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Crystal Nucleation in Sedimenting Colloidal Suspensions

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Homogeneous crystal nucleation from a metastable hard-sphere colloidal liquid has been extensively studied in simulations and experiments. A 12-order of magnitude difference between simulated and experimental nucleation rates is observed, the origin of which remains a puzzle. Here, we experimentally study crystal nucleation at the single particle level in suspensions of hard-sphere-like colloids under the influence of sedimentation. We find that sedimentation significantly enhances the nucleation rate, but contrary to what was previously thought, this is not due to simple density fluctuations, as the nucleation barriers become independent of the local density in a sedimentating fluid. Instead, we find an enhancement of the local dynamics, which suggests that local convective cells are responsible for shear-enhanced crystallization at low volume fractions.

I. INTRODUCTION

Homogeneous crystal nucleation from a colloidal hard sphere liquid is arguably the simplest crystallization process, encountered when small crystal nuclei form spontaneously in an existing metastable liquid phase1. To understand crystallization kinetics, this simplest case of nucleation has received enormous attention in both simulations and experiments. For the hard sphere (HS) fluid, the only control parameter is the volume fraction occupied by the spheres. Experimentally, the HS system is reasonably well realized with suspensions of monodisperse colloidal particles that interact through a steep repulsive potential2. In recent years, advances in imaging and particle tracking, which enabled the direct observation of crystal nucleation and growth in dense colloidal systems3, have deepened our understanding of crystal nucleation in model HS colloids. However, an important aspect, the absolute crystal nucleation rate, remains contentious.4

According to simulations, the absolute crystal nucleation rate of this system depends dramatically on φ (the effect of polydispersity and the slight softness in the interparticle potential is taken into account); a 15 order of magnitude increase of the rate is found simply going from φ =0.52 to φ =0.545. This behavior is corroborated by different computational techniques6–7. However, this behavior is not observed in experiments, where a rather weak dependence is found8–9. The comparison between simulated and experimental rates as functions of φ, reveals a 12 order of magnitude difference for = 0.52 that remains to be understood. Notable speculated causes for the discrepancy were, a two-step crystallization process10, difficulties in interpreting the experiments5, and hydrodynamic effects, neglected in the simulations6.

One important difference between experiments and simulations is the presence of gravity, which can induce sedimentation in real life experiments, but is typically not taken into account in simulations. Despite observations that colloidal crystallization differs between micro-gravity and normal gravity experiments, how sedimentation affects the kinetics of the crystals remained unclear11. Recently, the interplay between sedimentation and crystallization was studied via Brownian dynamics simulations12. The findings showed that in systems influenced by sedimentation, gradients in volume fraction make that locally in the sample a high φ (φ ≈ 0.56) is reached, for which nucleation is rapid: from these regions crystallization starts. What has to be explained is the effect of hydrodynamic interactions, not just on the crystallization process itself, but also how they couple to the gravitational field to alter the fluctuations in a suspended fluid. The role of long-range hydrodynamic interactions on the bare crystallization process remains controversial, with studies suggesting either an enhancement13 or a suppression of the crystallization rate14.

In the limit of high Peclet numbers (where gravitational effects are much stronger than thermal effects), experiments on monodisperse polystyrene particles15 have shown that long-range correlations are formed, where the amplitude of the velocity fluctuations (relative to the bare sedimentation velocity) grows with increasing packing fraction. Fluctuations are non-universal, and depend strongly on the shape of the cells16, and point to the importance of both density fluctuations and recirculations that originate locally in the sedimenting fluid. The effect of shear has also attracted a lot of interest17, showing that the crystallization rate is a non-linear function of shear rate18–20. Thus, within our current understanding, the non-equilibrium fluctuations that arise from the coupling between hydrodynamic interactions and the gravitational field have the potential to alter the crystallization process, either by an enhancement of density fluctuations, or by shear-induced crystallization. In the following we will show how to disentangle these two effects.

In this paper, we investigate experimentally what the influence of sedimentation is on crystal nucleation in HS
colloids. To that end, we study both the crystallinity and the density at the single particle level by means of confocal microscopy. So far, few quantitative studies of the effect of gravity on crystal nucleation that stem from real time and space experiments have been reported\textsuperscript{21}. We find that crystal nucleation is significantly enhanced during sedimentation, irrespective of the initial volume fraction at which the system is prepared. Furthermore, we report that contrary to what was suggested before\textsuperscript{12}, the nucleation barrier does not depend significantly on the local density, showing that density fluctuations are not at the origin of the enhancement of crystal nucleation. During this regime, we observe an enhancement of the mean-square displacement of the particles, which originates from the interplay between thermal diffusion and inhomogeneous convective cells.

