Long-Wavelength Fluctuations in Quasi-2D Supercooled Liquids

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Ryan C. Roberts, Jeremy C. Palmer,* and Jacinta C. Conrad*

ABSTRACT: We use molecular simulation to characterize the dynamics of supercooled liquids confined in quasi-2D slit geometries. Similar to bulk supercooled liquids, the confined systems exhibit subdiffusive dynamics on intermediate time scales arising from particle localization inside their neighbor cages, followed by an eventual crossover to diffusive behavior as cage rearrangement occurs. The quasi-2D confined liquids also exhibit signatures of long-wavelength fluctuations (LWFs) in the lateral directions parallel to the confining walls, reminiscent of the collective displacements observed in 2D but not 3D systems. The magnitude of the LWFs increases with the lateral dimensions of systems with the same particle volume fraction and confinement length scale, consistent with the logarithmic scaling predicted for 2D Mermin-Wagner fluctuations. The amplitude of the fluctuations is a nonmonotonic function of the confinement length scale because of a competition between caging and strengthening LWFs upon approaching the 2D limit. Our findings suggest that LWFs may play an important role in understanding the behavior of confined supercooled liquids due to their prevalence over a surprisingly broad range of particle densities and confinement length scales.

INTRODUCTION

Strongly confined fluids exhibit properties that are distinct from those observed in bulk systems. For example, in a variety of fluids, including gases,\textsuperscript{1,2} liquids,\textsuperscript{3,4} polymers,\textsuperscript{5-7} and colloidal suspensions,\textsuperscript{8,9} confinement has been shown to induce solid-like behavior that strengthens as the characteristic confinement length scale is decreased. There remains significant interest in understanding how the properties of fluids change as they are confined from three toward two dimensions, with practical relevance in applications ranging from separations and porous media characterization to energy storage and production and materials synthesis.\textsuperscript{1,2,10-14}

The crossover from two to three dimensions has been extensively explored for one prototypical supercooled liquid, a concentrated suspension of nearly hard spheres. When confined in a quasi-2D slit geometry, in which the height of the slit is much smaller than its lateral dimensions, uniformly sized particles form layers at the walls\textsuperscript{15,16} and arrange into a variety of ordered crystal structures.\textsuperscript{17,18} Although introducing size dispersity can frustrate long-range crystalline ordering,\textsuperscript{19,20} these systems can still undergo a transition to a disordered solid-like phase, in which the particle dynamics become increasingly slow and eventually arrest, upon increasing confinement. This transition is reminiscent of the 3D glass transition, in which the viscosity dramatically increases as the liquid is cooled. The dynamical signature of the 3D glass transition is the emergence of a plateau in translational correlation functions, signaling the formation of transient local cages that constrain particle motions. This plateau extends to longer times as the temperature is decreased or the particle concentration is increased.\textsuperscript{1,8,2,12,22}

Liquids in 2D and 3D exhibit different dynamics\textsuperscript{23} and phase behavior, however, complicating understanding of the relationship between the 2D and 3D glass transitions. In 2D, the crystal phase is unstable in the thermodynamic limit due to long-wavelength Mermin-Wagner (MW) fluctuations that represent elastic modes.\textsuperscript{24-26} Evidence of MW fluctuations has been recently found in both 2D liquids\textsuperscript{27} and amorphous solids.\textsuperscript{28} For 2D supercooled liquids, MW fluctuations disrupt particle cages and enable particles to become delocalized.\textsuperscript{23,25-32} Thus, 2D supercooled liquids exhibit distinct dynamics from their 3D counterparts.\textsuperscript{23,25} However, as shown in experiments\textsuperscript{28,31,32} and simulations,\textsuperscript{27,30,33,34} the behavior in 2D and 3D systems is similar when particle motions are characterized relative to the cages formed by their nearest neighbors.\textsuperscript{30,33,34} These studies suggest that analysis of the cage-relative motions of particles can be used to investigate the behavior of confined supercooled liquids as the dimensionality...
continuously changes from 2D to 3D, which remains incompletely understood.\textsuperscript{35}

Here, we use molecular dynamics (MD) simulations to show that quasi-2D supercooled liquid systems of particles confined in slits exhibit long-wavelength fluctuations (LWFs) analogous to those previously observed in dense monolayers. The magnitude of collective particle displacements in the lateral directions parallel to the confining walls, obtained by removing the cage-relative contribution to the mean-square displacement, increases with system size, consistent with the logarithmic growth expected for MW fluctuations. Both the amplitude of the fluctuations and the scaling constant depend nonmonotonically on the confinement length scale, which we attribute to a competition between arrest-driven caging and strengthening LWFs as the 2D limit is approached. These results indicate that dynamics in confined quasi-2D supercooled liquids are affected by steric effects and LWFs.

