Direct Imaging of Vibrations in Colloidal Crystals: In Equilibrium and in a Steady Drift

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ABSTRACT: Crystals of colloids, micron-size particles in a solvent, typically contain high concentrations of structural defects, limiting their applicability in self-assembly of metamaterials. Defects and grain boundaries play an important role for most properties of these crystals. Most previous research of colloidal crystals, by experiment and theory, focused on spatially averaged vibrational spectra: the differences in local environment between the bulk crystal particles and those at a grain boundary were typically neglected. We employ direct confocal microscopy and recent more accurate particle tracking algorithms to study the potential wells of individual particles in thermally vibrating quasi-two-dimensional colloidal crystals. We demonstrate that the energy landscape probed by a particle sensitively depends on its local environment. Furthermore, we emphasize the commonly neglected role of slight out-of-equilibrium drift of colloidal crystals, demonstrating that particle vibrations depend significantly on the drift velocity, so that the drifting crystals are softer, allowing an effective “drift temperature” to be defined.

INTRODUCTION

Crowded colloids of a low polydispersity,1 thermalized in a molecular solvent by Brownian motion, minimize their free energy by crystal formation.2 Colloidal crystals occur in natural sediments and may possibly play an important role in biomedicine, where ordered arrays of virus particles spontaneously self-assemble inside an infected cell.3 Colloidal crystals are widely studied as potential self-assembled templates for photonic band gap materials, promising a wide range of future photonics applications.4,5 Recently, colloidal crystals have also been employed in nanosphere lithography.6

Colloidal dynamics is overdamped by the viscous drag of the suspending medium, contrasting with the dynamics in simple atomic solids. Yet, colloidal physics mimics many important aspects of the collective phenomena occurring in atomic and molecular systems. Moreover, with the relevant time and length scales in colloidal systems being much longer than with atoms and molecules,7 very detailed experimental information on colloids can be collected in real motion and with a single particle resolution, employing direct optical microscopy.8−14 In particular, intense studies of vibration spectra in colloidal crystals have been recently carried out,15−21 aiming at the fundamental understanding of the elastic properties of solid matter and the development of a common conceptual framework for crystal, glass, and granular matter.8,15,22

Most colloidal crystals are polycrystalline,23 composed of small grains, and have relatively high concentrations of defects and grain boundaries (GBs).9,20 This fact limits the applicability of colloidal crystals in photonics24 and challenges the distinction between colloidal crystals and glasses.22 Moreover, localized vibrations develop at defect locations,9,20 reminiscent of dynamic and elastic heterogeneities in glass matter.24,25

Thus, the scattering techniques15−17,21 limited to probe only the spatial average of the lattice vibration dynamics, provide an incomplete information for colloidal crystals, where spatial inhomogeneity is non-negligible. Moreover, in many of the optical microscopy studies of colloidal crystal vibrations the focus is on the density of states of the vibration modes, with less attention being devoted to the actual real-space motion of the particles. Almost without exceptions,9,12,13 these studies employ the displacement covariance analysis (DCA).8,10,17,18,20 The DCA approach is based on the harmonic approximation of the interparticle potentials, which may not be valid for the colloids.15 Furthermore, due to the complexity of the DCA, it is prone to numerical artifacts.27,28 The influence of common experimental issues, such as minor sample drifts, sliding of GBs, and relaxation of metastable states is currently under investigation.20,28 As a result, the potential wells of individual particles in colloidal crystals are still not fully characterized. In particular, the relation between the shape of the potential well and the local structure is not yet established.

To overcome these limitations of the previous studies, we carry out real space microscopy studies, correlating the local structure about each particle with its dynamics. In particular, we image by direct video-rate confocal microscopy the thermal motion of (many thousands of) individual colloidal spheres, forming a quasi-two-dimensional crystalline lattice (see raw confocal image in Figure 1). The distribution of displacements of a particle, with respect to its equilibrium position, allows the shape of its potential well to be characterized. We demonstrate
that the potential well depends on the local environment of the particle within the solid; in particular, the wells next to defects and GBs are appreciably shallower. Furthermore, we demonstrate that particle vibrations are significantly modified in the presence of even a minor drift of the crystal through the solution, emphasizing the nontrivial role played by structural heterogeneities in these crystals. 14,18,28

