A colloidal particle placed in an inhomogeneous solution of smaller non-adsorbing polymers will move towards regions of lower polymer concentration, in order to reduce the free energy of the interface between the surface of the particle and the solution. This phenomenon is known as diffusiophoresis. Treating the polymer as penetrable hard spheres, as in the Asakura-Oosawa model, a simple analytic expression for the diffusiophoretic drift velocity can be obtained. In the context of drying films we show that diffusiophoresis by this mechanism can lead to stratification under easily accessible experimental conditions. By stratification we mean spontaneous formation of a layer of polymer on top of a layer of the colloid. Transposed to the case of binary colloidal mixtures, this offers an explanation for the stratification observed recently in these systems [A. Fortini et al., Phys. Rev. Lett. 116, 118301 (2016)]. Our results emphasise the importance of treating solvent dynamics explicitly in these problems, and caution against the neglect of hydrodynamic interactions or the use of implicit solvent models in which the absence of solvent backflow results in an unbalanced osmotic force which gives rise to large but unphysical effects.

I. INTRODUCTION

Many coatings, from paints to cosmetics, form by the drying of a thin, initially liquid, film. The liquid film contains a dispersion of colloidal particles and other non-volatile species that are left behind after the liquid evaporates, and these form a solid film [1, 2]. Due to the importance of coatings made from drying colloidal suspensions, there is an extensive literature on this process [3–15]. As the solvent (usually water) evaporates, the liquid/air interface descends at some speed \( v_{ev} \), pushing the non-volatile species such as colloidal particles and polymer molecules, ahead of it (Fig. 1).

Drying is a non-equilibrium process, and it creates concentration gradients. The concentration of colloid and polymer particles is high in an accumulation zone just below the descending interface, and lower near the bottom of the film. When there are concentration gradients in a mixture, there will be diffusiophoresis. Diffusiophoresis is the motion of one species in response to a gradient in the concentration of another. In this work, we focus on the motion of colloidal particles in response to a gradient in the concentration of smaller polymer molecules. We find that this diffusiophoretic motion can be strong enough to exclude the colloidal particles from a top layer of the drying film, i.e. it can drive stratification into a layer of small polymer molecules on top of a layer of the larger colloid particles.

Recent experimental work [16–18] on making solid films via the evaporation of mixtures of colloidal particles, has found stratified dry films. Here we argue that diffusiophoresis provides a potential explanation for how this stratification occurs spontaneously during drying. There has also been recent computer simulation and modelling work on this phenomena [16, 18–22]. Indeed stratification was first discovered in simulations by Fortini et al. [16]. However, to our knowledge nearly all of the existing work used an implicit solvent, i.e. modelled or simulated the particles as diffusing in a uniform continuum of some viscosity \( \eta \), neglecting hydrodynamic interactions (a notable exception is Cheng and Grest [23] who use molecular dynamics with an explicit solvent). In models with an implicit solvent that is set to be uniform, only gradients in the chemical potentials of the colloidal particles, and hence in the osmotic pressure, are possible. It is the gradient in the chemical potential of the small colloidal species, or equivalently in the osmotic pressure, that drives stratification in these models [16, 19, 20]. Here we consider the solvent explicitly and find that in an evaporating film the solvent backflow leads to a counter-gradient in the solvent pressure, that we expect to balance the gradient in the osmotic

FIG. 1. A schematic of a drying liquid (cyan) film containing colloidal particles (dark blue) and smaller polymer molecules (orange). The evaporation speed is \( v_{ev} \).
FIG. 2. A schematic of a large particle (blue) immersed in a solution of AO-model polymer (orange) of radius $R$, in a solvent of smaller molecules (green). The particle excludes the polymer from layer of width $R$ (indicated by a dotted line). Along the horizontal direction, there are gradients of both the polymer contribution to the pressure ($\Pi$), and the solvent contribution ($p_s$).

pressure. Neglecting backflow, as has been done for simplicity in much of the current work, therefore appears to be an unjustified approximation.

The current resurgence of interest in diffusiophoresis has largely focused on particles driven by electrolyte gradients, where there are significant effects additionally arising from diffuse liquid junction potentials [24–36]. However, diffusiophoresis also occurs in gradients of uncharged solutes [37–42], and we will apply expressions from this literature to polymer solutions. Our model for a polymer is the Asakura-Oosawa (AO) model [43–45], where the polymer itself is ideal (no polymer-polymer interactions) but is excluded from a layer of solvent of thickness $R$ around each colloidal particle. The radius $R$ is comparable to the radius of gyration of the polymer. This is a very simple model, and we believe that this paper can also serve as a pedagogical introduction to diffusiophoresis, identifying the molecular origins of the phenomena and highlighting the importance of solvent backflow [39, 41, 46–48].

The remainder of this paper is divided into four sections. In the first section we obtain an expression for the diffusiophoretic drift velocity of a large colloidal particle in a solution of much smaller AO-model polymers. Then we compare our approach in which solvent flow is taken into account, with earlier work with an implicit solvent, where solvent flow was neglected. We then study diffusiophoresis in a drying film containing colloidal particles and smaller polymer molecules. Our final section is a conclusion.

II. DIFFUSIOPHORESIS IN THE ASAKURA-OOSAWA MODEL

We are interested in determining the diffusiophoretic drift velocity $U$ of a colloidal particle in a solution of much smaller polymer. There is a gradient $\nabla \rho_p$ in the concentration of the polymer. This is illustrated in Fig. 2.

