Interparticle Interactions and Direct Imaging of Colloidal Phases Assembled from Microsphere–Nanoparticle Mixtures‡

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We investigate the interparticle interactions, phase behavior, and structure of microsphere–nanoparticle mixtures that possess high size and charge asymmetry.1 We employ a novel Monte Carlo simulation scheme2 to calculate the effective microsphere interactions in suspension, yielding new insight into the origin of the experimentally observed behavior.2 The initial settling velocity, final sediment density, and three-dimensional structure of colloidal phases assembled from these binary mixtures via gravitational settling of silica microspheres in water and index-matched solutions exhibit a strong compositional dependence. Confocal laser scanning microscopy is used to directly image and quantify their structural evolution during assembly. Below a lower critical nanoparticle volume fraction (φnano < φC), the intrinsic van der Waals attraction between microspheres leads to the formation of colloidal gels. These gels exhibit enhanced consolidation as φnano approaches φC. When φnano exceeds φC, an effective repulsion arises between microspheres due to the formation of a dynamic nanoparticle halo around the colloids. From this stable fluid phase, the microspheres settle into a crystalline array. Finally, above an upper critical nanoparticle volume fraction (φnano > φC), colloidal gels form whose structure becomes more open with increasing nanoparticle concentration due to the emergence of an effective microsphere attraction,3 whose magnitude exhibits a superlinear dependence on φnano.

Introduction

Colloidal suspensions enjoy widespread use in applications ranging from advanced materials to drug delivery. By tailoring interparticle interactions, one can design colloidal systems for the assembly of ceramics,4 inks,5 coatings,6 photonic band-gap materials,7–12 and pharmaceutical13 materials. Despite their broad importance, the structure and phase transitions of the fluid, gel, crystalline, and glassy states have primarily been studied in hard-sphere systems that interact only through infinite repulsion on contact.14–20 Recently, these studies have been extended by introducing controlled attractions through either the addition of nonadsorbing polymers21 or temperature change.22 Confocal laser scanning microscopy (CLSM)23 is an invaluable tool in these investigations that allows individual particles to be imaged within a three-dimensional assembly in real space.

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Several pioneering studies have been carried out to directly probe the structure and dynamics of colloidal crystals.\textsuperscript{24,25} glasses,\textsuperscript{20,26} and gels\textsuperscript{27} using CLSM. van Blaaderen and Wiltzius have investigated the structure of colloidal crystals\textsuperscript{28} and glasses assembled from hard-sphere systems.\textsuperscript{20} Weitz and co-workers have studied the homogeneous nucleation and growth of colloidal crystals,\textsuperscript{25} the dynamics of hard-sphere suspensions near the glass transition,\textsuperscript{28,29} and aging phenomena in colloidal glasses,\textsuperscript{30} and the structure of colloidal gels formed via depletion attraction.\textsuperscript{27} Their observations revealed that the gel structure is systematically more open as the depletion attraction between colloids intensifies. Finally, Varadan and Solomon\textsuperscript{22} have investigated the structure of dense colloidal gels formed by temperature-induced attractive interactions. They found that while the short-range structure of these gels is similar to that of dense amorphous hard-sphere systems, their long-range structure differs remarkably, with regions exhibiting significant density fluctuations.

In most systems of practical importance, long-range van der Waals forces must be balanced by Coulombic, steric, or other repulsive interactions to provide control over phase behavior, structure, and assembly. Nanoparticle engineering is a new paradigm by which these interactions may be regulated.\textsuperscript{1} We recently demonstrated that binary microgels possessing both size and charge asymmetry, in which microspheres are negligibly charged and nanoparticles are highly charged, experience a rich phase behavior, transitioning from a colloidal gel to a stable fluid and subsequently to a colloidal gel with increasing nanoparticle concentration.\textsuperscript{1,30} We further demonstrated that nanoparticle-mediated assembly of dilute binary mixtures on epitaxially patterned substrates could be utilized to create robust three-dimensional (3-D) colloidal crystals.\textsuperscript{31} We attributed the stabilizing transition to nanoparticle “haloing” around the microspheres, which serves to mitigate their van der Waals attraction.\textsuperscript{1,30} The system stability is ultimately reversed at higher nanoparticle concentrations, where flocculation ensues. Utilizing a novel Monte Carlo algorithm,\textsuperscript{2,32} Liu and Luijten were able to study this system by means of computer simulations.\textsuperscript{3} They confirmed that the experimental findings could be quantitatively explained from the formation of a nanoparticle halo, resulting from a weak electrostatic colloid–nanoparticle attraction. At higher nanoparticle concentrations, they also recovered an effective colloidal attraction that increases in strength with nanoparticle concentration and ultimately leads to flocculation. Interestingly, the attraction was found to differ from the regular depletion attraction.\textsuperscript{33,34} Similar findings were also recently reported by Louis and co-workers.\textsuperscript{35}