## METHODS

**Simulations.** We used event-driven molecular dynamics (MD)\textsuperscript{19,20,36} to simulate $N = 6912$--42 592 hard spheres confined in a slit geometry, consisting of two repulsive parallel walls separated by a distance $W$ along the $z$-axis of the simulation cell (Figure 1(a)). The walls have edge lengths of $L$ along the $x$- and $y$-axes of the cell, and periodic boundary conditions are imposed in these directions to model an infinite slab. The particles have unit mass $m = 1$ and are polydisperse in size (ca. 23%) to prevent spontaneous crystallization. Polydispersity was introduced by assigning each particle a diameter $\sigma$ randomly drawn from the distribution $P(\sigma) = A/\sigma^3$ with $\sigma \in [0.7255, 1.6099]$ (Figure 1(b)), yielding an average size of $\bar{\sigma} = 1$. Following convention, we adopt a system of reduced units to describe the systems and our results, in which Boltzmann’s constant $k_B = 1$, and $\sigma$ and $\tau = \sigma^2 (m/k_B T)^{1/2}$ are the fundamental measures of length and time, respectively, where $T$ is the temperature.\textsuperscript{19} We define the confinement parameter $H = W - \bar{\sigma}$ based on the width of the slit geometry accessible to an average-sized particle; the volume fraction $\phi = N \bar{\sigma}^2/6WL^2$ is defined in terms of $W$.

Initial particle configurations were generated by gradually compressing a system with $\phi = 0.45$, in increments of $\Delta \phi = 0.01$, to achieve the final targeted $\phi$ and $H$. After each incremental compression step, stress in the system was relaxed using MD and/or swap Monte Carlo (SMC).\textsuperscript{37--39} In most cases, stress relaxation was achieved by a running short MD simulation 10 time units in duration. For very confined or dense systems, relaxation following each of the last few compression steps was achieved by performing $10^5$ sweeps with the SMC algorithm, where 1 sweep is $N$ attempted MC moves. Following compression, the systems were equilibrated using SMC, performing enough MC sweeps ($10^5$--$10^6$) to ensure that the particles moved at least twice the mean particle diameter on average. The final equilibrated configurations from the SMC simulations were used to propagate microcanonical (NVE) MD trajectories $10^3$--$10^4$ time units in duration for the analysis of dynamics. Initial particle momenta in the MD simulations were randomly sampled from the Maxwell–Boltzmann distribution specified by $T = 1$. Dynamical quantities and associated statistical uncertainties were estimated from the averages and standard errors, respectively, computed from MD trajectories propagated from 50 independent realizations of the system at each state point using the protocols described above. Thus, approximately 50 000 simulations in total were performed for this study.

**Mean-Square Displacements.** From the trajectories, we characterize lateral positional fluctuations in the plane parallel to the walls using the mean-square displacement (MSD),

$$M(t) = \frac{1}{N} \sum_{i=1}^{N} \langle \delta \mathbf{r}_i(t)^2 \rangle$$

(1)

where $\mathbf{r}_i(t) = \mathbf{r}_i(t) - \mathbf{r}_i(0)$, $\mathbf{r}_i = \{x_i, y_i\}$ is the position of particle $i$, $t$ is the observation time, and the brackets $\langle \cdots \rangle$ indicate an average over different observation windows.

The cage-relative MSD $M_{\text{CR}}(t)$ characterizes the mean-square displacement of a particle relative to the motion of the cage formed by its nearest neighbors and is defined as

$$M_{\text{CR}}(t) = \frac{1}{N} \sum_{i}^{N} \langle \delta \mathbf{r}_{i\text{CR}}(t)^2 \rangle$$

(2)

where

$$\delta \mathbf{r}_{i\text{CR}}(t) = [\mathbf{r}_i(t) - \mathbf{r}_i(0)] - \frac{1}{N_{\text{nn}}(i)} \sum_{n=1}^{N_{\text{nn}}(i)} [\mathbf{r}_n(t) - \mathbf{r}_n(0)]$$

(3)

and $N_{\text{nn}}(i)$ indexes the nearest neighbors that form the cage for particle $i$.