A. Slip velocity at flat hard wall

As we are in the $R_c \gg R$ limit, the curvature of the surface of the colloid can be neglected, and we can thus start by considering the relative, or slip, velocity $v_s$ between a hard wall (surface of large colloid) and the polymer solution. To calculate this we consider the geometry shown in Fig. 3, where the wall and gradient are along the $x$ axis. For diffusiophoresis of a solid particle, the stresses and hence the velocity gradients are localised to the interfacial region [39]. Here this is a region of width $R$. The standard theory of diffusiophoresis [37, 39] yields the following expression for the wall slip velocity

$$v_s = -\frac{k_B T}{\eta} \left[ \int_0^\infty z (e^{-\beta \varphi(z)} - 1) \, dz \right] \nabla_x \rho_p,$$

where $\eta$ is the viscosity (assumed equal to the bulk viscosity), $\varphi(z)$ is the potential between the solute molecules and the wall, and $\beta = 1/k_B T$. This result applies to an ideal solute whose concentration gradient in the fluid is $\nabla_x \rho_p$, and was first identified by Derjaguin and co-workers in 1947 [46].

For the AO model, the wall potential is just

$$\varphi(z) = \begin{cases} \infty & z < R, \\ 0 & z > R. \end{cases}$$

FIG. 3. A schematic of a flat wall (blue) in contact with AO-model polymer (orange) in a solvent of smaller molecules (green). The wall excludes the polymer from layer of width $R$, the top of this layer is indicated by a dotted line. There are gradients of both the polymer and solvent concentrations, parallel to the wall along the $x$ axis. The flat wall is assumed stationary, and then the fluid flows to the left. $\gamma$ is the wall/solution surface free energy.
With this potential, the integral in square brackets in Eq. (1) is just minus the first moment of the width of the exclusion region, i.e., \( \frac{1}{2} R^2 \). Then the slip velocity

\[
v_s = \frac{R^2 k_B T}{2\eta} \nabla_x \phi_p
\]  

(3)

(this basic result can be found in Anderson’s review [39]).

Note that the slip velocity results in flow away from low concentrations of the polymer and towards higher concentrations. This can be understood as a wall-bounded Marangoni-like flow away from where the surface tension is low, and towards where it is high [47]. This motion reduces the total wall/solution surface free energy, as the region of low AO-model polymer concentration expands.

We show in Appendix [A] that the AO-model polymer contributes \( \rho_p k_B T R \) to the surface tension, and so the surface tension is highest where the concentration of AO-model polymer is highest, driving flow of the solution to these high polymer concentration regions.

Eq. (1) rests on the fact that in a mixture relaxing by diffusion, the hydrostatic pressure \( p \) is uniform, so that a gradient in the osmotic pressure \( \Pi \) is balanced by a countergradient in the solvent contribution to the pressure. The hydrostatic pressure is uniform because it is a ‘fast’ variable which relaxes via solvent flow, and so gradients in \( p \) are quickly eliminated. (We also assume other gradients, for example, in the temperature are negligible.)

But the osmotic pressure (and hence counterbalancing solvent contribution to the pressure) only relax via diffusive motion of the polymer, which is much slower. This consideration will be crucial in considering implicit versus explicit solvent models. Details of the gradients in the interfacial region are in Appendix [B].

B. Diffusiophoretic drift of large colloidal particle

For a particle of size \( R_c \gg R \), curvature of the interface is negligible. Then, the diffusiophoretic drift velocity of such a particle in a stationary fluid is \( U = -v_s \), where \( v_s \) is the above slip velocity at particle’s surface. This apparently trivial result hides a great deal of subtlety in terms of the underlying low Reynolds number flow problem [37, 39, 49]. Thus one has

\[
U = \Gamma \nabla \ln \phi_p, \quad \Gamma = -\frac{\phi_p k_B T}{2\eta R} \sim -\phi_p D_p.
\]  

(4)

where we have introduced a diffusiophoretic drift coefficient \( \Gamma \). We also define \( \phi_p \equiv \rho_p R^3 \) as the dimensionless polymer concentration (i.e., packing fraction). The final order-of-magnitude scaling estimate rests on the Stokes-Einstein expression for the diffusion coefficient of the polymer, \( D_p \approx k_B T/(\eta R) \).

The drift is away from the high polymer concentration regions, i.e., it tends to cause the large colloid to segregate from the smaller polymer. This can be understood as being driven by the reduction in surface free energy when the colloid moves to regions of lower polymer concentration where the surface tension between the colloid particle and the polymer solution is lower.

Colloidal particles can also move under the action of gravity (i.e., in sedimentation). But it is important to note that diffusiophoretic motion and sedimentation are fundamentally different [48, 50]. In sedimentation there is an external force (gravity) acting on the particle and this causes the falling particle to set up a long-ranged \( 1/r \) flow field in the surrounding liquid. By contrast in diffusiophoresis, the stresses and velocity gradients are largely localised to the interfacial region (here of thickness \( R \)), and the flow outside the immediate interface is much weaker and decays as \( 1/r^3 \) [39, 48].