II. METHODS

We use sterically stabilized, fluorescent polymethylmethacrylate particles dispersed in a mixture of cis-decalin and cycloheptyl bromide (CHB), with 260 \( \mu \text{M} \) tetrabutylammonium bromide salt to screen the weak electrostatic interactions between the particles\textsuperscript{22}. The hydrodynamic radius of the particles is \( r=0.97 \mu \text{m} \) and the polydispersity in size is 2.1\% (see Sup. Mat.). Details on the particle synthesis and the phase behavior of the particles can be found in Ref.\textsuperscript{23}. We mix cis-decalin and CHB to adjust the density mismatch \( \Delta \rho \) between the solvent mixture and the particles, thereby changing the buoyant mass and the Peclet number \( \text{Pe} \) that describes the relative importance of diffusion compared to sedimentation and is defined as \( \text{Pe} = \frac{4 \pi \rho \Delta \rho r^3}{3 k_B T} \), where \( g \) is the gravitational acceleration, \( k_B \) the Boltzmann constant and \( T \) the absolute temperature. The Peclet number also gives the ratio of the particle size to the gravitational length \( l_g \). We prepare samples with solvent densities of 1.1g/ml (mismatched, \( \text{Pe}=0.9 \)), viscosity 2.3mPas estimated from values reported in Ref.\textsuperscript{22}. Samples with volume fractions in the range 0.52-0.56 are prepared by diluting sediments centrifuged to random close packing\textsuperscript{24}; in this reference also the systematic errors in volume fraction are discussed, which can be several percent. For the density matched sample (viscosity 2.217mPas), the solvent mixture was carefully adjusted until there was no visual sign of sedimentation or creaming after centrifuging for 6 hours (at 21°C, at 3000g). The cells for microscopy are completely filled with the suspension to avoid any suspension-air interface that we found promotes heterogeneous nucleation.

We shear melt the systems by stirring them with embedded stirrer bars, which provides a reproducible initial disordered state\textsuperscript{25}. We typically monitor \( \sim 55000 \) particles in a 70x70x80 \( \mu \text{m}^3 \) volume away from the walls of the cell and more than 30 \( \mu \text{m} \) away from its bottom to avoid boundary effects\textsuperscript{26}, the first 23 hours after melting by performing one scan per hour. We determine the particle positions from the confocal images using a standard particle locating software\textsuperscript{27}. We perform a local bond-order analysis on the particle positions, using the spherical harmonics analysis introduced by Steinhardt et al.\textsuperscript{28}, and apply the criteria established by Frenkel and co-workers\textsuperscript{29} to identify crystal particles, as well as a cluster algorithm to identify individual nuclei\textsuperscript{30}, in each configuration. To avoid boundary problems in the analysis, we only consider particles that are at least 2 \( \mu \text{m} \) away from the boundaries of the scanned volume. It is worth noting that this choice can underestimate the size of the nuclei in the case when these are partially inside and outside the imaged volumes, inducing a systematic error that equally affects all measurements. However, we anticipate that this does not have a strong effect here, since there seem to be few nuclei at the borders, when looking at the x-z reconstructed images in time.

III. RESULTS AND DISCUSSION

For all systems influenced by sedimentation, we already see the appearance of crystalline seeds during the first hours after shear melting. Figure 1 shows a typical example of the crystal nuclei found in a mismatched system at different times after the beginning of the measurement that we start 20 minutes after melting. On the other hand, in the density matched system of volume fraction 0.52 we waited 6 months and we still do not observe any crystallinity. Our experiments with sedimentation then give results similar to both previous experiments and the simulations with gravity included. However, if we carefully density match our system, the nucleation rate is orders of magnitude lower, and the bound that follows from not observing any nucleation during 6 months is already much closer to the original simulation without gravity. This suggests that sedimentation greatly enhances the nucleation events and may be at the origin of the earlier discrepancy between experiments and simulations.

We calculate the nucleation rate in our mismatched systems by defining the rate \( k \), as \( k = \frac{1}{2 \tau} \), with \(< \tau > \) the time we waited to find a nucleus of size 70 particles in the measured volume \( V \). The size is chosen as the size after which we typically observe crystalline growth in our experiments. We measured the time that passed until the appearance of a nucleus of size 70 particles in each sample. This was the biggest nucleus that we observed for the sample of volume fraction 0.52, 6 hours + 20 min after melting. For the 0.54 and 0.56 samples, we observed 70 particle nucleus \( 1h + 20min \) and \( 20min \) after melting,
FIG. 1: Time evolution of the crystallinity in a non-density matched system (Pe=0.9); the volume of the box is the same as the volume 70x70x80 $\mu$m$^3$ that we measure, far away from the walls and the bottom of the cell, which is rough to prevent heterogeneous nucleation there. Red dots correspond to the crystal particles identified (a) one hour after the beginning of the measurement and (b) after four hours. Green dots correspond to the particles identified as liquid; their size is reduced for clarity. The arrow indicates the direction of gravity $g$.

respectively. This allows us to obtain a lower bound on the rate, which does not change significantly with the choice of this size.