Finally, the effects of LWFs can be analyzed by removing the contributions from cage-rattling from $M(t)$:

$$\Gamma(t) = M(t) - M_{\text{CR}}(t)$$

(4)

**Neighbor Identification.** Evaluation of the cage-relative MSD using eqs 2 and 3 requires identifying particles that form
the local neighbor cages. To account for the strong polydispersity, neighbors were identified using the surface-to-surface distance

\[ s_{ij} = |\mathbf{x}_i - \mathbf{x}_j| - \sigma_i + \sigma_j \]

where \( \mathbf{x}_i = \{ x_i, y_i, z_i \} \). Two particles were considered to be neighbors if \( s_{ij} \leq 0.5 \); this value corresponds to approximately the first minimum in the probability density distribution of \( s_{ij} \).

**Dynamical Corrections.** For some liquid-like systems, we used a modified version of eq 4:

\[ \Gamma(t) = M(t) - M^{\text{CR}}(t) - \Delta_{\text{TDD}}(t) - \Delta_{\text{CB}}(t) \]

where \( \Delta_{\text{TDD}}(t) \) and \( \Delta_{\text{CB}}(t) \) are corrections accounting for the asymptotic decorrelation of particle displacements at long times and local rearrangements of neighboring particles at intermediate times, respectively.\(^{29,34}\) These corrections are needed when the time scales associated with LWFs and the \( \alpha \)-relaxation (\( t_{\text{LWF}} \) and \( t_{\alpha} \) respectively) are of similar order. For the systems considered here, \( t_{\text{LWF}} \approx 10^3-10^4 \) time units. Thus, we use the criterion \( t_{\alpha} \leq 10^2 t_{\text{LWF}} \approx 10^4 \) time units as an indicator that these corrections may be necessary, employing an operational definition for \( t_{\alpha} \) as the time required for particles to displace one diameter on average (i.e., \( M(t_{\alpha}) = \bar{v}_i^2 \)). For many of our systems, we note that \( t_{\alpha} \) moves outside the accessible window for MD simulations, indicating that these systems are effectively glasses on these time scales and preventing characterization of the terminal relaxation behavior.

For liquids, the long-time behavior of \( M^{\text{CR}} \) is given by

\[ M^{\text{CR}}(t) = M(t) + (N_{\text{nn}}(t))^{-1} M(t) \]

implying that \( M^{\text{CR}}(t) > M(t) \). This behavior arises because the displacements of particles become uncorrelated from those of other particles in liquids on long times.\(^{29}\) For systems where \( M^{\text{CR}}(t) > M(t) \), we follow ref \(^{29}\) and fit the long-time behavior of \( M^{\text{CR}}(t) \) and \( M(t) \) to power-law functions of the form \( C^{\alpha} + b \), where \( C, a, \) and \( b \) are fitting parameters and \( a \) is globally fit for systems of constant \( \phi \) and \( H \). The correction for displacement decorrelation accounts for differences in the long-time growth of \( M^{\text{CR}}(t) \) and \( M(t) \) and is given by

\[ \Delta_{\text{TDD}}(t) = (C_{M^{\text{CR}}} - C_M) t^a \]

Similarly, power-law growth of particle displacements at intermediate times can arise from local rearrangements in neighboring particles, which is not accounted for in the definition of \( M^{\text{CR}}(t) \). We correct for this behavior using\(^{29,34}\)

\[ \Delta_{\text{CB}}(t) = D t^c \]

where \( D \) and \( c < 1 \) are constants found by fitting the function obtained after correcting for displacement decorrelation.