Here, we investigate the interparticle interactions, phase behavior, and 3-D structure of microsphere–nanoparticle mixtures over a broad composition range by combining this novel Monte Carlo simulation scheme with direct experimental observations. Specifically, both amorphous (gel) and crystalline sediments are created via gravitational settling of microspheres from these mixtures onto flat substrates in aqueous and index-matched solutions. Confocal microscopy is utilized to monitor their assembly in situ as well as to assess their 3-D structure. The use of index-matched solutions, which is required to facilitate 3-D imaging, lessens the van der Waals attraction between silica microspheres in the absence of nanoparticles. Below a lower critical nanoparticle volume fraction ($\phi_{\text{c}}^{\text{NL}} \sim 10^{-4}$), dense colloidal gels form that exhibit enhanced consolidation with increasing nanoparticle concentration. Between $\phi_{\text{c}}^{\text{NL}}$ and an upper critical nanoparticle volume fraction ($\phi_{\text{c}}^{\text{UL}} \sim 2 \times 10^{-3}$), a stable fluid forms from which the microspheres settle into a crystalline array. Finally, above $\phi_{\text{c}}^{\text{UL}}$, dense colloidal gels are also produced whose structure is more open with increasing nanoparticle additions. Interestingly, the observed phase transitions occur at values of $\phi_{\text{c}}^{\text{NL}}$ and $\phi_{\text{c}}^{\text{UL}}$ that are independent of microsphere concentration. These experimental observations are compared to predictions obtained from Monte Carlo simulations, which for this purpose, are extended to treat binary mixtures with larger colloid–nanoparticle size ratio than in ref 3 and index-matched solvent conditions.

Monte Carlo Simulations of Binary Mixtures

Simulation Method. Monte Carlo simulations are carried out to determine the effective microsphere pair potential induced by the nanoparticles in the fluid phase. This effective potential corresponds to a system in which the degrees of freedom of the nanoparticles have been integrated out.\textsuperscript{36} Since the induced interaction is additive to the intrinsic van der Waals attraction between the colloids, it can be computed independently. If it is sufficiently repulsive, it can mitigate the attraction and thus stabilize the suspension. The effective interaction is computed from the colloidal pair potential in the dilute colloid limit. Accordingly, large simulation cells are required, resulting in very large numbers of nanoparticles. Indeed, the calculations presented here involved up to $2 \times 10^{4}$ nanoparticles. In addition, the size asymmetry $a_{\text{mixed}}/a_{\text{nano}} = 200$ between the colloidal microspheres and the nanoparticles makes these calculations virtually inaccessible by conventional simulation methods. The geometric cluster Monte Carlo method,\textsuperscript{2} which was applied to a similar system (albeit for a size asymmetry that was twice smaller) in ref 3, addresses both issues. The nonlocal character of the particle movements overcomes the jamming problems typically experienced by large species immersed in an environment of smaller particles, whereas the efficiency improvement resulting from the rejection-

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The corresponding nanoparticle charge, nanoparticle zeta potential (70 mV in aqueous solution, inverse screening length (2 nm in aqueous solution), at volume fraction 0.02. The nanoparticle volume fraction double-layer interactions are modeled through the Hogg–Healy–Fuerstenau expression.37,38 For two nanoparticles at a surface separation $D$, this equation, under constant-potential conditions, reduces to an exponentially decaying repulsive interaction39

$$V_{\text{nano-nano}} = \epsilon_0 \sigma_{\text{nano}}^2 \Psi_{\text{nano}}^2 \ln[1 + \exp(-\kappa D)] \approx \epsilon_0 \sigma_{\text{nano}}^2 \Psi_{\text{nano}}^2 \exp(-\kappa D)$$ (1)

where the approximation is valid for $\kappa D \gg 1$. $\Psi_{\text{nano}}$ is the nanoparticle zeta potential (70 mV in aqueous solution, corresponding to a nanoparticle charge $Q = 30e$), $\kappa$ the inverse screening length (2 nm in aqueous solution), $\epsilon_0$ the vacuum permittivity, and $\epsilon_r$ the dielectric constant of the solvent (80 for water and 50 for glycerol/water, respectively). Since the silica spheres are negligibly charged, we ignore the electrostatic interaction between them. However, an electrostatic double-layer interaction arises between the microspheres and nanoparticles, which for large size asymmetry is described by37

$$V_{\text{micro-nano}} = \frac{1}{2} \epsilon_0 \sigma_{\text{nano}}^2 \Psi_{\text{nano}}^2 \ln[1 - \exp(-2\kappa D)] \approx -\frac{1}{2} \epsilon_0 \sigma_{\text{nano}}^2 \Psi_{\text{nano}}^2 \exp(-2\kappa D)$$ (2)

where the approximation is again valid for $\kappa D \gg 1$. Thus, an attraction is induced that is responsible for the formation of a nanoparticle halo. It must be noted that we have employed the “bare” average nanoparticle diameter $d_{\text{nano}} = 6$ nm in eq 2, whereas in eq 1 we have accounted for the hydration repulsion between nanoparticles by increasing the diameter by 1.5 nm.

**Experimental Section**

**Materials System.** Uniform silica microspheres (Geltech, Alachua, FL) serve as the large colloidal species. These microspheres have an average radius $a_{\text{micro}}$ of 0.590 ± 0.01 µm, as determined from quantitative image analysis carried out on scanning electron microscopy (SEM) photomicrographs (15 000 × magnification), and a density of 2.25 g/cm$^3$, as determined by X-ray scattering measurements with a Debye length ($\lambda_\text{D}$) of 1.8 nm under the experimental conditions of interest ($\Phi = 1.5$).