■ EXPERIMENTAL SECTION

To form a colloidal crystal, we suspend PMMA (poly(methyl methacrylate)) spheres, fluorescently labeled by Nile Red dye for confocal imaging, in dodecane (CH3(CH2)10CH3, Sigma-Aldrich, ≤ 3%); the particle volume fraction is ~3%. The particles are sterically stabilized by a poly-12-hydroxystearic acid (PHSA) monolayer, minimizing the van der Waals attractions18,22 so that the interparticle interactions are dominated by hard repulsions. 29 The synthesis of these particles and their properties are described in the literature.18,22,29–31 The synthesis of PHSA and its properties are described in the literature30 as well. We determine the diameter of our particles by dynamic and static light scattering as σ = 2.4 μm; the size polydispersity of our particles is ≤4%. The colloidal suspension is loaded into a rectangular Vitrocom capillary (0.1 × 2 × 50 mm) and sealed with epoxy glue. The density mismatch between PMMA and dodecane is Δρ ≈ 0.3 g/cm³, so that the particles rapidly sediment, forming a crystalline bilayer at the bottom of the sample. In equilibrium conditions, this system is fully dominated by the gravity. In particular, the equilibrium sedimentation length32 δ0 = kBT/Δρvpg, where vpg = σg/6 is the particle volume and g = 9.8 m/s², is much smaller than σ, so that the thermal excitations are insufficient for a particle to leave the horizontal plane. To coat the capillary by PHSA adsorption, we add a small concentration (~4% w/w) of PHSA to the suspension, so that particle sticking to the capillary walls can be eliminated without changing significantly the colloid pair potentials. 33 Where complete coverage by PHSA has been achieved, even a tiny tilt of the sample led to a measurable drift of the crystalline layer, driven by the tiny sample-parallel component of the ambient gravity. Drift velocities were varied from 0.2 to 12 nm/s by accurately tuning the tilt angle of the sample; for that purpose, the sample was mounted on a special tilt stage, allowing a resolution of 0.05°. Where any drift of the crystals was to be avoided, we employed samples with an incomplete PHSA coverage. In these samples, while particles in the microscope-illuminated area were free to vibrate, enough particles were stuck to the capillary bottom outside of the illuminated area, so that any drift of the crystals was blocked.

To image our colloids, we employ the Nikon AIR confocal laser scanning microscope in the resonant scanning mode, with an oil-immersed 100× Plan Apo objective. To achieve sufficient statistics, the 512 × 512 frames are collected for about 30 min, with a digital resolution of 0.17 μm/pix at ~4 fps. We detect the center positions (x, y) of the particles within the bottom two-dimensional (2D) layer of our bilayer crystal, employing a modified version of the PLuTaRC image-analysis algorithm,34 based on the algorithm of Crocker and Grier.35 An iterative algorithm is employed36 to minimize the effect of the finite pixel size on the accuracy of particle localization. The accuracy of this algorithm in 2D has been previously estimated36 as ~4 nm, allowing small particle displacements to be accurately measured. The area fraction η of the particles in the sample is obtained directly from their number density n: η = πσg²/4. Finally, to follow the displacement of our particles between the subsequent frames, we track the particles by proximity.35 The fact that all samples dealt with in the current work are crystalline simplifies particle tracking; still, the tracked trajectories were visually examined, to confirm their validity. While particles next to the boundary of the illuminated area move occasionally out of the frame and could not be tracked, a typical experiment yields full 30-min-long trajectories for more than 10³ particles. After tracking, particle positions have been derivered by subtracting the average displacement of all particles; thus, the rest of the analysis is based only on particle displacements with respect to the center of mass (CM) of the sample.

■ RESULTS AND DISCUSSION

Crystal Vibrations in Thermodynamic Equilibrium. In the presence of GBs and defects, the structural order is imperfect, varying across the sample. To quantify the local crystallinity around a given particle, we employ the 6-fold symmetric orientational bond order parameter37,40 ψ6 = Z⁶ΣiRe{exp(i6iΔθi)}; here the summation is carried out over the bonds to all Z nearest neighbors (NNs) of the particle and the angle Δθi of the i-th bond is measured in a frame oriented with one of the NN bonds, chosen at random. A particle surrounded by Z = 6 NNs in a perfectly hexagonal arrangement has ψ6 = 1. Decreased ψ6 values are detected near static structural defects. 3,40 In addition, the instantaneous values of ψ6 are also reduced by thermal crystal vibrations, so that even the “perfect” experimental crystals typically have ψ6 < 0.98.