**Correlated Displacements.** Correlations between the displacements of particles and those of their surrounding neighbors were quantified using

\[ S_{\text{vec}}(t) = \frac{1}{N_{\text{nn}(t)}} \sum_{j=1}^{N_{\text{nn}(t)}} \frac{\delta \mathbf{r}_i(t) \cdot \delta \mathbf{r}_j(t)}{\max[|\delta \mathbf{r}_i(t)|^2, |\delta \mathbf{r}_j(t)|^2]} \]

This function is a modified, particle-resolved version of the spatial–temporal correlation function used in refs \(^{32,40,41}\). The maximum operation in the denominator ensures \( -1 \leq S_{\text{vec}} \leq 1 \) and that very small or large neighbor displacements are not disproportionately weighted. The function assumes values of 1 and –1 when the displacement of the central particle \( i \) is perfectly correlated and anticorrelated with the displacements of its surrounding neighbors, respectively. Nearest neighbors were identified using the same surface-to-surface distance criterion used in the evaluation of \( M^{\text{CR}} \).

To facilitate visualization, we coarse-grained the displacement vector correlation function into a continuous field using Gaussian functions centered at the location of each particle:

\[ S_{\text{vec}}(t, x, y) = \sum_{i=1}^{N_d} S_{\text{vec}}(t) e^{-\frac{((x-x_i)^2 + (y-y_i)^2)}{2\sigma^2(t)}} \]

where the sum is taken over the \( N_d \) particles in the contact layer near the lower wall.

### RESULTS AND DISCUSSION

**Dynamics of Confined Supercooled Liquids.** We analyze the dynamics of hard-sphere supercooled liquids confined in a quasi-2D slit geometry. The state of the systems is nominally specified by \{\( \phi, H \)\}. We study ranges of these parameters (0.550 \( \leq \phi \leq 0.600 \) and 1.0 \( \leq H \leq 14.0 \)) corresponding to high to moderately confined dense packings of particles that exhibit glassy dynamics.

The MSDs for a moderately confined system \( (H = 3.00) \) illustrate typical behaviors of a supercooled liquid (Figure 2(a)). At volume fraction \( \phi = 0.560 \), the MSD exhibits ballistic behavior on short time scales, subdiffusive behavior (power-
law scaling such that \( M(t) \approx t^\alpha \) with \( \alpha < 1 \) on intermediate time scales (\( t \approx 1–10 \)), and an upturn (\( t \approx 10 \)) and gradual crossover to diffusive scaling (\( M(t) \approx t \)) on long times (\( t > 10 \)). These features are common in dense particle suspensions and deeply supercooled liquids. The subdiffusive behavior at intermediate times arises from “cage-rattling”, in which particles are localized inside the cages formed by their surrounding neighbors. The upturn observed for liquid but not glassy samples indicates the onset of local rearrangements that eventually accumulate and allow particles to escape the cages formed by their neighbors.\(^{33} \)

Cage escape is associated with the \( \alpha \)- (or terminal-) relaxation process in the liquid and occurs on characteristic time scale \( t_{\alpha} \).

Upon increasing \( \phi \), the subdiffusive region becomes more pronounced and persists for longer duration, reflecting enhanced caging and slowing of the dynamics.\(^{30,33,34} \) The upturn in \( M(t) \) eventually moves outside of the observation window for \( H > 0.585 \), indicating that the systems behave as glassy solids on the simulated time scales. For a less-confined system (\( H = 8.00 \)), the MSDs also become more subdiffusive as \( \phi \) is increased and are larger at a given \( \phi \) than those measured at \( H = 3.00 \) (Figure 2(b)).

Similar trends are observed upon decreasing \( H \) when the volume fraction is held constant (Figure 3). Interestingly, below a particular value of \( H \), oscillations appear in the subdiffusive caging plateaus (\( t \approx 1–10 \)). These oscillations are not observed in 3D amorphous solids\(^{8,32,43} \) but have been previously reported in 2D glasses, where they have been interpreted as evidence of LWFs.\(^{30,33,34} \)

**Collective Dynamics.** In model systems with LWFs, measured properties often depend sensitively on system size due to the artificial suppression of fluctuations larger than the linear dimension \( L \) of the periodic simulation cell. Although finite-size effects are generally undesirable, they can be rigorously analyzed to detect and characterize LWFs in model systems.\(^{44,45} \) Accordingly, we examine the influence of \( L \) on \( M^{\text{cr}}(t) \) and \( \Gamma(t) \).