C. Typical diffusiophoretic drift velocities

We now estimate how fast are typical diffusiophoretic drift velocities. For water at room temperature, \( k_B T \approx 4 \times 10^{-21} \) J, and \( \eta \approx 10^{-3} \) Pa s. We consider a polymer of radius \( R \approx 10 \) nm, with a maximum concentration \( \phi_p \approx 1 \). Then the diffusiophoretic drift coefficient \( \Gamma \approx -200 \mu m^2 s^{-1} \). Thus if the length scale of the gradient \( \lambda_G = 100 \) \( \mu m \), then the drift velocity \( U \) is of order 1 \( \mu m \) per second, and independent of the size of the large particle [51].

As noted in recent experimental work [27, 32, 39], diffusiophoresis can be considerably more effective than diffusion at transporting micron-sized colloidal particles over large distances. Rates of transport by directed motion (diffusiophoresis, flow, etc) can be compared to rates by diffusion via Pécel number. Here the Pécel number comparing diffusion to diffusiophoretic motion for the colloidal particle is

\[
P_{\text{coll}} = \frac{U \lambda_G}{D_c} \sim \frac{\phi_p D_p}{D_c}
\]  

(5)

where \( D_c \) is the diffusion coefficient of the colloid. We used the scaling result for \( \Gamma \) in Eq. (4) to obtain the final expression, invoking \( U = \Gamma \nabla \ln \phi_p \) and \( \nabla \ln \phi_p \sim 1/\lambda_G \). Because of this the Pécel number is independent of the length scale of the gradient in the polymer concentration. For large colloidal particles \( D_c \ll D_p \) and if \( \phi_p \) is not too small, diffusiophoretic motion will always be faster over all length scales (larger than \( R_c \)) on which there is a gradient.

III. EARLIER WORK ON DRYING FILMS, WHERE AN IMPLICIT SOLVENT WAS USED

In earlier work that one of us (RPS) was involved in, an osmotic imbalance mechanism was proposed to explain the motion of large colloidal particles in a concentration gradient of smaller colloidal particles [16, 21]. This model was used as a possible explanation for the stratification seen in the drying film experiments [16, 18]. Zhou et al.
proposed what is in effect a simple dynamic density functional theory (DDFT), while Howard et al. proposed a more advanced DDFT; see Appendix D for a discussion of DDFT. In all these models, there is no explicit solvent. The Langevin dynamics simulations also use an implicit solvent. There the solvent is replaced by friction against a stationary background, plus a corresponding noise term. The models all use the Stokes expression \((6\pi \eta R_c v)\) at velocity \(v\) for the drag in a stationary fluid. In the experiments there is, of course, a solvent (water).

The osmotic imbalance mechanism argues that the gradient in the concentration of the smaller colloidal species gives rise to an imbalance in the osmotic pressure across the diameter of the large particles, leading to a drift velocity of the larger colloid. (The same argument would apply for a colloid particle in a polymer solution.) The work of Zhou et al. and of Howard et al. is similar in the sense that they too have models for stratification, in which there is an implicit solvent. Earlier work on colloids in evaporating films has also used an implicit solvent.

In Ref. the size of the effect was estimated as follows. The osmotic pressure difference across the particle diameter is of the order \(R_c \nabla (\rho_p k_B T)\). This gives rise to a force

\[
F \sim -R_c^2 \nabla (\rho_p k_B T) \tag{6}
\]

since the area over which the osmotic pressure difference acts is of order \(R_c^2\). We see that this is essentially equal to the volume of the large particle multiplied by the osmotic pressure gradient, as in the generalised Archimedes principle which applies in sedimentation equilibrium. Ref. then argued that the force \(F\) leads to a drift velocity \(U\) in accordance with the Stokes mobility, \(U = F/(6\pi \eta R_c)\). Using Eq. (6) this predicts a velocity

\[
U \sim \left(\frac{R_c}{R}\right)^2 \frac{\phi_p k_B T}{\eta R} \nabla \ln \rho_p \quad \text{[incorrect]} \tag{7}
\]

where we emphasise that we now regard this result as containing an incorrect scaling with particle size. Compared to the correct diffusiophoretic drift velocity in Eq. (1), the result above is a factor \((R_c/R)^2\) larger, and would overwhelm the former for \(R_c/R \gg 1\).

However, both this equation and the Langevin dynamics simulations neglect the solvent dynamics. We now discuss why this is incorrect, and in particular why solvent backflow critically modifies the above result. This observation is not new, and indeed Jülicher and Prost make the same point in a different context. The complete story can be found in a tour-de-force analysis by Brady. Our exposition takes a different, more informal approach, and we believe the essential point is the same.

The key point is that the force acting on the particle should additionally include a contribution from the solvent pressure gradient, as well as the osmotic pressure gradient from the solute. The solvent pressure gradient arises from solvent backflow, which must always be present in a counter-diffusing solute-plus-solvent mixture (i.e. relaxing by collective or mutual diffusion). Then, it is relatively simple to demonstrate that the solvent force apparently perfectly cancels the osmotic imbalance force.

We first recall the fundamental definition of the osmotic pressure \(\Pi\), as the difference between the actual (hydrostatic) pressure \(p\) in the system of interest, and the pressure \(p_s(\mu_s)\) in a system comprising pure solvent at the same chemical potential \([54, 55]\), viz.

\[
p = \Pi + p_s(\mu_s) \tag{8}
\]

When there is a gradient in the concentration of a colloidal or polymer species there will be a gradient in the osmotic pressure \(\Pi\). However, in the absence of an external body force (like gravity, in a sedimentation equilibrium), the hydrostatic pressure rapidly relaxes by bulk flow to become uniform (\(\nabla p = 0\)), on a time scale which is much faster than colloidal particles or polymers can diffuse to eliminate a gradient in the osmotic pressure. If \(p\) is uniform, we have that the osmotic pressure and the solvent pressure have equal and opposite gradients (\(\nabla \Pi = -\nabla p_s\)). Applying the above osmotic imbalance argument we conclude that the force arising from the solvent pressure gradient cancels that arising from the osmotic pressure gradient, or in other words the net force \(R_c^2 \nabla p = 0\) (since \(\nabla p = 0\)).