To map the structural heterogeneity in our quasi-2D crystals,38 we employ the time-averaged ⟨ψ6⟩ of all particles. Distinct individual grains are observed to be separated by low-ψ6 boundaries [Figure 2(a)]. On first sight, quite surprisingly, the average bond lengths in the bulk of the crystalline grains seem longer than the ones at the boundaries [Figure 2(b)]. It may seem that this observation contrasts with the paradigm of liquid-like GBs having a lower density compared to the adjacent crystals. 36 However, while glassy boundaries, such as between the three-dimensional (3D) crystalline grains in systems of soft colloids, 30 can possibly be denser than the crystals, in quasi-two-dimensional crystals a different scenario takes place. In

Figure 1. A raw confocal microscopy image of a polycrystalline colloidal lattice. Note the relatively large crystalline grains, separated by a clear grain boundary. The field of view is 85 × 85 μm.
particles are harmonic, as commonly assumed, we anticipate that \( P(r) \propto \exp\left(-\frac{1}{2}k r^2\right) \), where \( k \) is the harmonic constant in units of \( k_B T/\sigma^2 \). However, colloidal crystals exhibit a significant structural heterogeneity (Figure 2), implying that \( k \) values change across the sample. Spatially averaged \( P(r) \) includes \( \exp\left(-\frac{1}{2}k r^2\right) \) contributions with different \( k \), so that \( \log(P(r)) \) dramatically deviates from a linear dependence on \( r^2 \). The true local \( P(r) \) must be in a much better agreement with the harmonic approximation. To demonstrate this fact, we bin the \( P(r) \) data by the \( \langle \psi_6 \rangle \) values of the vibrating particles. Thus, only particles in regions of similar crystallinity are averaged together, so that the heterogeneity of \( k \) in each of such \( P(r) \) averages is reduced. The resulting \( P(r) \), for a range of different \( \langle \psi_6 \rangle \) values, are shown in Figure 3(a), plotted as a function of \( r^2 \).

Figure 3. (a) The probability \( P(r) \) for a particle displacement in a crystal is plotted on a semilogarithmic scale, as a function of \( r^2 \). On this scale, a linear scaling would correspond to the effective potential well being perfectly harmonic. The data are binned by particles’ crystallinity \( \langle \psi_6 \rangle \) (see curved arrow). The data for \( \langle \psi_6 \rangle = 0.95 \) are zoomed-in in the inset, demonstrating the harmonic regime. The error bars are omitted for clarity, with even the largest statistical uncertainties (in the bottom right angle of the plot) reaching only \( \sim 30\% \), of the same size as as the point scatter within each of the data sets (see Supporting Information). (b) The dependence of the apparent harmonic constant on the time span \( \tau \) of the trajectory (scatter) is fitted by \( k(\tau) = C_1(\tau/\Delta \tau - 1)^{C_2} + k(\tau \to \infty) \) (purple line), where \( \Delta \tau \approx 0.26 \) s is the time interval between frames, \( C_1 > 0 \) and \( C_2 < 0 \) are arbitrary constants, and the true harmonic constant (dashes) is obtained in the long-\( \tau \) limit. (c) Both in experiments (solid lines) and in MC simulations (dashes), the harmonic constant increases with the local crystallinity \( \langle \psi_6 \rangle \). For a given \( \langle \psi_6 \rangle \), a higher \( k \) occurs in a denser sample (see legend). The slope of the simulated data is higher than in the experiments, suggesting that the GBs in the experimental samples may have a nonlocal effect on the elastic properties of the samples. The experimental statistical errors are only slightly larger than the line width.