The cage-relative MSD \( M^{\text{cr}}(t) \) (eq 2) characterizes particle displacements in the reference frame of the center-of-mass of their surrounding nearest neighbors (Figure 4(a)). Hence, it describes the cage-rattling dynamics that arise from the relative motion of particles inside their local neighbor cages. We observe that \( M^{\text{cr}}(t) \) exhibits a crossover from ballistic scaling on short times to diffusive scaling on long times and collapses for all \( L \), indicating that the local dynamics do not exhibit finite-size effects (Figure 4(b)).

By contrast, the contribution remaining after \( M^{\text{cr}}(t) \) is subtracted from the MSD, \( \Gamma(t) \) (eq 4), retains information about collective translational particle motions (Figure 4(a)). This function varies with system size on intermediate time scales, increasing with \( L \) for a given observation time \( t \) (Figure 4(b)). This system-size dependence indicates the presence of LWFs with characteristic size >\( O(L) \).\(^{30,34} \)

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**Figure 3.** Mean-square displacement \( M(t) \) functions for systems with various confinement parameters \( H \) and particle volume fractions of (a) \( \phi = 0.570 \) and (b) \( \phi = 0.600 \). The arrow indicates the direction of decreasing \( H \). The confinement parameters in (a) are \( H = 1.00, 1.20, 1.34, 1.70, 2.00, 2.50, 3.00, 4.00, 5.00, 6.00, 7.00, 8.00, 9.00, 10.00, 12.00, 13.00, 14.00 \). The confinement parameters in (b) are \( H = 2.00, 2.50, 3.00, 4.00, 5.00, 6.00, 7.00, 8.00, 9.00, 10.00, 11.00, 12.00, 13.00, 14.00 \).

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**Figure 4.** (a) Mean-square displacement \( M(t) \), cage-relative MSD \( M^{\text{cr}}(t) \), and \( \Gamma(t) = M(t) - M^{\text{cr}}(t) \) for a system with (\( \phi, H \)) = (0.570, 3.00) and periodic edge length \( L = 54.3 \). (b) \( M^{\text{cr}}(t) \) and \( \Gamma(t) \) for systems with various \( L \) and (\( \phi, H \)) = (0.570, 5.00).
Logarithmically Divergent Fluctuations. Similar finite-size effects are absent in 3D systems but have been observed in 2D solids where they have been attributed to long-wavelength Mermin-Wagner fluctuations. These LW acoustic modes result in collective translational particle displacements, which are isolated in the function $\Gamma(t)$. Thus, the time scale associated with the local maximum in $\Gamma$ on intermediate time scales is an estimate of the characteristic time scale $t_{LWF}$ at which LWFs emerge, and the corresponding magnitude $\Gamma(t_{LWF})$ provides an estimate of the squared amplitude of the LWFs.

To determine whether the LWFs observed in our quasi-2D systems are Mermin-Wagner-like, we examine the L-dependence of $\Gamma(t_{LWF})$ for systems with $0.550 \leq \phi \leq 0.600$. Specifically, for an elastic medium, the squared amplitude of the Mermin-Wagner fluctuations diverges logarithmically with system size $A^2 \sim B \log(L)$, where the scaling constant $B$ reflects the elastic contributions to the thermal vibrations and is a function of the shear and bulk moduli of the system. In agreement with this expectation, we observe that the increase in $\Gamma(t_{LWF})$ with $L$ is consistent with logarithmic scaling over distinct ranges of system size and $H$ for each isochore (Figure 5(a)), suggesting the presence of MW fluctuations. Further, $\Gamma(t_{LWF})$ varies nonmonotonically with both $\phi$ (constant $H$, Figure 5(a)) and $H$ (constant $\phi$, Figure 5(b)).

We note that the logarithmic scaling law above is derived in the continuum limit for an elastic medium and hence only weakly dependent on $H$. However, confinement drives slowing of dynamics $H$ reduces variation in local structure due to steric and packing constraints. To distinguish the effects of structure on $\Gamma$ and $\Gamma(t_{LWF})$ from those of confinement, we normalized $B$ by a structural metric. To select an appropriate metric, we note that finite variance in the distance between nearest neighbors is a necessary condition to obtain LWFs.

Further, for 2D crystals, the long-time limit of $M^{ck}$ is twice the variance of the neighbor–neighbor distance. We therefore hypothesize that a metric based upon $M^{ck}$ can be used to remove variation in $B$ that arises from local structure. We select as the metric $M^{ck}(t_p)$, where $t_p$ is the time at which $M^{ck}$ attains an inflection point in its caging plateau. This metric is invariant with $L$ and has been associated with cage size in 3D systems. Thus, we define a normalized scaling coefficient $B_n = B/M^{ck}(t_p)$.