At this point we have apparently argued ourselves into a corner: if there is no net force, there can be no drift, and no stratification mechanism. However such a conclusion is incorrect, as the theory of diffusiophoresis shows. Whilst there is indeed no net force, there is a force dipole at the wall, arising from the solute structuring by the wall potential. In the present case (AO model) this can be turned into a quantitative argument which recovers the classical diffusiophoretic drift velocity, highlighting that it is in fact the gradient in the solvent pressure, in the exclusion layer adjacent to the particle, that causes the drift in the AO model case. This is explained in Appendix D.

To complete the story we return briefly to sedimentation equilibrium. In this case, it is the solvent pressure \(p_s(\mu_s)\) that must be constant, not the hydrostatic pressure \(p\). The former is constant because in equilibrium the solvent chemical potential \(\mu_s\) is everywhere the same; the latter (hydrostatic pressure) is not constant because there is a body force acting on the fluid as a whole, due to gravity. Invoking Eq. (8) once more, we see that \(\nabla p = \nabla \Pi\) in this case, and thus the integrated effect of the hydrostatic pressure gradient can be accounted to an osmotic pressure gradient, which leads to a force on the particle. Balancing this force against gravity recovers the generalised Archimedes principle. Thus in sedimentation equilibrium it is possible to ignore the solvent degrees of freedom.

In implicit solvent models, such as the Langevin dynamics simulations and the DDFT models used to de-
scribe film drying [19, 20, 22], solvent backflow is ignored. Zhou et al.’s [20] model is effectively a simple DDFT. The result is of course that the effect of $\nabla p_s$ is thrown out, leaving the osmotic pressure imbalance force $R_s^3 \nabla \Pi$. Put another way, if we simplify a model by using an implicit solvent that is defined to be uniform (and indeed static), then unbalanced gradients in the osmotic pressure appear as though they arise from an external force such as gravity. As we have just argued, this mistake leads to a grossly overestimated drift velocity, with the ‘wrong’ size scaling, as in Eq. (7). Therefore, implicit-solvent models should be used with caution when applied to dynamical situations, including interpreting experimental data on drying films.

Although the above story is complete, there is an alternative viewpoint that is worth touching upon. As Brady has emphasised [50], another way to think about the problem is to retain an explicit representation of the solute particles (polymers, in the AO model), and incorporate the correct physics by properly accounting for hydrodynamic interactions. From this viewpoint, there is indeed an imbalanced osmotic pressure gradient force on a colloid particle, and this indeed leads to a ‘primary’ drift velocity in accord with the Stokes mobility. However, this primary effect is almost completely cancelled by the long-range hydrodynamic interactions, which account for the solvent backflow.

Returning momentarily to the DDFT modeling, we show in Appendix D that this formalism gives rise to chemical potential gradients which can essentially be ascribed to unbalanced osmotic pressure gradients. Since the underlying density functional theories are known to be very accurate [50, 57], it would be nice to find a way to incorporate this into the modeling. The ‘missing link’ appears to be in the specification of the Onsager coefficients which relate the chemical potential gradients to the fluxes (see Appendix E). In the recent work [19, 20, 22] the Onsager coefficients were chosen to correspond to independent Stokes mobilities (i.e. friction against a stationary background), neglecting hydrodynamic interactions as an obvious simplification. With hindsight, it is clear that much more careful attention needs to be paid to this aspect. We leave this for future work.

IV. APPLICATION OF ASAKURA-OOSAWA MODEL TO FILM DRYING

We now turn to the application of the theory in Section I to the problem of stratification of colloid particles in a drying polymer solution, first considering the subproblem of the distribution of the polymer.

A. Distribution of Asakura-Oosawa model polymer in a drying film

The presence or absence of gradients in the concentration of the polymer, is determined by a competition between the evaporation speed $v_{ev}$, and diffusion of the polymer. This is quantified by the Péclet number [2, 11, 16, 21] for the AO-model polymer molecules in the drying film, $P_{film}$, which compares the time scale for diffusion to that for drying of the film. This is defined by

$$P_{film} = \frac{v_{ev} H}{D_p}$$

where $H$ is the film thickness. If $P_{film} < 1$ drying is slower than diffusion. Diffusion smooths out concentration gradients, and so here there are only weak gradients. But if $P_{film} \gtrsim O(1)$, diffusion cannot keep up with accumulation below the descending interface. Hence gradients will form and diffusiophoresis will occur.

The analysis is facilitated by a happy coincidence: since the polymer molecules in the AO model are by definition ideal, we can use the exact solution for a diffusing ideal gas in front of a moving impermeable wall to represent the effect of the drying solvent front. This leads us to consider a diffusing ideal gas between two impenetrable walls. A diffusing ideal gas satisfies

$$\frac{\partial \phi_p}{\partial t} = D_p \frac{\partial^2 \phi_p}{\partial z^2}$$

with the $z$ axis normal to the walls. The boundary conditions are the two walls, plus the initial condition that at time $t = 0$, we have a uniform distribution of the polymer, at an initial packing fraction $\phi_0$.