To test the impact of the crystals’ structural heterogeneity on particle dynamics, we track the trajectories of all particles. In nondrifting dynamics, such as in Figure 2, all trajectories correspond to vibrations of particles about their average lattice positions. For our range of particle area fractions \( \eta \) and interparticle potentials, the distribution of particle displacements \( r \) is rotationally symmetric within the available statistics. The radial average \( \langle \psi_6 \rangle \) of the distribution decays with \( r \), indicating that the energetic penalty for large displacements is significant. If the effective potential wells probed by the these quasi-2D crystals, a significant disorder in plane-normal \( z \)-coordinates of the particles at the GBs leads to a decreased in-plane separation between the particle centers; particles capable to move in the \( z \)-direction can have their \( x,y \)-coordinates closer together. We confirmed this mechanism by a full 3D confocal reconstruction of the sample, where the \( z \)-coordinates of the particles at the GBs are observed to be disordered and the average \( z \) is larger, compared to its value within the adjacent crystalline grains. The motion of GB particles along \( z \) makes the particle tracking more challenging: with the currently employed high-quality 2D-imaging of particle positions (Figure 1) limiting our image acquisition rate, we ultimately lose track of a particle, once it leaves the focal \( xy \) plane. In Figure 2, particles tracked in different time periods are overlaid. Thus, the GBs, where the motion along \( z \) is significant, appear more crowded; importantly this artificial crowding has no effect on instantaneous particle separations and \( \psi_6 \) values, the averages of which are indicated by colors in Figure 2(a,b).

Figure 2. (a) The time-averaged particle positions in the quasi-two-dimensional colloidal crystal. The color of each particle indicates its average local bond-orientational order parameter \( \langle \psi_6 \rangle \). Note the significant structural heterogeneity, with the crystal grains separated by disordered grain boundaries (GBs). (b) The same crystal, with the color indicating the local in-plane density of particle centers. The increased density (shown in green) at the GBs is due to the increased fluctuations of plane-normal particle positions outside of the crystalline grains. By exploiting the freedom in the (plane-normal) \( z \) direction, \( (\text{g})r\text{ains} \). By exploiting the freedom in the (plane-normal) \( z \) direction, \( (\text{g})r\text{ains} \).
The true thermodynamic increase of the apparent potential wells of the diatomic gas lead to an incorrect probing by the particles. Clearly, such stretching of the particle displacements with respect to the average (dedrifted) value of the distribution is stretched in a certain direction in the plane of all particles by their spatial proximity (instead of the currently used \( ⟨r⟩ \) -binning) would not lead to a similar agreement with the harmonic approximation. Remarkably, for a given range of \( r \), the deviations of \( P(r) \) from the harmonic approximation are the smallest for particles located in the high-crystallinity regions \( \{ \text{green diamonds in Figure 3(a)} \} \), suggesting that the appealing mean-field concept of a particle in an effective potential well loses its meaning for the highly disordered spots of the sample, where the effective potential is itself changing on a similar time scale.

For an accurate determination of the local \( \kappa \), the trajectory of each particle should be sufficiently long, allowing all accessible regions of the potential well to be probed. When the trajectory is too short, some regions of the well are not probed, so that in these regions \( P(r) = 0 \). Thus, finite trajectories overestimate the value of \( \kappa \), as shown in Figure 3(b), where \( r \) is the time span of a trajectory. The limiting values \( \kappa(\tau \to \infty) \) corresponding to the true thermodynamic \( \kappa \) are shown in Figure 3(c). While the increase of \( \kappa \) with \( ⟨r⟩ \) and \( \eta \) is not surprising, our Monte Carlo (MC) simulations of perfect crystals of hard disks (see Supporting Information) yield a higher \( \psi \)-slope for a similar \( \kappa \) (cf. green dashed simulated curve in Figure 3(c) to the experimental data sets). While Coulomb forces may increase the effective radius of the particles, leading to an increased effective \( \eta \) in the simulations (see legend), such inflation of the particles\(^{42} \) is still unable to explain the discrepancy between theory and simulations observed in Figure 3(c). We speculate that the stronger \( \kappa(⟨r⟩) \) dependence in simulations is due to the absence of GBs in the simulated samples.\(^{41} \)