Along the isochores $\phi = 0.570$ and $\phi = 0.600$, $B_n$ increases monotonically as $H$ is decreased (Figure 6(b)). This monotonically increasing $B_n$ is expected for a system transitioning from 3D, for which $B_n$ would be effectively zero due to the

![Figure 5](https://doi.org/10.1021/acs.jpcb.2c07417)  
2D LWFs leads to the local maxima in $\Gamma(t_{LWF})$ and $B_n$ at an intermediate value of $H$. 

Along the $\phi = 0.600$ isochores, by contrast, all samples exhibit glassy dynamics, and the elastic coefficient $B$ is weakly dependent on $H$. The relatively small change in $B$ with $H$ likely reflects the weaker variation in local structure at higher $\phi$ due to steric and packing constraints. To distinguish the effects of structure on $\Gamma$ and $\Gamma(t_{LWF})$ from those of confinement, we normalized $B$ by a structural metric.
absence of logarithmic scaling, to 2D LWFs. The similarity in the values of $B_n$ for the two $\phi$ at fixed confinement suggests that normalization by the cage size indeed removes much of the variation in $B$ due to changes in local structure.

To further confirm this idea, we examine the $H$-dependence of $B_n$ in a system for which the local structure is nearly independent of confinement. To this end, we simulated a series of crystals of hexagonally close-packed, monodisperse hard spheres confined between two hard walls. The confined crystals are generated such that the average wall–particle and nearest-neighbor separations are $1.07\sigma$, which ensures that their local structure and dynamics do not vary significantly with $H$. For the crystalline systems, the increase of $\Gamma(t_{\text{LWF}})$ with $L$ is consistent with logarithmic scaling, indicating that they exhibit LWFs. Both the elastic coefficient $B$ and the normalized coefficient $B_n$ of the crystal systems increase monotonically as $H$ is decreased (Figure 6), as expected for a 3D-to-2D transition in LWFs, and are very close in magnitude. Hence, we conclude that the growth of $\Gamma(t_{\text{LWF}})$ with $L$ is indicative of MW fluctuations in quasi-2D confined supercooled liquids and that the nonmonotonic variation of $B$ with $H$ in some samples arises from confinement-induced changes in local structure.

State-Space Dependence. As the confinement length scale $H$ decreases, the nonlocal MW fluctuations are expected to increasingly dominate particle motion. In unconfined 3D geometries, directional correlations between the motions of particles in supercooled liquids and those of their neighbors are small on intermediate to long time scales. By contrast, in a 2D system with LWFs, the displacements of particles are highly directionally correlated with those of the neighbors in their surrounding cage. The transition from 3D- to 2D-like dynamics can be characterized by examining the two-particle correlation function $S_{\text{rec}}$, which quantifies the degree of directional correlation between the displacement of a central particle and those of its surrounding neighbors. The function $S_{\text{rec}}$ assumes values of 1 and $-1$ when the displacement direction and magnitude of the central particle

Figure 6. (a) Non-normalized $B$ and (b) normalized $B_n$ scaling coefficients for selected glassy isochores. For comparison, scaling coefficients are also shown for crystalline systems of confined hexagonally close-packed hard spheres. The average particle–particle and particle–wall surface-to-surface separations for the crystalline systems are $1.07\sigma$ and constant for all $H$. The maximum in $B$ for $\phi < 0.60$ reflects the competition between strengthening LWFs and variation in local structure. The $B_n$ scaling isolates the growth of LWFs relative to variation in local structure.

