphere system means that the particles do not interact with one another beyond their radius and are infinitely repulsive at contact [41–43]. An advantage is that the particle size can be selected to be \( \sim 1 \, \mu m \) in radius: small enough to undergo random Brownian motion, yet still large enough to be imaged using microscopy [9].

We use poly(methyl methacrylate) (PMMA) spheres coated with a polymer brush layer that sterically stabilizes the particles, preventing them from aggregating [44]. We use a bidisperse mixture with particles of two different radii, large particles with radius \( a_L = 1.08 \, \mu m \) and small particles with radius \( a_S = 0.532 \, \mu m \), helping us avoid crystallization [37]. The particles have a polydispersity of approximately 6% and additionally their mean radii \( a_L \) and \( a_S \) are each uncertain by 1%. Our particles are fluorescently dyed so that we can observe their motion with confocal microscopy. It is desirable to reduce the influence of gravity in a colloidal suspension by density matching the colloid with the surrounding fluid. To do this we use a standard mixture of 85% (weight) cyclohexyl bromide and 15% decahydronaphthalene (decalin, mixture of cis- and trans-). This solvent mixture also matches the particles’ refractive index, which is necessary for the microscopy. To reduce the influence of electrostatic repulsion between the particles, we saturate the solvent mixture with tetrabutylammonium bromide (Aldrich, 98%) with a resulting concentration of \( \sim 190 \, \mu M \) [45].

![FIGURE 1. Top: Composite photograph of capillary tube. The scale bar is 500 \( \mu m \). Bottom: 2D confocal image from experiment 55, with volume fraction \( \phi_{\text{tot}} = 0.49 \). The scale bar is 10 \( \mu m \).](attachment:image.png)
confined sample, such as shown in Fig 1(bottom), which can be studied at different locations to give different tube sizes. The capillary tubes vary slightly in radius as a function of length (slope of about 1°), and the slight taper does not seem to affect our overall results. The taper is slight enough that we cannot observe it in any of our confocal images. Due to the pipette puller protocol we used, the capillary tubes sag slightly, and so their cross-section is slightly elliptical rather than circular. This does not seem to influence our results and will be discussed further below. When the tubes are filled with the sample, some of the particles stick irreversibly to the tube walls due to van der Waals forces. This typically resulted in one complete layer of particles (of both sizes) coating the wall. During the course of our experiments, we do not observe any new particle sticking to the walls, nor do we observe any stuck particles becoming unstick.

We use a confocal microscope (Visitech vt-Eye) to study our samples. In a confocal microscope, laser light is scanned across the fluorescent sample and excites the dye to emit a different color light. The emitted light passes through a pinhole to remove the out-of-focus light and then is measured by a detector, a photomultiplier tube in our microscope. The sample is quickly scanned in x and y to acquire a two-dimensional (2D) image. Then the microscope focus is adjusted to scan more 2D images at different depths z, thus building up a 3D image. Being able to create 3D images makes confocal microscopy a powerful tool for studying particle dynamics in the system. Each 3D scan takes about 2-3 seconds, and we typically take movies comprised of 400 of these 3D images. The particles are tracked in 3D using standard tracking techniques [46, 47]. Within each confocal image, the small and large particles are easy to tell apart [see Fig. 1(bottom)] and so we can distinguish between them in our data [48].

We have two control parameters, the tube radius and the volume fraction. Complete descriptions of how these are measured are given in the Appendix; here, we summarize the key points.

Our first control parameter is the tube radius. The positions of the particles give us an accurate idea where the tube surface is. However, the tubes have an elliptical cross-section rather than a circular cross section. We measure the major and minor axes \( R_{\text{max}} \) and \( R_{\text{min}} \) for each experiment. The ratio \( R_{\text{max}} / R_{\text{min}} \) ranges from 1.14 to 1.39, with mean 1.24. Because we are concerned with confinement effects, we report our data in terms of \( R_{\text{min}} \) in general, although both radii are listed for all experiments in Table I. Note that we report the radii corresponding to the maximum positions of the observed particle centers: in general the particles at these maximum positions are the small ones (whose centers can get closer to the tube walls) and so the true tube sizes are larger by \( a_s = 0.532 \mu m \).

