Velocity fluctuations of initially stratified sedimenting spheres

Shang-You Tee  
Department of Physics and Division of Engineering and Applied Sciences, Harvard University, Cambridge, Massachusetts 02138, USA  

P. J. Mucha  
Department of Mathematics and Institute for Advanced Materials, University of North Carolina, Chapel Hill, North Carolina 27599, USA  

M. P. Brenner  
Division of Engineering and Applied Sciences, Harvard University, Cambridge, Massachusetts 02138, USA  

D. A. Weitz  
Department of Physics and Division of Engineering and Applied Sciences, Harvard University, Cambridge, Massachusetts 02138, USA

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The study of velocity fluctuations in the sedimentation of spheres is complicated by the time evolution of the underlying particle distribution, both at the microscale and in the bulk. We perform a series of experiments and simulations to isolate the effect of an initial, stable stratification in the particle concentration. The directly observed dependence of velocity fluctuations on stratification agrees with a previously obtained scaling theory. © 2007 American Institute of Physics. [DOI: 10.1063/1.2806597]

Sedimentation is a fundamental fluids process of broad importance that possesses interesting physics and perplexing behaviors. Even the seemingly simplest situation, the slow sedimentation of a dilute collection of monodisperse spheres in a Newtonian fluid, continues to generate questions of intriguing difficulty. While the mean sedimentation velocity is well understood, the velocity fluctuations about this mean velocity are more complicated. Caflisch and Luke found that stratification effects have been shown to play a role in wavelet-scale selection in the instability of sedimenting spheroids. Amid such uncertainties, a more controlled investigation is warranted to quantify the dependence of velocity fluctuations on stratification in situations where stratification is the dominant controlling mechanism.

In this article, we demonstrate through experiments and simulations that the velocity fluctuations decrease with increasing stratification, in quantitative agreement with an existing scaling model. To avoid complications from evolving density profiles, we choose a series of experiments and simulations initialized with dilute, linear particle concentration profiles, at physical parameters minimizing mixing so the initial profile is well-maintained in time, up to the near-constant average downward transport. Because of the absence of appreciable time evolution, we can effectively isolate and quantify the dependence of velocity fluctuations on stratification.

Velocity fluctuations are driven by particle density fluctuations, as demonstrated by a physical argument due to Hinch, assuming that a vertically homogeneous dilute random sample obeys Poisson statistics, a region of size \( \ell \) with particles of radius \( a \) at volume fraction \( \phi \) has a typical concentration fluctuation \( \Delta \phi \sim \sqrt{3\phi a^3}/\ell^3 \). The velocity of such a region is determined by balancing its buoyant weight, \( \Delta \phi \ell^3 \Delta \rho g \sim (\Delta \phi) \ell^3 (\eta V_0 / a^2) \), with its Stokes drag, \( \sim \eta \ell \Delta V \), giving \( \Delta V = CV_0 \sqrt{\phi \ell / a} \), with \( V_0 \) the Stokes velocity of an isolated particle. The dominant \( \Delta V \) in homogeneous sedimentation is then set by the smallest container dimension, \( d \), because the above arguments break down for \( \ell > d \), giving

\[
\Delta V_{\text{Poisson}} = CV_0 \sqrt{\phi d / a},
\]

in agreement with Caflisch and Luke. These fluctuations lead to long-time diffusive mean squared displacements with effective diffusivity, \( D \sim \ell \Delta V \). Density fluctuations on a given length scale are then produced and destroyed due to randomness on the same time scale, \( \tau_\phi \sim \ell^2 / D = \ell / \Delta V \), advecting a distance \( \sim \ell \) in their lifetime. The competition between fluctuation creation and destruction drives almost any

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vertically homogeneous system to Poisson statistics at long length scales, consistent with the assumed statistically steady density fluctuations.

Even a very small stable particle density stratification modifies the above argument. The buoyancy mismatch of density fluctuations is lost if the $\Delta \phi$ density fluctuations are smaller than the stratification concentration change over a swirl lifetime translating distance $\sim \ell$, meaning such fluctuations can no longer advect as far as in the vertically homogeneous case. For a locally linear decrease in $\phi$ in the direction opposite to the average motion, $\beta = \phi_0 (1 - \beta z)$, with $\beta > 0$, the stratification suppresses fluctuations for scales bigger than those where the two changes in concentration balance, $\beta \phi \ell \sim \Delta \phi$, that is, $\ell \sim a \phi^{-1/5} (\beta a)^{-2/5}$, with velocities $\Delta V \sim V_0 a^{2/5} (\beta a)^{-1/5}$. These arguments for stratification control of fluctuations apply only when $\ell < d$; otherwise, in the absence of other controlling mechanisms, the fluctuations are set by the container size $d$. The same scalings are obtained from analysis of a model stochastic continuum equation.\(^\text{12}\)