Figure 7. Contour maps of the coarse-grained directional displacement field $S_{\text{rec}}$ within the contact layers for systems with approximately the same periodic side length $L$ and cage-relative MSD $M^{\text{CR}}$ at the selected lag time but varying levels of confinement $H$. The constant $L$ controls the maximum possible wavelength of density fluctuations, and the constant $M^{\text{CR}}$ controls for variation in local structure. The state points are $(\phi, H) = (a) \{0.59, 13.00\}$, $(b) \{0.57, 4.00\}$, and $(c) \{0.55, 2.00\}$. Darkly colored regions in which $S_{\text{rec}} \gg 0$ exhibit displacements that are correlated in both direction and magnitude. In brief, $S_{\text{rec}}$ is obtained by a sum of Gaussians centered at $r_i$ that smear the $S_{\text{rec},i}$ of each particle. The maximum value for $S_{\text{rec}}$ of 6 reflects the fact that only the Gaussians of the nearest neighbors contribute to the sum. To account for directional correlation alongside significant mobility, the Gaussian variance and height are the MSD and $S_{\text{rec},i}$ of particle $i$, respectively. The lag time of each displacement is $t = 10$ simulation time units, which is approximately the time for the first minimum in $\Gamma^{\text{CR}}$. The extent of spatial correlations decreases with decreasing confinement, i.e., the dark regions are smaller for larger $H$. 

966
i and all of its nearest neighbors are perfectly correlated and anticorrelated, respectively. To isolate the effects of confinement, we control for variations in local structure by examining three state points with different \( H \) but similar \( L \) and equal \( M_C^R \) at lag time \( t = 10 \approx t_{1,\text{SC}} \).

A weakly confined system (\( (\phi, H) = (0.59, 13.00) \)) exhibits modest correlations in directional displacement between neighboring particles (Figure 7). As \( H \) is decreased, the correlations become stronger (i.e., \( S_{\text{sc}} \) increases), and the size of regions of correlated displacement increases. Together, these observations suggest that increasing 2D LWFs in supercooled liquids leads to enhanced collective displacement within regions of particles. This result is consistent with experimental observations of larger displacement correlations in colloids at a 2D oil–aqueous interface compared to 3D systems.

Lastly, to characterize how LWFs vary across the \( \{ \phi, H \} \) state space, we calculated the elastic coefficient \( B \) for supercooled liquids across a range of confinement spanning bulk to quasi-2D systems (Figure 8). LWFs are detectable across a broad range of \( H \). As \( \phi \) is increased from 0.550, state points with measurable values of \( B \) (by our methods) extend to larger values of \( H \). This result is consistent with the expectation that \( B \) (which is a measure of the elastic contributions to the thermal vibrations) should increase as systems become denser and hence more solid-like. Indeed, nonzero values of \( B \) are observed up to the largest \( H \) studied (\( H = 14 \)) for \( \phi \geq 0.580 \). For \( \phi < 0.580 \), our methods fail to detect LWFs beyond \( H \approx 10 \). We note, however, that 2D LWFs can be present and influence dynamics within normal 2D liquids.\(^{27,54} \) Thus, alternative approaches may reveal LWFs in more weakly confined systems.

Along isochores with \( \phi \leq 0.590 \), \( B \) attains a maximum at an intermediate value of \( H \) (as also seen in Figure 6). This value of \( B \) signifies the maximum magnitude of the LWFs induced by finite-size effects. The largest maximum in \( B \) occurs at the lowest value of \( \phi \) examined (i.e., in the most liquid-like system). Further, because LWFs are detectable across a narrower range as \( \phi \) is decreased, the change in \( B \) with \( H \) is steeper. Thus, finite-size effects arising from LWFs are most pronounced for low-\( \phi \) systems in the quasi-2D limit.

While writing this article, Ciamarra and collaborators published a study in which they investigated the change in dynamics for increasingly confined systems containing a 80:20 Kob-Andersen Lennard-Jones glass-forming mixture.\(^{25,56} \) Using periodic boundary conditions to avoid layering in the confinement direction, they showed that LWFs affected the relaxation dynamics of quasi-2D supercooled liquid by examining their dependence on lateral size, similar to what is reported here. Further, they compared the time scales associated with structural relaxation and the longest-wavelength vibrational modes and thereby determined whether systems exhibited 2D-like (in which dynamics were dominated by LWFs) or 3D-like (in which dynamics were dominated by structural relaxation) behavior. Interestingly, ref 55 also examined the behavior of the Kob-Andersen mixture confined in a slit geometry, but they did not report observing a nonmonotonic dependence of MW fluctuations on \( H \), as was found in our study. This apparent discrepancy may be due to the fact that the particle volume fraction was not held constant as \( H \) was varied in their study nor were the effects of varying \( \phi \) systematically explored. Alternatively, it may arise from the use of different model liquids. In either case, our study presents a complementary analysis of a different model, in which the effects of both particle volume fraction \( \phi \) and the confinement length scale \( H \) were systematically explored, revealing distinct behavior in different regions of the parameter space.