**Vibrations in Drifting Crystals.** To test for a possible nonlocal effect of the grain boundaries, we increase the average grain size in our samples. For that to be achieved, the capillary is fully coated with the PHSA, so that particle immobilization at the bottom of the capillary is fully eliminated. As our crystals are formed by particle sedimentation from bulk suspension, any immobilized particles at the bottom of the sample severely limit structural relaxation during crystal growth, leading to a reduced size of the crystalline domains. However, with the immobilized particles completely eliminated, the crystals drift under very small tilts of the sample-containing capillary. Fine-tuning of the tilt angle (see Experimental Section) allows the drift velocity \( ν_d \) to be adjusted, with the amplitude of the nonequilibrium contributions to the dynamics determined by \( ν_k \). Remarkably, while the interparticle potentials in our system are purely repulsive,\(^{15,34,42} \) all particles belonging to a given crystalline grain drift together, so that the shear is mostly concentrated at the GBs. With the drift velocities of the individual grains being slightly different, the grains are not at rest in the reference frame colloving with the CM of the polycrystalline sample; thus, dedrifting does not bring the effective potential wells of the particles to rest. Grain motion in the CM-frame is complex and may involve rotation (Figure 4(a)), challenging the quantitative analysis of \( P(r) \). In particular, the distribution of particle displacements with respect to the average (dedrifted) particle positions is no longer symmetric under rotation. The distribution is stretched in a certain direction in the \( xy \) plane, leading to a similar apparent stretching of the potential wells probed by the particles. Clearly, such stretching of \( P(r) \) would lead to an incorrect \( \kappa \) if not taking appropriately into account. The apparent potential wells of the different particles stretch, in general, in different directions; see for example Figure 4(a), where the arrows indicate the stretching directions of the individual potential wells. The motion of grains with respect to the CM-frame is very slow, so that on the experimentally relevant time scale, each potential well is only stretched along a certain axis. Thus, the apparent equipotentials assume a shape of a rectangle capped by semicircles (the so-called, “Bunimovich stadium” potential).

To resolve \( \kappa \) for such a potential, we split the trajectory of each particle into time intervals of length \( τ \). For each value of \( τ \), the average \( P(r) \) is calculated. Then, we fit the \( P(r) \) for each \( τ \) by

\[
A \exp\left(-\frac{1}{2}\kappa(τ)r^2\right)I_2\left[-\frac{1}{2}\kappa(τ)r^2\right]^{3/2}
\]

where \( A \) and \( B \) are free parameters and \( I_2 \) is the modified Bessel function of the first kind. This choice is motivated by the radially averaged distribution in a well infinitely stretched along the \( x \) axis, being

\[
P(r) = (2\pi)^{-1} \int_0^{2\pi} \exp\left(-\frac{1}{2}\kappa(τ)y^2\right)dy \propto \exp\left(-\frac{1}{4}\kappa r^2\right)I_2\left(-\frac{1}{4}\kappa r^2\right)
\]

which for a finite stretching of a well should be combined with the expression used with a static well. To the second order in \( r \), the logarithm of our fitting function scales as \( -1/\kappa r^2 \), allowing \( \kappa(τ \to \infty) \) to be identified with the conventional harmonic constant in a nondrifting crystal. We test our approach by fitting a simulated \( P(r) \) of a particle moving in a slowly drifting harmonic well. The fitting expression allows the correct \( \kappa \) of the well to be reproduced, indicating that the distortion of the potential well due to the drift is taken correctly into account; incorrect \( \kappa \) values are obtained when this drift-related correction is neglected. Thus, our approach accounts correctly for the imperfect dedrifting of particle positions resulted by the different drift velocities of the individual grains in a polycrystalline sample. We note in passing, that the same approach may be...
beneficial for other fields where tracking of an individual nanocolloidal particles in drifting wells is employed.\textsuperscript{33}

The drift of crystalline domains through the suspending medium, while increasing the complexity of the $P(r)$ analysis, results also into the emergence of complex out-of-equilibrium phenomena, which may possibly have a nontrivial effect on crystal vibrations. To test for a possible influence of such phenomena on $\kappa$, we measure $P(r)$ for a range of different average drift velocities $v_{d}$. Strikingly, $v_{d}$ as low as $\sim2.5$ nm/s is sufficient for $\kappa$ to drop by about a factor of 2 for a given $\langle \psi_{6} \rangle$ [see Figure 5(a,b)]. This drop in $\kappa$, indicating that the amplitude of particle vibrations grows with the drift, has two contributions: (a) a slight increase of the lattice constant with the drift; (b) energy fluctuations associated with the drift. To decouple these contributions, we extrapolate the data for different samples to $v_{d} = 0$. The corresponding harmonic constants $k_{0}$ scale with $d_{NN}$ [solid symbols in Figure 4(b)], extrapolating to zero at $\eta \approx 0.68$, slightly lower than the melting of a hard-disk hexatic phase.\textsuperscript{44} We employ this $k_{0}(d_{NN})$ scaling and the measured $d_{NN}(v_{d})$ to predict the $\kappa(v_{d})$ variation in an absence of drift-associated energy fluctuations. The resulting $\kappa(v_{d})$ [pink circles in Figure 5(b)] significantly overestimate the experimental values (brown squares), indicating that much of the drift-induced softening of the crystals can be attributed directly to the energy fluctuations, which are missing for the pink circles.