Hydrous zirconia nanoparticles (Zr 10/20, Nyacol Products, Ashland, MA) serve as the small colloidal species. These nanoparticles have an average radius $a_{\text{nano}}$ of 3 nm (see Figure 1), as determined by X-ray scattering measurements with a reported radius range of 0.5–11 nm.40,41 Their reported density is 3.65 g/cm$^3$,42 and they are supplied in an acidic solution (pH = 0.5) at a volumetric solids loading of 7.4%. At pH = 1.5, they possess a zeta potential of 63 ± 12 mV, as determined by microelectrophoresis (Malvern Zetasizer 3000). This value is in reasonable agreement with the zeta potential of approximately 70 mV estimated from the reported effective charge determined from titration studies40 using the approach outlined by Gisler et al.43

**Suspension Preparation.** Aqueous binary suspensions are prepared by adding an appropriate volume fraction of nanoparticles to deionized water predosed with nitric acid (reagent grade, Fisher Scientific) to pH = 1.5. An appropriate volume fraction of a stock solution (pH = 1.5) of SiO$_2$ microspheres ($\Phi_{\text{micro}} = 2.5 \times 10^{-2}$) is then added followed by ultrasonication (model 550 sonic dismembrator, Fisher Scientific, Pittsburgh, PA). The suspensions are then stirred for 0.5 h followed by another ultrasonication step.

Binary suspensions are also prepared in an index-matched solution by first mixing an appropriate volume of deionized water (pH = 1.5) with a stock solution of glycerol that contained fluorescent dye (Rhodamine B, Sigma-Aldrich, St. Louis, MO). The ratio of water-to-glycerol is 1:2.8, as given by the Lorenz–Lorentz index of refraction mixing rule. An appropriate volume fraction of nanoparticles is then added, followed by stirring for 10 min. Next, an appropriate volume fraction of colloidal microspheres is added, followed by ultrasonication. The suspensions are then stirred for 5 min followed by ultrasonication. Each sonication step consists of 2.5 min pulsed at a 1 s on/off sequence at 20 kHz.

**Bulk Sedimentation.** Colloidal silica microspheres ($\Phi_{\text{micro}} = 1.0 \times 10^{-3}$) suspended in binary mixtures undergo gravity-driven sedimentation in glass cylinders (8 cm (H), 0.4 cm (D)) that are capped to minimize solvent evaporation. Initially, the samples are opaque (even those suspended in the index-matched solution)

due to the scattering of visible light from the colloidal microspheres. As the samples settle, the suspension separates into a clear, microsphere-free supernatant, a cloudy solution, and an opaque sediment. This process is imaged using time-lapse photography with a video camcorder (Digital Handycam Camcorder, Sony, Japan) controlled by SupervisionCam software (Peter Kirst, Herrsching, Germany).

The initial microsphere settling velocity ($V_{\text{micro}}$) is determined by tracking the interface separating the clear and cloudy solutions. Normalized values ($V/\text{V}_{\text{micro}}$) are reported to account for the large differences in solution viscosity between the aqueous and index-matched media and thereby facilitate direct comparison between these two systems. One can calculate $V_{\text{micro}}$ using the Richard–Zaki equation:

$$V_{\text{micro}} = V_0 (1 - \phi_{\text{micro}})^{\frac{3}{2}}$$  \hspace{1cm} (3)

where $V_0$ is the settling velocity for an isolated microsphere determined by Stokes’ law

$$V_0 = \frac{2}{9} \frac{\rho_m - \rho}{\eta} g$$  \hspace{1cm} (4)

$\rho_m$ is the microsphere density, $\rho$ is the density of the liquid phase, $g$ is the apparent solution viscosity ($\eta = 1$ mPa$s$ for water, 40 mPa$s$ for glycerol/water), $g$ is the gravitational constant, and $n$ is a power law exponent ($n = 4.65$). $V_{\text{micro}}$ is found to be $5.8 \times 10^{-4} \text{ mm/s}$ for microsphere sedimentation from concentrated binary mixtures ($\phi_{\text{micro}} = 0.1$) in aqueous and index-matched solutions, respectively. Microsphere settling velocities ($V_{\text{micro}}$) are also determined from experimental observations for mixtures residing in the stable binary fluid phase, where settling of individual microspheres from suspension is observed. The minimum values of $V_{\text{micro}}$ are found to be $\sim 3 \times 10^{-4} \text{ and } 1 \times 10^{-3} \text{ mm/s}$ for aqueous and index-matched solutions, respectively. These experimental values, which are in good agreement with those predicted from eq 3, are utilized to determine the reported normalized settling velocity ($V/\text{V}_{\text{micro}}$). Finally, the sediment volume fraction ($\phi_{\text{volumetric}}$) is determined by measuring the final sediment height at long times ($\geq$ several weeks).

**Colloidal Assembly.** Colloidal silica microspheres suspended in binary mixtures are assembled under gravity-driven sedimentation in custom sample cells designed for confocal imaging. An appropriate volume of a given binary suspension is used as required to yield a final sediment height of nominally 150 μm for suspensions with $\phi_{\text{micro}} = 2.5 \times 10^{-3}$ and $2.5 \times 10^{-2}$ and 1 mm for suspensions with $\phi_{\text{micro}} = 2.5 \times 10^{-4}$ (the sample volume required to create 150 μm sediments is too low at this $\phi_{\text{micro}}$). Each cell is fabricated by attaching a glass tube ($40 \text{ mm (H)} \times 8 \text{ mm i.d.})$ to a glass cover slip using poly(dimethylsiloxane) (PDMS) (Sylgard 184, Dow Corning, Midland, MI). Both the tubes and cover slips are cleaned prior to bonding by immersion in a 3:1 sulfuric acid (reagent grade, Fisher Scientific)/hydrogen peroxide (30% in water, Fisher Scientific) solution for 1 h. They are then rinsed repeatedly with deionized water, dried under a nitrogen stream, and bonded to one another using PDMS.