In Figure 2, we compare our nucleation rates under gravity (normalized with the short-term diffusion coefficient, $D_0$, and the particle diameter, $d$) with previously measured $8, 9, 31, 32, 33, 34$ and predicted $6, 7, 35$ rates. In experiments, $D_0$ is typically calculated using Stokes-Einstein formula at dilute concentrations, $D_0 = \frac{k_B T}{6 \pi \eta r}$, with $\eta$ the viscosity of the total solvent and $r$ the hydrodynamic radius. For the density mismatched sample initially prepared at $\phi = 0.52$ the rate is of the order of $10^{-8}$ in the dimensionless units of Figure 2. For the density matched sample of $\phi = 0.52$ that has not shown nucleation events so far, we calculate a bound for the rate, which corresponds to $10^{-11}$. Therefore, we find that there is at least a 3 order of magnitude difference between mismatched and matched systems in the lower liquid-crystal coexistence regime, in line with what was discussed in the introduction. This is further supported by our experiments in a sample with a higher density mismatch (solvent density 1g/ml, that corresponds to Pe=1.8) prepared at $\phi = 0.52$, where we again observe enhanced nucleation compared to the density matched case (not shown here). Moreover, when looking at the rates from previous (light scattering) experiments, shown in green in Figure 2, one notices roughly two trends: the upper curve where the measured rates approximate the rates predicted by the simulations with gravity in Ref. $12$ and the lower curve where the measured rates are closer to the rates predicted by the gravity-free simulations in Ref. $6, 7, 35$. The data following the former stem from experiments affected by sedimentation, with Pe number 0.43 (Ref.$8$) and 0.24 (Ref.$32$), while the data following the latter stem from density matched experiments. Indeed, gravitational effects affecting previous experiments explain the enhancement of the nucleation rate, while improved density matching already shows a decrease in the rates in the low liquid-crystal coexistence regime. To our knowledge, the only measured rates not in line with this observation are the ones stemming from the density matched experiments reported in Ref.$34$, also presented in Figure 2.

FIG. 2: Dimensionless crystal nucleation rates as function of $\phi$ as predicted by simulations and measured in experiments. Red lines correspond to gravity-free simulations, green symbols to previous experimental rates, the blue line to simulations under gravity and the black line to systems influenced by sedimentation studied here. The Peclet number $Pe = 0.9$ here; it is 0.43 in Ref.$8$, 0.24 in Ref.$32$ and more than an order of magnitude smaller in the other experiments.
To gain insight into why nucleation is enhanced in the mismatched systems, we first look at the number of crystal nuclei, computed as the total number of nuclei detected inside the volume at each scan; we find that it increases considerably during the first hours after melting, and thereafter decreases, as shown in Figure 3a (see also Sup. Mat. for the time evolution of both the crystalline particles and the largest nucleus size). It is worthwhile noting that these results concern all the nuclei, not just the supercritical ones. To understand how the changes in crystallinity relate to sedimentation, we first study the density order parameter $\rho_i$. This order parameter is calculated via Voronoi tessellation, which assigns each particle a local volume $v_{vor}=\frac{1}{\rho_i}$, called the Voronoi volume; the Voronoi volume is then used to measure the local volume fraction, $\phi_i=\frac{V_p}{v_{vor}}$, with $V_p$ the volume of the particle. For all mismatched systems, we observe a gradual decrease of the local volume fraction with height during the first hours of the measurements and the establishment of a z-profile after six hours (Figure 3b), which indicates that sedimentation has stopped. Here the height is measured within the observation volume, which is kept fixed at about 30 microns from the bottom of the container.

We find that crystallization is much enhanced during sedimentation, and slows down when the systems have settled at longer times; after 5 hours the density profile hardly changes anymore, and the number of nuclei starts to decrease. From the time evolution of the 2D (x-z) projections of the systems we notice that the growth of the existing nuclei is suppressed once sedimentation stops; the bigger nuclei become smaller and the small pre-critical nuclei dissolve. It should be noted that we observed qualitatively similar behavior in samples with higher density mismatch (Pe=1.8). The suppression of growth once sedimentation stops has been discussed before in simulations under gravity, see Ref. 12. This gives a strong indication that the increased nucleation probability is due to hydrodynamic effects during sedimentation, rather than changes in the local volume fraction (and consequently local changes in the supersaturation). The big and anomalous fluctuations of the z-dependent local volume fraction profiles within each time frame are also likely due to collective hydrodynamic effects; it is generally invoked (in simulation studies) that at small Pe numbers (Pe< 1) hydrodynamic interactions should be irrelevant, but there are studies that found important effects already for very low densities 36. The non-equilibrium z-dependent profiles extracted from our measurements seem to agree with the latter and probably cause convection. On the contrary, in the case of diffusive settling (sedimentation without hydrodynamics), a smooth decrease of the local volume fraction with height was observed 12. We notice that no correlation between the density inhomogeneity and the spatial distribution of crystalline nuclei is observed, as in Fig. 1.