In setting of glass beads in 3D, the hydrodynamically induced spatial fluctuations of particle settling velocity have been demonstrated to thermalize the system, providing an effective temperature scale, called “the gravitational temperature for sedimentation”.\textsuperscript{11,35} In our case, since $\kappa$ is expressed in units of $k_{B}T/\sigma^{2}$, an increase of the effective “drift temperature” by a factor of $\sim1.4$, compared to the common thermodynamic temperature, could justify the observed softening of the drifting crystals, for $v_{d} \approx 4$ nm/s. However, the effective thermalization mechanism is very different from that in the fluid 3D sediments: the settling velocities of our crystals are lower by orders of magnitude, the backflow of the solvent is negligible, and particles belonging to each crystalline domain drift together at roughly the same velocity. Importantly, while the Péclet number\textsuperscript{15} ($Pe$) of an individual PMMA sphere in dodecane, in the relevant range of substrate tilt angles, is of the order of 0.1, a much larger $Pe \approx 10^{5}$ is associated with the motion of a typical crystalline grain, so that the non-Brownian effects may be very significant. In view of the increased motion of the GB particles along $z$, discussed above, we suggest that for a crystalline domain to drift faster by some $\Delta v_{d}$ compared to its neighboring domains, it must occasionally push the particles at its GB up (along $z$) by a significant fraction of $\sigma_{i}$ with the GBs in our samples being curved and the interparticle potentials almost perfectly hard, any relative motion between the crystals would otherwise be prohibited by geometry. Thus, the nonequilibrium sedimentation length $\delta_{neq}$ of the particles at the GBs must be of the order of $\sigma/2$. Expressing the effective thermal energy due to the drift by $k_{B}T_{neq}$ we obtain $k_{B}T_{neq} = \delta_{neq}/\Delta v_{d}g \approx \sigma \Delta v_{d}g/2$, so that $k_{B}T_{neq} \approx 3k_{B}T$. A 2D crystal of $N \approx 900$ particles has $N_{gb} \approx 100$ particles at its boundary. Assuming that the nonequilibrium energy fluctuations of $\sim3k_{B}T$ per GB particle are equally distributed throughout the crystal, each particle in the crystal gets an extra energy of $\sim0.3k_{B}T$, which may possibly account for the reduction of $\kappa/k_{0}$ by a factor of $\sim1.4$ (cf. brown squares and pink circles in Figure 5(b)). A direct measurement of particle dynamics along $z$ at the GBs may allow, in the future, this somewhat speculative, hand-waving argument to be substantiated and the full $\kappa(v_{d})$ to be understood.

Finally, we note that the $\kappa$ values shown in the solid symbols in Figure 4(b) are lower (for a given $d_{NN}$), compared to Figure 3(c) [reproduced in open symbols in Figure 4(b)], where the crystalline grains were smaller. This observation suggests that the GBs, where much of the crystal dynamics is concentrated, may have an unexpected constraining effect on the adjacent high-$\langle \psi_{6} \rangle$ particles: the vibration amplitude of these particles is severely reduced, further emphasizing the complex outcomes of structural heterogeneities in these crystals.

\section{Conclusions}

We have demonstrated by a real-space analysis of particle vibrations in a colloidal crystal that the amplitude of particle vibration is a sensitive function of the local crystallinity. In particular, we detect and quantify the decrease in this amplitude for higher values of the bond orientational crystalline order parameter. Employing a controlled drift of crystals on tilted substrates, we have demonstrated a drift-induced softening of crystal vibrations. Our experiments provide a wealth of local information on particle vibrations, complementing the commonly used displacement covariance analysis. Future experimental and theoretical studies should allow a direct connection to be developed between these two different approaches, en route to a deeper understanding of vibration modes in ordered and disordered solids.

\section{Associated Content}

\section{Supporting Information}

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.6b02070.
Details of the computer simulation procedures and of the drift measurements (PDF)

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Notes
The authors declare no competing financial interest.

■ ACKNOWLEDGMENTS

The authors thank P. J. Lu for sharing his PLuTARC codes and Y. Rabin for the fruitful discussions. This research was supported by the Israel Science Foundation (no. 85/10, no. 1668/10). Some of the equipment was funded by the Kahn Foundation. A.B.S. is partially funded by the UK Engineering and Physical Sciences Research Council grant EP/J007404/1.