**Image Analysis**}

The confocal images are analyzed using IDL routines to filter the images and to locate particle positions in three dimensions. The images are first filtered using a spatial band-pass filter to remove noise at high and low spatial frequencies. Each particle center is then located based on the regional intensity maximum. When the pixel intensity is averaged across the entire particle, its location can be identified with subpixel accuracy ($\sim 50 \text{ nm}$). 3-D reconstructions ($23 \times 21 \times 12.5 \text{ μm}^3$) are obtained for representative sediment structures. The sediment volume fraction is determined by dividing the total microsphere volume ($V_{\text{tot}} \times 4/3\pi a^3$, where $V_{\text{tot}}$ is the total number of microspheres and $a$ is the sphere radius) by the imaged volume and averaging over that obtained from five different scanned regions per sample. The pair correlation function $g(r)$ is determined by finding the number of microsphere centers inside a volumetric shell created a distance r away from a given microsphere center, relative to a uniform distribution of the same density. Each microsphere is checked against all other microspheres in a given scan volume over a total radius of 6 μm using $dr = 0.01 \text{ μm}$ as the shell thickness. We next identify nearest neighbors using the Delaunay triangulation method and calculate the Voronoi polyhedron volume associated with each particle. By examining the orientation (bond angle and distance) between a given microsphere and its Delaunay nearest neighbors, one can also determine whether it resides in an amorphous or crystalline structure. Adjacent particles with similar orientations of their neighbors are termed “ordered neighbors”, and a particle with eight or more ordered neighbors is considered to be a crystalline particle.

**Results and Discussion**

**Microsphere–Nanoparticle Interactions.** We first determine the effective interactions induced between microspheres in the presence of nanoparticle species in both aqueous and index-matched solutions in order to predict their respective phase behavior. Figures 2 and 3 show the effective microsphere potential $V_{\text{eff}}kT$ as determined from the colloidal pair correlation function obtained in the simulations. Colloidal many-body effects have been eliminated by performing these simulations in the dilute colloid limit. At low nanoparticle concentrations in water (Figure 2a), a repulsive barrier appears that increases with nanoparticle volume fraction. This is similar to what has been reported in ref 3 for a microsphere-nanoparticle size ratio of 100, although in the present case lower nanoparticle volume fractions are needed to achieve a comparable effective repulsion, due to the larger size ratio. Upon further increase of the nanoparticle concentration (Figure 2b), an attractive minimum appears in the potential, preceding the repulsive barrier. It was found in ref 3 that the depth of this minimum grows quadratically with $\phi_{\text{micro}}$, indicating a qualitative difference with the linear concentration dependence exhibited by regular depletion and repulsive barrier.

*Examination of microsphere/nanoparticle configurations indicates that, upon approach of two microspheres, the halos remain largely unperturbed, and at a separation
corresponding to the attractive minimum of the effective potential only limited nanoparticle depletion is observed in the region between the opposing halos.

In the index-matched solvent, $V_{\text{eff}}$ varies in a similar way with nanoparticle volume fraction. For $\phi_{\text{nano}}$ as low as $10^{-4}$, an effective barrier of approximately $5.5\ k_B T$ is observed in the simulations (Figure 3a). Compared to the colloids suspended in water, lower nanoparticle concentrations are required to achieve a given induced repulsion. This is consistent with an enhancement in nanoparticle haloing around the microspheres, which in turn stems from the higher nanoparticle zeta potential employed in eq 2. The zeta potential is determined from the corresponding potential in aqueous solution under the assumption that the nanoparticle surface charge is independent of solvent composition. Because of our inability to assess the zeta potential in glycerol-water solutions, a more accurate value could not be obtained under these conditions. At higher $\phi_{\text{nano}}$ the attractive minimum is observed again (Figure 3b). While the concentration at which a certain potential minimum is achieved is also lower for the index-matched solution than under aqueous conditions, the difference between the two cases amounts to more than a simple shift of both phase boundaries. If we tentatively associate a barrier of $5\ k_B T$ with the lowest nanoparticle volume fraction $\phi_{\text{nano}}$ required to mitigate the intrinsic van der Waals attraction between the colloids, and a minimum of $-2\ k_B T$ with the nanoparticle volume fraction $\phi_{\text{C}}$ beyond which gelation occurs, we obtain a window of stability $\phi_{\text{C}}^{\text{U,C}}/\phi_{\text{C}}^{\text{L,C}} \approx 17$ in aqueous solution, whereas we find $\phi_{\text{C}}^{\text{U,C}}/\phi_{\text{C}}^{\text{L,C}} \approx 60$ for the index-matched solution. We note that the actual window for the latter is possibly even larger, as we have not taken into account the lowering of $\phi_{\text{C}}^{\text{L,C}}$ due to the weakening of the van der Waals attractions.