Our other key control parameter is the volume fraction \( \phi \). We measure this in each data set by counting the numbers of small and large particles observed within a subvolume of the tube, and converting this to the volume fraction using the known particle sizes. Note that 1% uncertainties in the particle radii translate to 3% uncertainties of the volume fraction, and since each particle radius is uncertain, we have an overall systematic volume fraction uncertainty of at least 5% [49].

In summary, our experimental method is to image different portions of the same tube in hopes to get a constant volume fraction with differing tube radii, and to study different tubes with different volume fractions to understand the role of volume fraction. In practice, we determine these variables when the data are post-processed, and report the measured values in Table I.

### 3. RESULTS

#### 3.1. Motion slows in confined samples

By following the motion of all of the particles in 3D, we can observe how the motion depends on confinement. We quantify this by calculating the mean square displacement, defined as

\[
\langle \Delta z^2 \rangle = \langle (z(t+\Delta t) - z(t))^2 \rangle
\]

where \( \langle \Delta z^2 \rangle \) is a function of the lag time \( \Delta t \), and the angle brackets indicate an average over all times \( t \) and...
all particles. Here we are considering the \( z \) direction to be along the axis of the tube, primarily because this axis is perpendicular to the optical axis of the microscope and therefore has less position uncertainty\(^1\). The data are plotted in Fig. 2, where each family of curves correspond to a different volume fraction. Within each family, the slower curves correspond to narrower tubes: confinement induces glassier behavior. To further quantify this, we consider the specific value of \( \langle \Delta x^2 \rangle \) at a lag time \( \Delta t = 100 \text{ s} \) (which is arbitrary but chosen to match prior work [22]). This value is plotted as a function of minimum tube radius \( R_{\text{min}} \) in Fig. 3. The different symbols correspond to different ranges of volume fractions, and in general within each range, smaller tubes correspond to slower motion. Plotting the data against \( R_{\text{min}} \) within each range, smaller tubes correspond to narrower tubes: confinement induces glassier behavior. To further quantify this, we consider the specific value of \( \langle \Delta x^2 \rangle \) at a lag time \( \Delta t = 100 \text{ s} \) (which is arbitrary but chosen to match prior work [22]). This value is plotted as a function of minimum tube radius \( R_{\text{min}} \) in Fig. 3. The different symbols correspond to different ranges of volume fractions, and in general within each range, smaller tubes correspond to slower motion. Plotting the data against \( R_{\text{max}} \), \( (R_{\text{min}} + R_{\text{max}})/2 \), or \( \sqrt{R_{\text{min}}R_{\text{max}}} \) does not change the overall appearance of this graph significantly. It is intriguing to note that the magnitude of the effect is less significant than was seen in similar experiments with parallel plates [22]. Here, the strongest influence of confinement is to lower mobility by a factor of \( \approx 3 \) (for circles, squares, and triangles in Fig. 3). With parallel plates, mobility was lowered by a factor of \( \approx 40 \) for data with \( \phi \approx 0.46 \) [22], a volume fraction in the same range as our cylindrical data. We are unsure why the two experiments differ in the magnitude of mobility reduction.

Microscopy allows us to spatially resolve details of the motion. If confinement-induced slowing is a finite size effect, then the motion might be spatially homogenoue: the whole sample feels that it is small [14, 50]. If instead the confinement-induced slowing is an interfacial effect (due to the sample-wall interface), then particle motion would depend on where each particle is relative to the boundary. Of course, both effects could be present simultaneously [28]. We check this by plotting the particle mobility \( \langle \Delta x^2 \rangle \) as a function of the distance \( s \) to the nearest wall in Fig. 4(a). It is immediately apparent that particles move slower when they are close to the wall (\( s \to 0 \)), and a plateau value for the mobility isn’t reached until several particle radii into the sample. Note that we are using the full 3D mobility \( \langle \Delta x^2 \rangle = \langle \Delta x^2 + \Delta y^2 + \Delta z^2 \rangle \), for a fixed lag time \( \Delta t = 30 \text{ s} \) (for which we have more statistics), and now the angle brackets indicate an average over those particles with the specific value of \( s \). The value of \( s \) is based on the initial position of the particle, at time \( t \) rather than \( t + \Delta t \). It is probable that the mobility near the wall is very slightly enhanced, due to particles which diffuse away from the wall during \( \Delta t \) and thus enhance their mobility [38]. The different curves are for small and large particles as indicated. Not surprisingly, the smaller particles are more mobile than the large ones. Our directly observed gradient in mobility is similar to that inferred from experiments on small molecule glasses [50–52], polymer glasses [53], and seen directly in simulations [30, 31, 54]. Our data show that the mobility changes over a distance