Setting $\ell \sim d$, we define the critical stratification,

$$
\beta_{\text{crit}} = \frac{d^{3/2} \phi^{5/2}}{12V \phi_0^{2/5} \Delta \phi^{1/5}} = (d \sqrt{N_d})^{-1},
$$

where $N_d$ is the number of particles in a volume of radius $d$, with $\Delta V^2 / \Delta V_{\text{Poisson}}^2 \sim (\beta / \beta_{\text{crit}})^{-2/5}$ for $\beta$ above the critical stratification. This critical stratification decreases rapidly as the cell thickness $d$ increases, with the actual crossover at an unknown but presumably $O(1)$ scalar multiple of the $\beta_{\text{crit}}$ defined in Eq. (2).\(^\text{12}\) Earlier studies\(^\text{1,12}\) indicated this crossover to be near $\frac{1}{2} \beta_{\text{crit}}$, though the crossover was not systematically studied before.

In our experiments, polystyrene beads of radius $a = 14.3 \pm 0.1$ $\mu$m and volume fractions $\phi$ of 0.01% and 0.1% are suspended in sucrose solution in a $d \times w \times h$ cell with horizontal dimensions $d = 2$ cm and $w = 3$ cm, and height $h = 30$ cm. The viscometer-measured viscosities of the sucrose solutions range from $\eta = 1$ to 10 cP with varying sugar concentration. The cell is immersed in a stirred water bath at a temperature $22.0 \pm 0.1$ $^\circ$C. The nearly uniform sizes of the particles limit particle concentration gradients due to polydisperse effects. Likewise the small cell dimension is chosen to limit bulk transport due to internal mixing and hydrodynamic diffusion of the front. Low Reynolds numbers (Re $\sim 10^{-3}$) and high Peclet numbers (Pe $\sim 10^3$) indicate that inertia and Brownian motion can be neglected. Since the density of the sucrose solutions is higher than that of the particles, the mean velocity of the particles in these experiments is always upwards to the top of the cell. Velocity fluctuations are measured by particle image velocimetry (PIV) with a CCD camera imaging an area 11.3 $\times$ 8.5 mm, with 5 mm depth of focus, $= h/4$ away from the top boundary. We determine the vertical velocity fluctuations by $\Delta V = (\langle V \rangle^2 - \langle V \rangle^2)^{1/2}$, where $V$ is the local vertical velocity component and $\langle V \rangle$ is the mean velocity in the window.

A linear stratification is established using the Oster double bucket technique.\(^\text{17}\) The first bucket is filled with dense sucrose solution with a known volume fraction of particles while the second bucket is filled with a higher volume fraction of particles in a less dense sucrose solution. The suspension from the first bucket is stirred continuously and passed through a flow diffuser onto the existing fluid surface in the cell; simultaneously the suspension from the second bucket is allowed to flow into the first bucket. By keeping these two flow rates equal, a stable linear particle concentration profile is created with larger (smaller) volume fraction at the top (bottom) of the cell. A side effect is that a stable sucrose density gradient is also established; but the sucrose density difference in the imaged area varies less than 5% across all runs and therefore should not appreciably affect the fluctuation measurements here. In principle, no sucrose variation is strictly necessary, but we found it difficult to establish small stable particle concentration stratifications without an accompanying sucrose stratification, the differences in suspension density due to the particle concentration stratification alone insufficient to prevent mixing near the flow diffuser in our setup.

Local initial volume fractions at selected heights are determined by measuring the transmittance of a laser beam through the cell with a photodiode and comparing against known transmittance-volume fraction calibration curves. This measurement is further augmented by pipetting small amounts of fluid from different heights of the cell after the experiment and weighing with a scale balance. Assuming complete mixing in the first bucket yields volume fractions and densities in good agreement with the transmittance measurements and with the measured weights of the pipetted volumes.