**Phase Behavior of Microsphere–Nanoparticle Mixtures.** The phase behavior of microsphere–nanoparticle mixtures of varying composition in aqueous and index-matched solutions is determined experimentally in order to make comparisons to the behavior predicted by simulations. The phase diagram, shown in Figure 4, is obtained by monitoring microsphere sedimentation in bulk samples and directly by using confocal microscopy. In the absence of nanoparticles, negligibly charged silica microspheres flocculate due to van der Waals interactions yielding clusters that settle rapidly to form a dense,
amorphous sediment. The van der Waals potential $V_{\text{vdW}}$ for silica microsphere–microsphere interactions is calculated to be approximately $-60 k_B T$ in aqueous and $-16 k_B T$ in index-matched solutions, respectively, at a minimum separation distance of approximately 1 nm (cf. the approach outlined in ref 28). Below a lower critical nanoparticle volume fraction $\phi_{\text{L,C}}$, the system remains unstable, yielding a dense, amorphous sediment. This observation suggests that the microsphere interactions are indeed influenced by van der Waals forces, even in index-matched systems where they are substantially weakened. However, these forces decay very rapidly with increasing microsphere separation, and a nanoparticle-induced repulsive barrier, as demonstrated in Figures 2a and 3a, is sufficient to prevent aggregation. Indeed, at intermediate nanoparticle volume fractions ($\phi_{\text{L,C}} < \phi_{\text{L,C}}$), the system resides in the stable fluid region from which individual colloidal microspheres settle under gravity to produce a crystalline sediment. The value of $\phi_{\text{L,C}}$ appears to be relatively insensitive to microsphere volume fraction or the solution in which they interact. Finally, above the upper critical volume fraction ($\phi_{\text{L,C}} \geq \phi_{\text{L,C}}$), the system becomes unstable, giving rise to a new amorphous structure upon sedimentation. This is consistent with the attractive minima observed in the simulations (Figures 2b and 3b) at higher nanoparticle concentrations. Upon quantitative comparison with the simulation results, we note a number of points. In view of the relatively coarse-grained model employed and the degree of arbitrariness involved (both in the simulations and experimentally) in locating the phase boundaries, the lower phase boundary $\phi_{\text{L,C}}$ for both solution compositions may be considered quite close to the concentrations in Figures 2a and 3a, although the simulations exhibit a solvent dependence for $\phi_{\text{L,C}}$ that is not observed in the experiments. The simulational results for the estimated upper phase boundary are rather similar for both cases and lie somewhat above the experimental estimate for $\phi_{\text{L,C}}$. As a result, the window of stability $\phi_{\text{L,C}}/\phi_{\text{L,C}}$ for the aqueous case has almost the same value in the simulations ($\approx 17$) as in the experiments ($\approx 20$), and the difference between simulation and experiment essentially amounts to a shift in nanoparticle concentration. In contrast, for the index-matched case there is a greater discrepancy between the $\phi_{\text{L,C}}/\phi_{\text{L,C}}$ value predicted by simulations ($\approx 60$) and that observed experimentally ($\approx 20$). This suggests that the extent of nanoparticle haloing, which the simulations find to be stronger in the index-matched solution, is actually rather similar in magnitude for both experimental systems. As mentioned above, this reflects our assumption regarding the nanoparticle charge in the index-matched case.

Interestingly, a broadening of the stable fluid region at lower microsphere volume fractions, as originally suggested by Töhrer et al., is not observed. There is, however, a modest narrowing of this region at the highest microsphere volume fraction studied, in good accord with prior observations.

**Direct Imaging of Colloidal Assembly from Microsphere–Nanoparticle Mixtures.** The structural evolution observed during gravity-driven assembly of colloidal microspheres from index-matched binary mixtures is monitored in situ using confocal microscopy over a broad concentration range. We report only the time evolution sequence for binary mixtures comprised of the lowest microsphere volume fraction studied ($\phi_{\text{micro}} = 2.5 \times 10^{-3}$), because of the fast dynamics at higher concentrations. The $x$–$y$ scans, shown in Figure 5, reveal the dramatic differences in structure observed for binary mixtures assembled from four different regions within the phase diagram, where $\phi_{\text{nano}} = 0$ (microspheres only), $5 \times 10^{-5}$ (lower unstable region), $5 \times 10^{-4}$ (stable fluid region), and $5 \times 10^{-3}$ (upper unstable region). In these images, the colloidal microspheres appear as dark circular features against a surrounding lighter background that corresponds to the index-matched, fluorescent dye solution. In the absence of nanoparticle additions, the microspheres aggregate into stringlike clusters that are elongated in shape. This type of cluster formation leads to an initially open sediment structure, as shown in the $x$–$z$ scan depicted in Figure 6. However, this amorphous sediment ultimately densifies over time under its own weight, as microspheres continue to settle from solution. In sharp contrast, there is little evidence of cluster formation in binary mixtures that reside in the lower unstable region ($\phi_{\text{nano}} = 10^{-5}$). Despite being unable to visualize microsphere clusters in this sample, we observe that they exhibit a faster sedimentation rate than individual microspheres settling from the stable fluid phase ($\phi_{\text{nano}} = 5 \times 10^{-3}$). This is perhaps most obvious in the $x$–$y$ scans acquired at times of 120 and 180 min, in which there is a greater accumulation of colloidal microspheres at the substrate–sediment interface for the unstable mixture. A finite number of small clusters must form in mixtures that reside in the lower unstable region—otherwise there would be no difference in the rate of microsphere accumulation relative to the settling rate observed for individual microspheres in the stable region. This is confirmed by bulk sedimentation results (see Figure 5b) that reveal significantly higher settling velocities for microspheres in mixtures that reside in the lower (and upper) unstable regions than for those in the stable fluid phase. Finally, in the upper unstable region ($\phi_{\text{nano}} = 5 \times 10^{-3}$), there is again visual evidence in both the $x$–$y$ and $x$–$z$ scans that the microspheres form clusters, albeit far more compact in nature than those formed in the absence of nanoparticle additions. In this region, the microsphere clusters settle from suspension to form an amorphous sediment that is initially more open than that formed from either of the other mixtures, but significantly less open than that produced in the absence of added nanoparticles. Below, we comment on the origin of this difference. At the longest times probed (> 3 weeks), only sediments assembled from the stable fluid phase are observed to crystallize. In all other cases, the final sediment consists of a dense, amorphous structure.