Another, rather surprising, observation is that nucleation events happen at all heights inside the measured volumes that correspond to different volume fractions, during sedimentation. This suggests that the barrier for nucleation to occur does not significantly depend on the volume fraction, which is in sharp contrast with simulation results so far; in gravity-free simulations, the barrier was shown to drop from 40 to 20 ($k_B T$) just going from volume fraction 0.5207 to 0.5343 5 and simulations with gravity also showed that there is an optimum volume fraction $\phi \approx 0.56$ for nucleation to start. To test this hypothesis, we extract from experiments the size distribution functions of crystalline nuclei. We identify crystalline particles via bond-order parameters as described in Section II, and compute the average volume fraction dependence of the cluster size distribution, $N_n$, by grouping clusters together based on their local average density, irrespective of the global volume fraction or Peclet number. Examples of $N_n$ are shown in the inset of

![Figure 3](image-url)
Figure 4. We average configurations at least 8 hours after shear melting, when the density profile has settled. In the framework of Classical Nucleation Theory (CNT), the negative logarithm of the size distribution function is interpreted as an effective free energy \( \Delta G(n) = -k_B T \ln(N_n) \). In this context, the barriers are a measure of the probability function to find a nucleus of a certain size at each local volume fraction. For a discussion on the applications of CNT to equilibrium and non-equilibrium systems, see Ref. 5 and Ref. 37, respectively. Our results for the nucleation barrier at local volume fractions in the range 0.52-0.58 are shown as symbols in Figure 4; to our knowledge nucleation barriers stemming from experiments are shown here for the first time. We find that the nucleation barriers have only a weak dependence on the local volume fraction. These nucleation barriers explain the weak \( \phi \) dependence and the rapid nucleation seen in previous experiments in Figure 2. To contrast this result with the expectations in absence of gravity, we run Monte Carlo simulations of hard spheres at the same volume fractions considered above. To extract free-energy barriers we use a variant of Umbrella Sampling, called CNT-US, which is described in detail in Ref. 38. The results are shown in Fig. 4 with dashed lines. As expected, the volume fraction dependence of the barriers is much stronger than what we find in experiments. Interestingly the experimental barriers are close to the numerical barrier computed at \( \phi = 0.54 \), which is the same volume fraction below which the discrepancy between simulations and experiments appears in Fig. 2.

Since the density fluctuations are not responsible for the enhanced nucleation during sedimentation, we would like to gather insight into the nature of the hydrodynamic effects; we acquire two dimensional images of our systems over a single plane. This allows to track the particle trajectories during sedimentation in the mismatched system and compare to the trajectories in the matched system. From the trajectories, we calculate the mean squared displacement \( \langle \delta r^2 \rangle \) to quantify the particle mobility. On a log-log scale, we find a slope of 1 for the density matched system and a slope of \( \frac{3}{2} \) for the density mismatched system as seen in Figure 5. This type of super-diffusive behavior typically arises from the interplay between thermal diffusion and convective cells.

![FIG. 4: Free energy in terms of thermal energy as a function of nucleus size for local volume fractions in the range 0.52-0.58 from experiments with gravity (symbols) and simulations without gravity (dashed lines). The inset shows the cluster size distribution corresponding to the free energy profiles that are shown by symbols in the main panel.](image)

**FIG. 5: Mean square displacement perpendicular to gravity as a function of time for a density mismatched system and a density matched system.**
where spatially inhomogeneous (but correlated) velocity fields alter the power-law scaling of mass transport. So hydrodynamic interactions dominate the Brownian forces in our sedimenting suspensions. This in general does not come as a surprise in the case when a system is subjected to a gravitational field, and was expected here already from the z-dependent local volume fraction profiles in Figure 3b. It is a clear indication of flow (convection) inside the system in the direction perpendicular to gravity, which leads us to speculate that the observed enhancement of nucleation rates originates from shear-induced crystallization.

IV. CONCLUSION

We examine crystal nucleation in HS colloids at the single particle level. We find that sedimentation of the particles, due to the density mismatch between particles and solvent, greatly enhances the nucleation events and the nucleation rate. In addition, we show that the nucleation barrier does not significantly depend on the local volume fraction. This greatly differs from simulation results under gravity that explain the changes in nucleation rates as being due to the changes in local volume fraction; our results rather highlight the dominant role of hydrodynamics in sedimenting suspensions.

V. SUPPLEMENTARY MATERIAL

See the supplementary material for the determination of the size polydispersity and additional data on the crystallinity at different volume fractions as a function of time.

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REFERENCES