The bottom wall is fixed at $z = 0$. This models the substrate the film is on. The boundary condition here is just zero flux. The top wall is the solvent/air interface and is descending at speed $v_{ev}$. It therefore has a position

$$z_{int}(t) = H - v_{ev} t = (1 - t^*) H$$

at time $t$, where we have defined the reduced time

$$t^* = \frac{v_{ev} t}{H} \quad (\leq 1).$$

Here the boundary condition is

$$D_p \left( \frac{1}{\phi_p} \frac{\partial \phi_p}{\partial z} \right)_{z_{int}, t} = -v_{ev}$$

which is just the criterion for a diffusing ideal gas that the particles actually at the descending interface must be descending at the same speed as the interface.

In general we must solve for $\phi(z, t)$ numerically. However in the limit we are mostly interested in, where $P_{film} \gg 1$, there is range of times $t^*$ where a gradient is established but where the gradient is far from the bottom surface, i.e. where the accumulation zone beneath
the descending interface has not yet reached the bottom. Then the profile can be obtained by taking the $H \to \infty$ limit and using the exact solution in that limit of Fedorchenko and Chernov [58], see Appendix C. The full solution of Fedorchenko and Chernov is a little complicated, but after short time $\tau = t v^*_{ev}/D_p = 0.25$ when the gradient is being established, while the dashed curves are at the later time $\tau = 5$, when the exponential $z$ dependence is well established. At still later times, the height of the exponential continues to increase linearly, while the width remains constant.

The dynamics of the colloidal particles during drying of the film are determined by the competition between two velocities: $U$ and $v_{ev}$. To begin with we keep things as simple as possible and neglect diffusion (the $D_c \to 0$ limit). In this limit a particle below the interface moves down with speed $U(z, t)$, while actually at the interface the descending interface velocity $-v_{ev}$ forms an upper bound to the particle velocity. In other words, at the interface particles must descend at least as fast as the interface, but can go faster. Thus in a drying film the velocity of a colloidal particle is

$$v_c(z, t) \sim \begin{cases} 
\min(-v_{ev}, -v_{ev}P_{\text{film}}p_0 t^*) , & (z = z_{\text{int}}) \\
U(z, t) , & (z < z_{\text{int}})
\end{cases}$$

(17)

If we look at the behaviour actually at the interface, we have

$$v_c(z_{\text{int}}, t) \sim \begin{cases} 
-v_{ev} , & (P_{\text{film}}p_0 t^* \leq 1) \\
-v_{ev}P_{\text{film}}p_0 t^* , & (P_{\text{film}}p_0 t^* > 1)
\end{cases}$$

(18)

There are thus two regimes, the first is for $|U| < v_{ev}$, where the particles are swept up by the descending interface, forming a layer there. The second is for $|U| > v_{ev}$, where diffusiophoretic motion is faster than the speed of descent of the interface. Then the colloidal particles outrun the descending interface, and no particles accumulate at the interface. In this latter case the colloidal particles are depleted from a layer of thickness of order $D_p/v_{ev}$ below the interface, this is the thickness of the accumulated layer of polymer.

C. When $P_{\text{film}}$ and $p_0$ are large, stratification occurs with depletion of colloidal particles from a top layer

As the reduced time $t^* \leq 1$, then unless the product $P_{\text{film}}p_0 > 1$ depletion of the particles from the top surface will not occur during drying. In Fig. 5 we show...
Calculations are for $P_{\text{film}} = 20$ and $\phi_0 = 0.3$.

Note that there is always accumulation of the colloidal particles at early times as then there are no gradients. Gradients take time to build up. But at high enough polymer concentrations and Péclet numbers, this turns to depletion of the colloidal particles from the top layer, later on during drying.

In the absence of diffusion, the velocity $v_c(z,t)$ in Eq. (17) completely describes the time evolution of the positions of the particles during drying. The height of a particle at time $t$ is simply obtained by integrating $\frac{dz}{dt} = v_c$. Using this we further develop our understanding of the behaviour of the large particles in a drying film by considering the fate of a set of particles that at $t = 0$, are uniformly spaced across the height of the film. The results are shown as the red curves in Fig. 6 where each red curve followed upwards traces out the trajectory of a particle. The blue line is the position of the top interface. Trajectories that start near the bottom of the film ($i.e.$ left hand side of plot) are relatively unaffected and remain close to vertical, those in the middle of the film are pushed to the left away from the interface, while those near the top hit the descending interface for a time, and then later are pushed away from the top (assuming they can escape the interface).

Note that particles that start near the top of the initial wet film are caught by the descending interface early on in drying. But then later as the polymer gradient increases, these particles are pushed away from the descending interface. For the parameters used in Fig. 6 $U$ at the surface exceeds $v_{\text{ev}}$ at $t^* \approx 0.17$ (being the point where the green dashed separatrix touches the interface). At later times there is an expanding zone beneath the descending interface where $|U| > v_{\text{ev}}$ and so the colloidal particles are pushed down to further away from the interface. This exclusion region is shown in pink in Fig. 6 and there are no colloidal particles in it, but the concentration of the AO-model polymer is highest there. In this sense the AO-model polymer and the particles have kinetically segregated in the drying film, with the smaller AO-model polymer on top. This is as found in earlier work on mixtures of small and large colloidal particles [16] [21].