The microsphere–nanoparticle mixtures thus exhibit four distinct types of behavior during their assembly as the nanoparticle volume fraction increased: (1) aggregation leading to elongated clusters in the absence of nanoparticles, (2) aggregation leading to a large population of individual microspheres with some compact clusters at low nanoparticle concentrations, (3) a stable fluid phase at intermediate nanoparticle concentrations, and (4) aggregation leading to compact clusters at higher nanoparticle concentrations. These variations in cluster morphology stem from differences in the effective microsphere interactions as a function of $\phi_{\text{nano}}$, following the sequence depicted in the simulational results (Figures 2 and 3). As microsphere interactions become more attractive, their sticking probability increases resulting in clusters that are more branched in nature. For the binary mixtures studied here, the strongest attractive interactions between microspheres occur in the absence of nanoparticle additions, where van der Waals forces dominate. Initial sediments assembled after a few hours of settling from dilute microsphere suspensions are most open, since there is ample opportunity for cluster growth prior to further consolidation as settling proceeds. In contrast, as the
microsphere concentration increases, cluster growth appears to be suppressed by enhanced sedimentation leading to initial sediment structures (~few hours) that are far more compact in nature (data not shown). As discussed below, the final structure of colloidal assemblies formed from these binary mixtures under gravity-driven sedimentation is influenced by their microsphere and nanoparticle concentrations. The dense structures observed at high nanoparticle concentrations are compatible with the interaction profiles in Figures 2b and 3b, where it was observed that, in addition to the shallow attractive minimum (which is presumably responsible for the

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Figure 5. Confocal images of x–y scans acquired at the sediment–substrate interface for structures assembled from binary mixtures comprised of φ_{micro} = 2.5 \times 10^{-3} and varying nanoparticle volume fraction (φ_{nano}) at different times during sedimentation.

Figure 6. Confocal images of x–z scans acquired through structures assembled from binary mixtures composed of φ_{micro} = 2.5 \times 10^{-3} and varying nanoparticle volume fraction (φ_{nano}) at different times during sedimentation.
aggregation behavior), a strong repulsive barrier is still present, suggesting that the (considerably stronger) van der Waals interactions continue to be (partially) mitigated in the upper part of the phase diagram.

**Structure of Colloidal Phases Assembled from Microsphere–Nanoparticle Mixtures.** We systematically analyze the final sediment structures assembled from binary mixtures to quantify the effects of microsphere and nanoparticle concentration within each region of the phase diagram reported in Figure 4. 3-D reconstructions for representative sections of sediments assembled from these mixtures are shown in Figure 7. They are color-coded to distinguish colloidal particles in disordered (blue) and crystalline regions (red). Crystalline structures are observed only for samples that settle from binary mixtures that reside in the stable fluid region. The 3-D reconstructions reveal the polycrystalline nature of sediments formed from within this region, with blue particles highlighting domain boundaries that exist at the interface between single crystal domains. Outside this region, the sediments are comprised of dense, amorphous structures (or gels). Because it is difficult to discern the subtle effects of varying composition on the 3-D structure of either the crystalline or amorphous sediments solely from these images, we further analyze the imaged volumes to determine their sediment volume fraction, radial distribution function, Voronoi polyhedra volume, and local bonding structure.