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\(^1\) The mean square displacement for the other components are qualitatively similar, except that they have more uncertainty which artificially increases the data at small lag times; see [49] for a discussion.
fluence of the large particles (solid and dotted lines, respectively). The mobility data in Fig. 4(a) are anticoerlated with $n_{\text{large}}(s)$: the local minima of $n_{\text{large}}(s)$ are indicated by the vertical dashed lines, and correspond to local maxima of $\langle \Delta r^2 \rangle$. The influence of $n_{\text{small}}$ is not seen, probably because the number fraction of small particles is much less than that of the large particles ($N_{\text{small}}/N_{\text{large}} = 0.07$ for these data). The anticoerrelation between number density and mobility matches what has been seen in prior work on confined samples [22, 24].

Further insight into the mobility is found by splitting $\langle \Delta r^2 \rangle$ into components parallel to the tube ($z$ direction), tangential to the tube wall ($\theta$ direction), and in the radial direction ($s$ direction). These components of mobility are shown in Fig. 4(b). Here it appears that the radial mobility is the most influenced by the oscillations of $n_{\text{large}}(s)$. Again, this is the same result seen before with flat walls [22]. The particles at the maxima of $n_{\text{large}}(s)$ appear to be at favorable positions and their mobility is reduced, whereas those at the minima are in less favorable positions with higher mobility. Motions in the $z$ and $\theta$ directions are along contours of constant mean $n$ and do not fluctuate with $n(s)$.

All of the results of Fig. 4 are qualitatively replicated in Fig. 5, which is data from a smaller radius tube ($R_{\text{min}} = 7.8 \mu m$ compared to $R_{\text{min}} = 17.1 \mu m$) and volume fraction only slightly larger than that of Fig. 4 ($\phi_{\text{tot}} = 0.54$ compared to 0.50). In Fig. 5(a), the mobility is lower near the boundary, lower for large particles, and oscillates with higher mobility corresponding to minima of $n_{\text{large}}(s)$. The data for the components [Fig. 5(b)] are noisier due to less statistics in the smaller tube, but again $\langle \Delta r^2 \rangle$ shows a stronger anticoerrelation with $n_{\text{large}}(s)$. The oscillations of $n_{\text{large}}(s)$ in Fig. 5(c) are more complex than those seen in Fig. 4(c), probably due to packing constraints of a smaller tube.

4. CONCLUSIONS

Our results – in particular Figs. 4(a) and 5(a) – suggest that the slower motion of confined colloidal samples is due to an interfacial effect, where particles near the sample walls are slowed. This agrees with a prior observation that colloidal particle motion is slower near rougher walls [23], a result demonstrating that the nature of the confining walls plays a role and not merely the finite size of the sample chamber. It is unlikely this is merely a hydrodynamic effect, as the magnitude of such an effect would only be a factor of $\sim 2 - 4$ in mobility and would not depend on $R_{\text{min}}$ [24, 38].

The clear gradient seen in Figs. 4(a) and 5(a) was not seen in prior work where colloids were confined between parallel walls [22]. It may be that the influence of confinement is stronger in the cylindrically confined case: between parallel walls, there are two unconfined directions, whereas in the cylinder there is only one unconfined direction [55]. Another possibility is that the curvature of the cylindrical walls introduces an effect not present with flat walls. These geometrical differences are the only major differences between the experiments.