Unlike in earlier modelling work we have taken solvent flow into account. However, our model polymer is simple, and we have not considered colloid-colloid interactions, so our predictions may not apply for high colloid packing fractions. The width of the layer without colloidal particles is set by the width of the accumulation zone of the AO-model polymer, which is $D_p/v_{\text{ev}}$. For polymer of radius $R = 10 \text{nm}$ in water, $D_p \approx 20 \mu^2 \text{m}^{-1}$ and so at an evaporation rate of $v_{\text{ev}} = 0.1 \mu \text{m}^{-1}$ [8] [60], this gives a top layer $200 \mu \text{m}$ thick. This evaporation rate is typical for room temperature evaporation of water in still air, heating [60] can increase this by a factor of 10, which decreases the top layer thickness to $20 \mu \text{m}$.

It is worth noting that as typical film thicknesses are hundreds of microns [11] [8], and evaporation rates range from 0.1–1 $\mu \text{m}^{-1}$, that for polymers (or particles) $10 \text{nm}$ across, film Péclet numbers will be in the range 1–100. Film Péclet numbers for larger colloidal particles will then be in the range $10^7$. However, for small molecules ($e.g.$ ions, co-solvents) we expect film Péclet numbers $\ll O(1)$ under most circumstances. Thus diffusion due to gradients in small molecules will typically be small, as there will be no gradients to drive them. Our Eq. (19) encompasses this, as it predicts stratification occurs only for large Péclet numbers, which corresponds to large $R$ because $P_{\text{film}} \sim 1/D \sim R$ (with a Stokes-Einstein diffusion constant).

We note that significant concentration gradients of small molecules or ions can be produced by surface evaporation in saturated porous media [61]. This suggests interesting diffusiophoretic phenomenon may be observable in these situations, and indeed the partner phenomenon of diffusio-osmosis may be relevant ($i.e.$ pore-scale flow produced by a wall slip velocity, as in Eq. (1)).

We have not considered diffusion of the colloidal particles, but we expect this to only perturb our results. To observe diffusiophoretic effects we require for the polymer that $P_{\text{film}} > 1$. As the colloidal particle is larger than the polymer, the relevant Péclet number for the colloid particle is $P_{\text{film}} = v_{\text{ev}} H/D_c \gg 1$. Therefore, during drying, the colloidal particle will only be able to diffuse short distances, the ratio of the distance diffused to $H$ is...
\[(t^*/P_{\text{film}}')^{1/2} \ll 1.\]

V. CONCLUSION

Drying of a liquid film is an inherently out-of-equilibrium process. It will always lead to gradients, and these gradients can drive diffusiophoresis. To explore this we studied a very simple model system: diffusiophoretic drift of large colloidal particles in response to a gradient in polymer molecules. We used the Asakura-Oosawa polymer model. Due to its simplicity, we obtained a very simple analytic expression for the diffusiophoretic drift velocity, our Eq. (4). Also due to its simplicity, the physical mechanism is clear: the polymer increases the surface tension of the interface between the colloidal and the polymer solution. This drives motion of the particle towards lower polymer concentrations, as in a wall-bounded Marangoni effect. The velocity gradients associated with this motion are localised to the interface. This interface extends out from the particle surface to a distance of order the polymer size. The resulting diffusiophoretic drift is opposed by viscous dissipation largely localised to this interfacial region.

When there is diffusiophoresis in a drying liquid films there are two competing velocities: the evaporation velocity \(v_{\text{ev}}\), and the diffusiophoretic drift velocity \(U\). For large colloidal particles, when at the top interface \(v_{\text{ev}} > |U|\) the particles accumulate here, but when \(|U| > v_{\text{ev}}\) diffusiophoresis pushes particles away from the top interface. This creates a layer beneath the descending interface where there are no colloidal particles.

Since we expect the essential physics to apply also to mixtures of large and small colloid particles (see below), this provides an explanation for the experimental results of Fortini et al. [10]. Martin-Fabiani et al. [17] and Makepeace et al. [18]. They studied dried films made from mixtures of large and small particles, and found under some conditions that the large particles were excluded from the top of the dried film, and there were only small particles there. The films stratify during drying, with the small particles on top.

Earlier models [16–19, 21] also predicted this stratification but as discussed in section [11] these models neglected the effect of solvent backflow. This omission does not change the direction of stratification, but it does significantly overestimate the effects of diffusiophoretic drift. Neglecting backflow results in a drift velocity that increases as the square of the radius of the particle. When the solvent backflow is taken into account, the correct diffusiophoretic drift velocity is independent of colloid particle size, and much weaker. Although this is the case, we have demonstrated in section [14] that the correct diffusiophoretic drift velocity still predicts stratification, under experimentally accessible conditions, and in accord with experimental observations.

There is growing interest in diffusiophoresis, and not only is the AO model an especially simple example, polymers have a number of advantages for engineering controlled diffusiophoretic motion. Their effect can be tuned by varying their size, and as they are larger than ions, they diffuse more slowly, making it easier to establish the concentration gradients needed.

In this study we have chosen the AO model due to its simplicity, but the experiments on drying films use near-hard-sphere mixtures [16–18]. The diffusiophoretic drift coefficient \(\Gamma\) scales as \(\gamma/\eta\) (where note \(\eta\) should be the viscosity in the structured layer adjacent to the large colloid surface). Hard sphere interactions increase both \(\gamma\) and \((\text{bulk}) \eta\), but neither effect is dramatic at volume fractions up to around 20–30%. So, at all but high packing fractions, we expect the AO model to be roughly quantitatively predictive of the behaviour of colloidal suspensions at comparable packing fractions.