The sediment volume fraction, $\phi_{\text{sediment}}$, is plotted in Figure 8a for samples assembled from binary mixtures of varying composition. For comparison, data acquired directly from confocal images of samples formed from binary mixtures suspended in the index-matched medium is shown along with data determined by bulk sedimentation of binary mixtures in both aqueous and index-matched solutions. Not surprisingly, the most open sediment structure ($\sim 30\%$ vol) forms for pure microspheres settling in an aqueous solution. When the microspheres settle in an index-matched solution, open sediments of density ($\sim 35\%$ vol) similar to those reported for the aqueous system are observed only for the most dilute sample studied. Enhanced consolidation occurs with increasing microsphere volume fraction leading to sediment densities of $35\%$, $43\%$, and $51\%$ vol for sediments produced from initial microsphere suspensions of $2.5 \times 10^{-3}$, $2.5 \times 10^{-2}$, and $2.5 \times 10^{-1}$, respectively. With increasing nanoparticle volume fraction, the sediment density increases until a plateau value of roughly $60 \pm 3\%$ vol is reached for mixtures of all compositions settling within the stable fluid phase. In contrast, the maximum densities ($\sim 53–57\%$ vol) observed for sediments formed during bulk sedimentation (initial $\phi_{\text{micro}} = 10^{-1}$) from both aqueous and index-matched systems are considerably lower in this region. In the upper unstable region, the sediments become increasingly more open with further increases in nanoparticle volume fraction. At a given nanoparticle concentration, denser sediments are observed for mixtures with higher initial microsphere volume fraction in both the lower and upper unstable regions of the phase diagram.
several index-matched systems, suggesting that the strength of the interaction is most strongly on both nanoparticle volume fraction and solution composition. As stated previously, these data points correspond to bulk sedimentation results. (b) Normalized settling velocity ($V/V_{\text{micro}}$) as a function of nanoparticle volume fraction ($\phi_{\text{nano}}$) during initial sedimentation of binary mixtures ($\phi_{\text{micro}} = 1 \times 10^{-1}$) suspended in aqueous and index-matched solutions ($\phi_{\text{micro}} = 1 \times 10^{-1}$ suspended in an aqueous solution, where triangular data points correspond to bulk sedimentation results. Interestingly, the sediment densities in the absence of nanoparticles and at the highest nanoparticle volume fraction ($\sim 10^{-2}$) investigated are quite comparable for the index-matched systems, suggesting that the strength of the attraction in both limits is similar (of the order of several $k_B T$), despite the fundamentally different origin of the interaction. The initial microsphere settling velocities depend strongly on both nanoparticle volume fraction and solution composition, as shown in Figure 8b. As stated previously, these data are presented in a normalized fashion to facilitate direct comparison between aqueous and index-matched binary mixtures. In the absence of nanoparticle additions, the normalized settling velocity is approximately 100- and 6-fold higher than for individual microsphere sedimentation from the stable fluid phase in aqueous and index-matched systems, respectively. These values reflect the formation of particle clusters in suspension driven by van der Waals attractions between microspheres. Because the microsphere volume fraction is identical in both systems, the higher settling rates observed for aqueous suspensions are attributed to enhanced cluster formation compared to the index-matched suspensions. This is not unexpected given the suppression of the van der Waals attractions in the latter system. In the lower unstable region, the normalized settling velocity decreases as a function of increasing nanoparticle volume fraction until a value of unity is approached at the critical lower nanoparticle volume fraction $\phi_{L,C}$. This decrease in velocity is directly linked to diminished cluster formation, resulting from the effective potential barrier (Figures 2a and 3a) that in turn arises from nanoparticle halo formation. Within the stable fluid phase, both systems exhibit normalized settling velocities of roughly unity, corresponding to the sedimentation of individual microspheres in these samples. Finally, at higher nanoparticle volume fractions ($\phi_{\text{nano}} > \phi_{L,C}$), the normalized settling velocity increases with increasing nanoparticle volume fraction, as the effective microsphere attraction intensifies, leading to enhanced cluster formation. The rapid nature of this increase can be understood from the simulation results (cf. Figures 2b and 3b and ref 3), which indicate that the microsphere attraction increases superlinearly with nanoparticle concentration.

The microsphere pair correlation functions $g(r)$ of sediments assembled from binary mixtures of varying composition are reported in Figure 9. The peaks in $g(r)$ indicate ordering of particles into coordination shells. The data for crystalline sediments assembled from binary mixtures that reside in the stable fluid phase possess features characteristic of a well-ordered structure; i.e., the particles are organized in multiple coordination shells around a given central particle. As the microsphere volume fraction increases, the degree of order observed for these crystalline sediments lessens slightly, as indicated by a modest broadening of the first and higher order peaks. In contrast, the $g(r)$ shown for sediments assembled from binary mixtures that reside in the lower or upper unstable regions of the phase diagram possess features characteristic of a dense liquid structure. There are subtle differences between these data. For example, the first peak in $g(r)$ is broadest in the amorphous sediment structures formed in the absence of nanoparticles or at the highest nanoparticle volume fraction probed for systems with the lowest microsphere volume fraction. However, more striking is the similarity in $g(r)$ for sediments assembled within both the lower and upper unstable regions near their respective phase boundaries. Important differences do emerge, however, when analyzing the heterogeneity of these weak gels via Voronoi tessellation.

van Blaaderen and Wiltzius20 and Varadan and Solomon22 were first to apply Voronoi tessellation to spatially characterize the respective structures of colloidal glasses and dense colloidal gels. The peak value in the Voronoi polyhedral (VP) volume distribution reflects the average VP volume available to colloidal microspheres within these dense sediment structures, whereas the standard deviation and skewness of the distribution are indicative of their structural inhomogeneity, as shown in Figure 10. The VP volume distributions for amorphous sediments formed from binary mixtures in the lower or upper unstable regions exhibit a distribution that is broader than that observed for crystalline sediments formed from binary mixtures that reside in the stable fluid phase. Correspondingly, their values are also higher, indicative of greater structural heterogeneity. It is illustrative to first examine these data for sediments formed from pure microsphere suspensions of varying $\phi_{\text{micro}}$. The VP volume distribution is broadest for sediments produced from the most dilute microsphere suspension and narrows significantly with increasing $\phi_{\text{micro}}$. Correspondingly, the $\alpha$ values reported for the pure microsphere sediments systematically decrease from 0.7 to 0.29 $\mu m^3$ with increasing $\phi_{\text{micro}}$, which reflects the enhanced structural homogeneity of gels at higher density (or $\phi_{\text{sediment}}$). As the nanoparticle volume fraction is increased in the lower unstable region, the sediment structures density to yield more homoge-
neous structures, as reflected by a shift in their VP peak to lower values and a simultaneous reduction in their $\sigma$ values. The VP volume distributions for crystalline sediments assembled from binary mixtures that reside in the stable fluid region possess features characteristic of a well-ordered structure, i.e., a very narrow distribution with a peak at $1.2 \text{ m}^3$ and values of $0.1 \text{ m}^3$. This is close to the minimum value of $1.16 \text{ m}^3$ expected for microsphere crystals that adopt an face-centered cubic (or hexagonal close packed) packing geometry. As the microsphere volume fraction increases, these crystalline sediments become less ordered, as indicated by a decrease in VP peak intensity. Finally, in the upper unstable region, there is a systematic shift to a broader VP volume distribution and higher values with increasing nanoparticle volume fraction. In this region, the observed changes appear to be less sensitive to microsphere volume fraction.