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2 Although the hydrodynamic effect of the wall should be the same for flat or curved walls, sufficiently close to the walls. Prior simulations found essentially no difference in the diffusion constant between flat and curved walls, for particles within 1.5 diameters of the wall [38].
in this paper and those published earlier. As noted in Sec. 3.1, the other observed difference is that the slowing of motion in cylindrically confined samples (Fig. 3) is less pronounced compared to the observations between parallel plates [22]. This is counterintuitive given the argument above that cylindrical confinement is “stronger” than parallel-plate confinement. This trend is the opposite of that seen for small molecule liquids [55]. Future experiments may be able to elaborate on the question for colloids: at one extreme, particle motion could be studied in a half-infinite system near a wall, whereas at the other extreme, particle motion could be studied in a spherical pore. The data presented in this paper suggest that changing the dimensionality of the confinement in this way can result in interesting and qualitatively distinct behavior, in other words, enhancing a mobility gradient near walls while diminishing the overall confinement effect.

Our results also imply that flowing colloidal suspensions through small cylindrical tubes will be harder for smaller tube radii. Additionally, the decrease in mobility near the walls perhaps implies an increase in the apparent viscosity of the sample near the walls, thus modifying the flow velocity profile in a nontrivial fashion.

Note that we do not see any quantization effects: we do not see any particular change in the dynamics at any special ratios of particle sizes to tube sizes. This is in contrast to some theoretical predictions [56–58]. However, our data are only at a limited number of tube sizes, as shown in Fig. 3; our tubes are elliptical in cross section and so the ratio of particle size to tube size is not a constant for any given data set; and it is likely that such quantization effects are more subtle than we would be able to see in an experiment.

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APPENDIX

We explain in more detail how our experimental parameters are measured. In each case, we prepare samples and take data, measuring the tube radii and volume fraction from the data.

Tube radius: As noted in Sec. 2, the tubes do not typically have a circular cross-section. Also, in general, the images of the tube are not precisely aligned with the $xyz$ laboratory reference frame. To determine the radius, the position data are first rotated by $0 – 4^\circ$ around the $x$ and $y$ axes as necessary so that the $z$ axis of the data corresponds with the tube axis. (Note that the $z$ axis is not the optical axis of the microscope; rather, the $z$ axis is within a few degrees of perpendicular to the optical axis.) Then the data are projected onto the $xy$ plane and their center of mass is found. The $x$, $y$ coordinates are converted to $r, \theta$ and the maximum $r$ is found as a function of $\theta$. Following the procedure of Eral et al., in order to smooth the measured contour we fit $r(\theta)$ to a Fourier series up to $m = 4$ modes [38]. The function $r(\theta)$ is found to be well-described by an ellipse in all cases, and accordingly it is easy to determine the major and minor axes.

Volume fraction: For our experiments, we take movies from different portions of several tubes. Ideally each tube is filled with a sample of homogeneous volume fraction, but in practice the volume fraction varies slightly from region to region. Additionally, defining volume fractions in confined sample chambers is a little problematic as the concentration is inherently smaller...
at the walls simply due to packing constraints [57] even if the interparticle spacing is spatially homogeneous. To define volume fraction, we integrate the \( r(\theta) \) data described in the previous paragraph to determine the cross sectional area (adding on \( a_0^2 \) to determine the physical wall boundary). The length of the observed region is known, so therefore we know the volume \( V_{\text{tot}} \) of the tube that is imaged. Likewise we know the numbers of the small and large particles \( N_S \) and \( N_L \) that are in the image, and so the volume fraction can be determined from \( (N_S V_S + N_L V_L)/V_{\text{tot}} \) in terms of the individual particle volumes \( V_S \) and \( V_L \). \( V_S \sim \alpha_S^3 \) and likewise for \( V_L \), so 1% uncertainties in the particle radii lead to 3% uncertainties of the volume fraction. Since each particle radius is uncertain, we have an overall systematic volume fraction uncertainty of at least 5% [49]. There is also some uncertainty between samples as the different samples are observed to have different number ratios of small and large particles (see Table I), and so errors in small and large particle radii will affect the different volume fraction calculations in different amounts. Encouragingly, visual inspection of the images suggests that the calculated volume fractions listed in Table I are at least qualitatively in correct order. Samples with volume fractions within 0.02 of each other appear visually to be the same volume fraction, and samples with greater differences are visually distinct.

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