The resulting velocity fluctuations with time are plotted in Fig. 1. For a $\phi = 0.01\%$ suspension that is shaken carefully
and checked by transmittance and pipetting to verify there is no stratification, the velocity fluctuations are measured to be steady for the duration of the experiment. When stable linear concentration gradients are created, \( \phi = \phi_0 (1 - \beta z) \), the fluctuation each achieve steady values (sometimes after an initial transient), but have larger magnitudes. The experiment at \( \beta / \beta_{\text{crit}} = 59 \) appears to decay over a longer time scale, but still appears to reach a steady value that is then maintained until the front gets close to the imaged measurement area (not shown in the figure). In each of these cases, the velocity fluctuations and polydispersities are so small that the sedimentation front does not appreciably spread during the experiments, thus ensuring that the initial concentration gradients persist over time. When thevolume fraction is increased to \( \phi = 0.1\% \), the nonstratified suspension has fluctuations that decay for most of the duration of the experiment, in agreement with previous measurements, the difference between the 0.01% and 0.1% behaviors also being described by the convective model of, e.g., Refs. 7, 13, and 15. However, when a steep volume fraction gradient is initially imposed, the fluctuations for \( \phi = 0.1\% \) are both suppressed and relatively steady in time. As steeper stratifications are introduced, the fluctuations are suppressed further. We emphasize that each symbol denoting the velocity fluctuations plotted in Fig. 1 corresponds to a single experiment, because of the inherent difficulty in setting up the particle concentration gradient (cf. repeatedly stirring and observing the same suspension). Our data thus includes outliers such as the early times of the diamond symbols in the top half of the figure; nevertheless, we see that even that experiment appears to settle down to statistically steady fluctuations.

Figure 2 shows the time-averaged velocity fluctuations from the experiments versus the initially imposed stratification (ignoring data from the initial transients, and including additional experiments beyond those in Fig. 1). In the main figure, the measured \( \Delta V / |V| \) are plotted, and there is unsurprising separation between the data at \( \phi = 0.01\% \) and \( \phi = 0.1\% \), because the Poisson-level fluctuations (1) are larger at higher volume fraction. Because the unstratified experiment at \( \phi = 0.01\% \) yields relatively constant fluctuations, we use that experiment to determine the constant in Eq. (1), giving \( C = 0.39 \), which is of the same order as the scalar premultipliers for square-based cells. The vertical velocity variances for both concentrations collapse when rescaled according to the Poisson prediction (with this \( C \)), as shown in the inset. A power-law fit to the \( \Delta V^2 \) in the Fig. 2 inset indicates an exponent \(-0.43 \pm 0.07 \), in good agreement with the \(-2/5 \) model prediction.

To further test these results, we performed simulations of sedimenting point particles interacting with wall-modified dilute-limit hydrodynamics, using the technique described in Ref. 12. Each simulation conducted for this study consists of between \( 10^5 \) and \( 2 \times 10^5 \) point particles of hydrodynamic particle radius \( a \) settling in a \( d \times w \times h \) cell, with no-slip boundary conditions along the parallel side walls (separated by \( d = 150a \) or \( 200a \)), periodic boundary conditions in the long \( x \) direction (with \( w = 4d \) or \( 8d \) ), an impenetrable container bottom and top imposed by simple images (separated by \( h = 8d \) ), and hydrodynamic truncations of Ref. 12 of \( N_c = 31 \) or 63. No hindered settling was included here, nor was the effect of the experimental sucrose density gradients modeled in any way. Various linear particle concentration stratifications along the vertical direction were initialized with particle concentrations at the top of the cell between 0 and 100% of the bottom concentration. In all, 112 individual simulations were computed for initially stratified conditions representing 14 different \( \beta / \beta_{\text{crit}} \) values. For comparison, an additional 29 simulations at similar physical parameters were conducted with no initially imposed stratification.

Velocity fluctuations were measured in a window \( 2 \sim 3.2d \) from the container bottom, that is, from 0.25 to 0.4h. The velocity fluctuations in the simulations behaved similarly to the experimental measurements in Fig. 1, except that the variation with time is larger in the simulations here because of the smaller number of particles; for comparison, the experiments here contain \( O(3 \times 10^5 \sim 10^7) \) particles. We define the initial concentration at the middle height of the cell to be the reference \( \phi_0 \), and the total sedimentation time \( \tau = h/|V| \) (where \(|V| \) varies with the sucrose density gradient in the experiments, it is constant across the height of the simulated cells). We time-average between the mean settling velocity transports that concentration to the top of the measurement window (\( t = 0.25 \tau \)) and when this mean transport reaches the bottom window (\( t = 0.75 \tau \)), with the resulting time-averaged velocity variances in each direction for each simulation plotted versus stratification in Fig. 3. These velocity variances have been rescaled by the no-free-parameters Poisson prediction for each simulation geometry, consistent with the predicted transition from \( \Delta V^2 \sim \Delta V_{\text{Poisson}}^2 \) for \( \beta \leq \beta_{\text{crit}} \) to \( \Delta V^2 / \Delta V_{\text{Poisson}}^2 = (\beta / \beta_{\text{crit}})^{-2/5} \) for \( \beta > \beta_{\text{crit}} \).