The number of nearest-neighbor (#NN) bonds per particle is depicted in Figure 11 for sediments assembled from binary mixtures of varying composition. This quantity provides insight into the degree of structural rearrangement at the local scale. The local topology of colloidal gels should be dominated by chainlike aggregates...
shifting toward lower peak values with the opposite trend emerges with the \( \phi_{\text{nano}} \) distribution conditions. Finally, in the upper unstable phase region, the positional positions during sedimentation under these microspheres become increasingly trapped in nonequilibrium positions with increasing \( \phi_{\text{nano}} \) in this region, but there is a modest broadening of the \( \text{#NN} \) distribution with increasing microsphere volume fraction in this region, which spans over a decade in \( \phi_{\text{nano}} \) as specified. Arrows denote direction of increasing nanoparticle volume fraction.

(\( \text{#NN} = 2 \)) if the particle–particle bonds are rigid and irreversible. In sharp contrast, if particles within a given cluster can rearrange by, for example, rotation, then significant compaction and reorganization can occur. In the absence of nanoparticles, a chainlike morphology is observed during the initial stage of sedimentation of dilute microsphere suspensions (\( \phi_{\text{micro}} = 2.5 \times 10^{-3} \)); see Figure 5. However, at longer sedimentation times (>3 weeks), these sediment structures experience substantial consolidation yielding a final \( \text{#NN} \) bonds of \( \sim 4 \) and a \( \phi_{\text{sediment}} \) of \( \sim 0.35 \). In the lower unstable region, the peak in the \( \text{#NN} \) distribution increases with increasing nanoparticle volume fraction within these dense amorphous sediments. This observation suggests an overall weakening of microsphere–microsphere bonds with increasing \( \phi_{\text{nano}} \) in this region of the phase diagram. In the stable fluid region, the peak in the \( \text{#NN} \) distribution occurs at a value of 12, which is expected given their crystalline nature. There is no discernible dependence on nanoparticle volume fraction in this region, but there is a modest broadening of the \( \text{#NN} \) distribution with increasing microsphere volume fraction. This trend likely reflects the fact that the microspheres become increasingly trapped in nonequilibrium positions during sedimentation under these conditions. Finally, in the upper unstable phase region, the opposite trend emerges with the \( \text{#NN} \) distribution shifting toward lower peak values with \( \phi_{\text{nano}} \). Suppressed bond rotation is expected due to intensified microsphere–microsphere attractions deeper within the upper phase region. Our observations are in good agreement with those reported by Weitz and co-workers\(^2\) for depletion gels formed from colloidal–polymer mixtures.

## Conclusions

We have investigated the interparticle interactions, phase behavior, and structure of binary microsphere–nanoparticle mixtures that possess both high charge and size asymmetry. Through compositional tailoring, we have created both colloidal gels and crystals by gravity-driven sedimentation of colloidal microspheres in aqueous and index-matched solutions on flat substrates. The use of an index-matched, fluorescent dye solution is required to facilitate 3-D imaging and structural analysis of the assembled colloidal phases. However, a direct consequence of index matching is that the strength of the van der Waals attraction between microspheres is lessened considerably relative to their interaction in aqueous solution. Thus, the colloidal gels that form in the absence of nanoparticle additions (or at low concentrations) are relatively weak yielding amorphous structures that densify considerably under their own weight. At intermediate nanoparticle concentrations, the colloidal microspheres are fully stabilized and form a polycrystalline array upon sedimentation. Within this region (which spans over a decade in nanoparticle volume fraction), there is surprisingly little effect of nanoparticle concentration on the observed 3-D crystal structure or sediment density. In sharp contrast, the degree of crystallinity declines with increasing microsphere volume fraction, as evidenced by a substantial broadening of the distributions of both the Voronoi polyhedra volumes and number of nearest neighbors. These observations have important implications for our parallel efforts, which exploit the nanoparticle-mediated assembly of colloidal microspheres on epitaxially patterned substrates.\(^3\) Namely, the structural evolution of colloidal crystals assembled from stable (fluid) mixtures depends far more critically on microsphere concentration than on nanoparticle concentration. In the lower and upper unstable regions, the driving force for microsphere flocculation stems from van der Waals interactions and effective, nanoparticle-induced interactions, respectively, as we have demonstrated by direct simulation of microspheres in the presence of various nanoparticle concentrations. The nanoparticles initially only induce a repulsive barrier that increases in strength with increasing nanoparticle concentration, but at higher concentrations an effective attraction appears as well. Therefore, the structural evolution of colloidal gels assembled from binary mixtures in these two regions exhibits the opposite dependence on nanoparticle concentration, such that denser gels form with increasing nanoparticle volume fraction in the lower unstable region whereas more open

![Figure 11](image-url)

**Figure 11.** Probability distribution of the number of nearest neighbors for structures assembled from binary mixtures comprised of (a) \( \phi_{\text{micro}} = 2.5 \times 10^{-3} \), (b) \( \phi_{\text{micro}} = 2.5 \times 10^{-5} \), and (c) \( \phi_{\text{micro}} = 2.5 \times 10^{-7} \) with varying nanoparticle volume fraction (\( \phi_{\text{nano}} \), as specified).
gels form with increasing nanoparticle volume fraction in
the upper unstable region. The ability to tune the phase
behavior and structure of colloidal phases through nano-
particle engineering broadens the palette of systems that
can be used as precursors for assembling ceramic, photonic
band-gap, and pharmaceutical materials, as well as novel
inks and coatings.

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