The approximations involved in this assumption can be addressed in part by using more accurate models of hard sphere suspensions, such as are available from modern density functional theory (DFT). The resulting dynamic DFT models are very appealing [19, 22], but it is clear, as we have identified at the end of section [11] that work needs to be done to address the neglect of solvent backflow effects in this approach.

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Appendix A: Wall surface tension in the Asakura-Oosawa model

Here we calculate the surface tension \(\gamma\) in the AO model, for a polymer solution against a non-adsorbing hard wall. The surface tension is the excess (over bulk) of the grand potential per unit area of the surface

\[
\gamma = \gamma_0 + \int_0^\infty (\omega(z) - \omega_b) dz
\]

where \(\gamma_0\) is the surface tension of pure solvent, \(\omega(z)\) is the grand potential density at a height \(z\) above the wall, and \(\omega_b\) is the bulk grand potential density.

As the AO model is ideal, the grand potential density \(\omega[\rho(z)]\) is just the ideal term plus the interaction with the wall

\[
\omega(z) = \rho k_B T [\ln(\rho/\rho_b) - 1] + \rho \varphi
\]

where \(\rho_b\) is the bulk polymer density. The variational principle \(\delta \omega/\delta \rho(z) = 0\) is satisfied by the Boltzmann distribution, \(\rho(z) = \rho_b \exp(-\beta \varphi)\). So, we have

\[
\gamma = \gamma_0 - \rho_b k_B T \int_0^\infty (e^{-\beta \varphi} - 1) dz.
\]
In the AO model this is easy to evaluate, and we get
\[ \gamma = \gamma_0 + \rho_b k_B T R. \] (A4)

Thus non-adsorbing polymers increase the surface tension by an amount equal to the osmotic pressure \( \rho_b k_B T \), multiplied by the depletion layer thickness. Note that the surface excess is
\[ \int_0^\infty (\rho - \rho_s) \, dz = \rho_b \int_0^\infty (e^{-\beta \varphi} - 1) \, dz = -\rho_b R. \] (A5)

This is negative, thus the increased surface tension can be viewed as a result of the generic Gibbs adsorption isotherm result, i.e. \( \partial \gamma / \partial \mu = \rho_b R \) where \( \mu = k_B T \ln \rho_b \).

Diffusiophoretic drift can now be understood as a consequence of a wall-bounded surface tension gradient [47]. In particular we can write \( v_s = (L^*/\eta) \nabla \gamma \), as though the surface tension gradient (force per unit area) is localised at a height \( L^* \) above the actual surface. In this simplified picture, the fluid undergoes uniform shear in the interfacial region \( 0 < z < L^* \). For the AO model, \( L^* = \frac{1}{2} R \). Compared to the actual flow field solved next, this picture is certainly oversimplified, but nevertheless is useful for gaining ad hoc insights.

**Appendix B: Flow field in interfacial region**

To obtain a better understanding of the molecular origins of diffusiophoretic drift we consider a polymer solution gradient next to a non-adsorbing wall, as shown in Fig. [3] and solve for the flow field. In this geometry the velocity is only along \( x \)-direction, and is only a function of \( z \), i.e. we have to compute \( v_z \). The relevant component of the Stokes equation is
\[ \eta \frac{d^2 v_x}{dz^2} = \nabla_z p. \] (B1)

In the bulk fluid (\( z > R \)), the hydrostatic pressure \( p \) is uniform, but in the interfacial region (\( z < R \)) the AO-model polymer is excluded and so in our model the relevant pressure is that of the solvent. Thus although in the bulk there is no gradient in the hydrostatic pressure, in the interface there is a gradient in the hydrostatic pressure, and this gradient is equal to the gradient in the solvent pressure in the bulk. As we have asserted in section [11] the solvent pressure gradient satisfies \( \nabla_z p_b = -\nabla_z H = -k_B T \nabla_z p_b \) where \( \nabla_z p_b \) is the polymer concentration gradient in the bulk. So we have
\[ \nabla_z p = \begin{cases} -k_B T \nabla_z p_b, & z < R \\ 0, & z \geq R \end{cases} \] (B2)

We insert Eq. (B2) in Eq. (B1), and integrate twice applying the boundary conditions \( v_z = 0 \) at \( z = 0 \) and \( dv_x/dz \to 0 \) as \( z \to \infty \) (in practice, this holds at \( z = R \) in this model). This yields the velocity profile
\[ v_x(z) = \begin{cases} [z(2R - z)k_B T/(2\eta)] \nabla_z p_b, & z < R \\ [R^2 k_B T/(2\eta)] \nabla_z p_b, & z \geq R \end{cases} \] (B3)

This comprises parabolic (half-Poiseuille) flow in the depletion layer, continuous with plug flow in the bulk. The resulting wall slip velocity \( v_s = v_x(\infty) = v_x(R) \) is identical to Eq. (3).

To anyone familiar with the derivation of the classic result in Eq. (1) this would not be surprising, but it perhaps sheds an interesting light on the mechanism in the present problem. In words: the gradient in polymer concentration is necessarily associated with a counter-gradient in the solvent chemical potential (to maintain constancy of overall bulk hydrostatic pressure). This generates a real solvent pressure gradient in the depletion layer adjacent to the large particle surface. This pressure gradient drives a thin film flow, resulting in an effective wall slip velocity on the scale of the particle. This leads to diffusiophoretic drift of a suspended particle. As noted, this drift is in the direction of reducing the interfacial free energy of the colloid particle in the polymer solution.