■ REFERENCES

(5) Chakim, P.; Kriegs, H.; Petekidis, G.; Fytas, G.; Economou, E. Al2O3 inverse opals.


Supporting Information

Direct Imaging of Vibrations in Colloidal Crystals: in Equilibrium and in a Steady Drift

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Dependence of Thermal Vibrations on Local Environment

To emphasize the dramatic dependence of thermal vibrations on the local environment, we calculate the ratio \( \Pi(r) \) between the \( P(r) \) measured for the local orientational bond order parameter values of \( \langle \psi_6 \rangle = 0.82 \) and \( \langle \psi_6 \rangle = 0.95 \) [Fig. 3(a) of the main text]. The ratios, for a range of \( r^2/\sigma^2 \) values, are shown in Fig. S1. Note, at the right hand side of the plot, a difference of only \( \sim 16\% \) in \( \langle \psi_6 \rangle \) corresponds to a difference of a factor of \( \sim 25 \) in \( P(r) \).

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Figure S1: The probability $P(r)$ for large particle excursions is strongly dependent on the local environment. The ratio $\Pi(r)$ between the $P(r)$ at $\langle \psi_6 \rangle = 0.82$ and $\langle \psi_6 \rangle = 0.95$ strongly increases with $r$. 
Crystal Drift Measurements

The crystal drift velocities, appearing in the main text have been measured by direct confocal microscopy. The drift is almost perfectly monotonic in time, as demonstrated in Fig. S2, where the black solid line are the experimental data and the red dashes are a linear fit. The slope of the linear fit is the drift velocity. All the parts of the analysis in the main text are based on the de-drifted particle coordinates, where the linear trend shown in Fig. S2 is subtracted.

![Graph showing drift velocity over time](image)

Figure S2: A typical plot presenting the drift of a crystal. Note that the drift velocity is almost perfectly constant in time.

Crystal Simulations

To simulate the $P(r)$ in crystals [see Fig. 3(c) in the main text], we employ the Monte-Carlo (MC) algorithm, where the initial state is a perfect hexagonally-packed two-dimensional sys-
tem of hard disks, with cyclic boundary conditions being used. The lattice constant is chosen to be between 1.042 and 1.063, in particle diameter $\sigma$ units, to match the experimentally-relevant range of particle densities. The simulation cell includes $\sim 7000$ particles. A single particle step length is $0.04\sigma$. All particles are relaxed by 1000 MC steps, prior to collecting the $(x(t), y(t))$ data, where $x(t)$ and $y(t)$ are the coordinates of each particle at time $t$. Particle positions are recorded once in 15 MC steps to avoid significant correlations between their subsequent positions.

**Particle in a Well**

To test the performance of the phenomenological expression used for fitting of $P(r)$ in drifting crystals, we fit this expression to simulated $P(r)$ of a single particle in a drifting harmonic well, where the precise $\kappa$ value is known. Our MC simulations allow the particle to make $5 \times 10^3$ relaxation steps, starting from the center of the well, with the $(x(t), y(t))$ coordinates during the subsequent $10^7$ steps kept for the $P(r)$ analysis. The length of a single MC step was chosen as $0.03(k_B T/\kappa)^{1/2}$. In each simulation, we fix the drift of the energy well position; the drift values were in the range between 0 and $3 \times 10^{-4} (k_B T/\kappa)^{1/2}$ per MC step, covering the experimentally-relevant range of $v_d$. As for the experimental data, we split the particle trajectory into time intervals of length $\tau$. For each value of $\tau$, the average $P(r)$ is calculated. Then, we fit the average $P(r)$, for each value of $\tau$ by $A \exp(-\frac{1}{2}\kappa(\tau)r^2)I_0[-\frac{1}{2}\kappa(\tau)r^2]B$, where $A$, $B$, and $\kappa$ are free parameters and $I_0$ is the modified Bessel function of the first kind. The fitted $\kappa(t \to \infty)$ agrees with the value used in the simulation. In this range of $v_d$, fitting the data by either $A \exp(-\frac{1}{2}\kappa r^2)$ or a linear combination of this function with $\exp(-\frac{1}{4}\kappa r^2)I_0(-\frac{1}{4}\kappa r^2)$ (see main text), yields $\kappa$ values which are less precise, supporting our choice of the fitting function in experiments on drifting crystals.