Additionally, our analysis of LWFs in the quasi-2D systems is based on global dynamical quantities calculated by averaging over all of the particles in the systems. Nonetheless, the dynamics of confined liquids may be heterogeneous and vary as a function of distance from the bounding walls due to the organization of particles into distinct layers.\(^{16} \) Thus, we anticipate that additional insights into LWFs in these systems can be obtained by examining the local dynamics in different particle layers. We expect that the dense contact layers start to dominate the average MSD as \( H \) is decreased. Indeed, our preliminary examination of local dynamics indicates that long-wavelength MW fluctuations may be enhanced in the dense contact layers near the walls relative to those in the center of the pores, but this topic merits further investigation in future studies. Another promising avenue for future study is the development of structural metrics that provide an improved description of the changes in dynamics observed as the confinement length scale is changed. There has been recent success, for example, in identifying metrics that connect the dynamics of 2D and quasi-2D liquids consisting of bidisperse, nearly hard-sphere particles.\(^{56} \) Although these structural descriptors do not appear to be directly extensible to polydisperse systems in this study, we anticipate that similar approaches may prove successful in developing metrics that better account for the variations in the strength of the LWFs with confining length scale than those presented here. A second potential structural metric is the hexatic order parameter, which has been shown to describe melting processes\(^ {57,58} \) and nanoparticle adsorption\(^ {59} \) in two dimensions. Our previous computational study showed that the in-plane relaxation time increased exponentially with the correlation length associated with the hexatic order parameter.
for confinements whose widths were commensurate with the average particle size.\textsuperscript{20} Thus, an interesting avenue for future study is to examine if there is a connection between LWFs and crystalline order.

\section*{CONCLUSIONS}

Using event-driven molecular dynamics simulations, we showed that confined supercooled liquids exhibit long-wavelength MW fluctuations in the lateral directions parallel to the confining walls, similar to those observed in 2D but not 3D systems. Competition between caging and MW fluctuations as the liquids are increasingly confined leads to a nonmonotonic dependence of the fluctuations on the confinement length scale. Thus, the fluctuation amplitude can be modulated by controlling the particle volume fraction and extent of confinement. The results reveal that LWFs are detectable for volume fractions below random close-packed as long as the confinement is sufficiently strong (in Figure 8); below $\phi < 0.58$, bulk behavior appears for $H > 14$. Thus, choosing $H \geq 20$ appears to be sufficient to avoid LWFs for systems far from random close packing (RCP). For $\phi$ closer to RCP, however, we are able to detect LWFs even at widths up to $H = 14$, the maximum examined in this study due to the need to simulate systems with large $L$ to determine the scaling behavior. Although their strength appears to decrease as $\phi$ is increased toward RCP, additional simulations closer to RCP are needed to quantitatively test this speculation.

The detectability of MW fluctuations even in systems where the confinement length scale is an order of magnitude larger than the particle diameter suggests that they may influence the properties of confined liquids in practical settings. Collective displacements in quasi-2D paramagnetic colloids, for example, were found to smear cage rearrangement and enhance particle mobility\textsuperscript{23} and to accelerate viscoelastic relaxation and reduce viscosity.\textsuperscript{29} Thus, improved understanding of dynamical processes in quasi-2D systems may open new routes to tuning the meso- and macroscopic properties of confined liquids.

\section*{AUTHOR INFORMATION}

Corresponding Authors

Jeremy C. Palmer — Department of Chemical and Biomolecular Engineering, University of Houston, Houston, Texas 77204-4004, United States; orcid.org/0000-0003-0856-4743; Email: jcpalmer@uh.edu

Jacinta C. Conrad — Department of Chemical and Biomolecular Engineering, University of Houston, Houston, Texas 77204-4004, United States; orcid.org/0000-0001-6084-4772; Email: jconrad@uh.edu

Author

Ryan C. Roberts — Department of Chemical and Biomolecular Engineering, University of Houston, Houston, Texas 77204-4004, United States

Complete contact information is available at: https://pubs.acs.org/10.1021/acs.jpcb.2c07417

Notes

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