In Fig. 4, we combine the experimental data with the
vertical velocity fluctuations from the simulations, the latter now averaged over 5 simulation runs at identical physical parameters. Interestingly, while the experiments and simulations are each separately in reasonable agreement with the theoretical \(-2/5\) scaling, the experimental and computational results do not lie on the same power-law fit, the experimental data instead shifted to smaller fluctuations at the same \(\beta/\beta_{\text{crit}}\). The data indicates that the experiments crossover to stratification control of the fluctuations at stratifications \(\sim 32\) times smaller than in the simulations. That is, comparing the dashed lines in Fig. 4, the line close to the stratification-controlled simulation data indicates crossover from Poisson statistics to stratification control at the \(\beta_{\text{crit}}\) defined in Eq. (2), while the line passing close to the experimental data extrapolates to indicate crossover from \(\Delta \bar{V}^2 = \Delta \bar{V}^2_{\text{Poison}}\) to stratification suppression at \(\frac{1}{32} \beta_{\text{crit}}\). We do not attempt further quantification here of the precise scalar multiple of the \(\beta_{\text{crit}}\) defined in Eq. (2) at which crossover to stratification suppression of velocity fluctuations occurs, in light of the discovery that the crossover can shift so significantly between our simulations and experiments.

Various differences between the experiments and simulations could be responsible for this shift in the crossover stratification. The simulations include only point-force hydrodynamics, but that approximation should be reasonable at the low \(\phi\) of the experiments. The simulated point-forces are additionally smoothed by the Fourier truncation, but results from different truncations do not appear to correlate in a noticeable way. Meanwhile, the procedure used to set up the initial stratified conditions in the experiments additionally introduced sucrose density gradients, yielding gradients in \(\langle V \rangle\) and viscosity, while the simulations modeled the motion of initially stratified particles through a homogeneous suspending fluid. The dimensionless velocities considered eliminate any direct influence of different local viscosities; but it remains possible that the sucrose density gradients influenced the experiments in some unforeseen way. Finally, a major difference is geometrical; the experiments have a ratio of horizontal dimensions \(w/d = 1.5\), while the simulations are in cells with \(w/d = 4\) and 8. Moreover the simulations are periodic in the long horizontal dimension, whereas the experiment is obviously influenced by the walls in the long horizontal dimension.

We hypothesize that the differences between the experimental and computational results here might be primarily driven by geometrical differences. Different geometries may also be relevant to understanding differences between various experiments. We note in particular that stratification control does not explain recent results with \(h/d = 2\) and square cross sections \((w = d)\),\(^{14}\) nor with \(h/d = 4\),\(^{15}\) while Nguyen and Ladd\(^{13}\) suggest (using simulations with \(h/d\) large and square cross sections) that different controlling mechanisms may be at work depending on experimental details. It is particularly likely that the statistically steady arguments underlying the stratification suppression scaling break down without \(h/d\) large, because the time scale of the initial swirl turnover becomes comparable to that of advection across the cell. Meanwhile, cross-sectional details might influence the precise crossover stratification because of differences in local mixing.

Ideally, the experimental and computational conditions would correspond more closely, but the simulation method used only provides a pair of no-slip side walls, and thus corresponds more closely to physical conditions with \(w \gg d\) so as to minimize the influence of periodic boundary conditions in the long horizontal dimension, while the smaller \(w/d\) ratio and the presence of the sucrose aided in setting up the initially imposed gradients in the experiments. In principle, future experiments could be conducted in cells with larger...
$w/d$ or further minimizing or even eliminating the sucrose density gradient.

Our initially stratified experiments and simulations are each consistent with the model for stratification suppression of velocity fluctuations, but with different constants setting the crossover stratifications. Other controlling mechanisms not studied here could of course prevail in other settings. Moreover, it is natural to question whether this scaling even makes sense at large stratifications where the suppressed correlation length becomes less than the interparticle scale. We note that the smallest swirl sizes deduced from $\ell/d = \Delta V^2/\Delta V^2_{\text{Poisson}}$, here correspond to a couple mean interparticle spacings in the simulations and are even below the mean spacings in the most stratified experiments. Meanwhile, we observe that the simulation data appears to indicate greater suppression of fluctuations beyond the model scaling at the largest stratifications considered. It remains possible that the $\Delta V^2/\Delta V^2_{\text{Poisson}} \sim (B/\beta_{\text{crit}})^{2/5}$ model needs modification for larger stratifications and cell-geometry effects, though it appears reasonably consistent with the experiments and simulations considered here.

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