**Appendix C: Fedorchenko and Chernov solution**

Fedorchenko and Chernov obtained an analytic solution [58, 59] for a diffusing ideal gas in an infinite system below a descending wall. We are interested in a thin film not an infinite (\( H \to \infty \) system), however we can apply their result here, but only for \( P_{film} \gg 1 \).

Fedorchenko and Chernov’s solution [58, 59], applied to our AO-model polymer, can be written in terms of
\[ \phi_b(z, t) = \phi_0 [1 + \Delta(z, t)] \] (C1)

for \( \phi_0 \) the initial uniform concentration of polymer in the film. Here \( \Delta(z, t) \) is given by
\[ \Delta(z, \tau) = \left( \frac{\tau}{\tau} \right)^{1/2} \exp \left( -\frac{(z' + \tau)^2}{4\tau} \right) \]
\[ + \frac{1}{2} (1 - z' + \tau) e^{-z'} \text{erfc} \left( \frac{z' - \tau}{2\tau^{1/2}} \right) \]
\[ - \frac{1}{2} \text{erfc} \left( \frac{z' + \tau}{2\tau^{1/2}} \right) \] (C2)

with \( z' = |z - z_{int}|/(D_p/v_{ev}) \) and \( \tau = tv_{ev}^2/D_p = t^* P_{film} \).

After an initial transient, it simplifies to
\[ \Delta(z, t) \approx \frac{tv_{ev}}{D_p} \exp \left( -\frac{|z - z_{int}|}{D_p/v_{ev}} \right). \] (C3)

This holds for \( \tau \gg 1 \) (i.e. \( t^* P_{film} \gg 1 \)).

Note that this gives an accumulation zone below the interface of width \( D_p/v_{ev} = H/\rho_{film} \). When \( P_{film} \gg 1 \), this is much less than the initial film thickness. Thus the solution in an infinite system is very close to that in a thin film, except when \( t^* \) is close to one, because the accumulation zone ends far above the bottom substrate at \( z = 0 \). The time to establish this profile is \( D_p/v_{ev}^2 \), or in reduced units \( t^* = 1/P_{film} \). Note that at \( t = 0 \) the concentration is uniform, so there is no gradient.
Appendix D: Dynamic density functional theory

To see how the problem appears from the point of view of dynamic density functional theory (DFT), we start with an exact DFT for tracer (i.e. dilute) colloids in an ideal polymer solution [22].

\[
\beta f = \int \! d\mathbf{r} \rho_c(\ln \rho_c - 1) + \int \! d\mathbf{r} \rho_p(\ln \rho_p - 1) - \int \! d\mathbf{r} \! d\mathbf{r}' \rho_c(\mathbf{r}) \rho_p(\mathbf{r}') f_{cp}(\mathbf{r} - \mathbf{r}'),
\]

(D1)

where \( \beta = 1/k_B T \). The second term in this accounts for the colloid-polymer interaction, and features the Mayer function \( f_{cp} = e^{-\beta\phi_{cp}} - 1 \) which we leave general for the time being. This is essentially also the model proposed by Zhou et al. [20]. From this the colloid chemical potential is

\[
\beta \mu_c = \frac{\delta \beta f}{\delta \rho_c(\mathbf{r})} = \ln \rho_c(\mathbf{r}) - \int \! d\mathbf{r}' \rho_p(\mathbf{r}') f_{cp}(\mathbf{r} - \mathbf{r}').
\]

(D2)

A similar expression obtains for the polymer chemical potential. Taking the gradient of \( \beta \mu_c \) we find

\[
\beta \nabla \mu_c = (1/\rho_c) \nabla \rho_c - \int \! d\mathbf{r}' \rho_p(\mathbf{r}') \nabla f_{cp}(\mathbf{r} - \mathbf{r}'),
\]

\[
= (1/\rho_c) \nabla \rho_c + \int \! d\mathbf{r}' \rho_p(\mathbf{r}') \nabla f_{cp}(\mathbf{r} - \mathbf{r}'),
\]

\[
= (1/\rho_c) \nabla \rho_c + (1/\rho_c) \nabla \nabla \rho_p + \rho_c V_{cp} \nabla \rho_p,
\]

(D3)

where \( V_{cp} = -\int \! d\mathbf{r} \rho_c f_{cp}(\mathbf{r}) \) is the excluded volume between the colloid and polymer, specialising to the AO model case for which \( f_{cp}(\mathbf{r}) = -1 \) for \(|\mathbf{r}| \leq R + R_c\) and is zero otherwise. In the last step we assume that the polymer concentration is weakly varying on the scale of the colloid diameter. In a similar manner we find

\[
\nabla \rho_p \approx (1/\rho_p) \nabla \rho_p + \rho_c V_{cp} \nabla \rho_c.
\]

Now consider the matrix of Onsager coefficients which relate chemical potential gradients to fluxes, \( \mathbf{J}_i = L_{ij} \nabla \mu_j \). We shall suppose that the leading diagonal elements are \( L_{cc} = -\rho_c D_c/k_B T \) and \( L_{pp} = -\rho_p D_p/k_B T \), but as an ansatz keep the leading-order off-diagonal effect \( L_{cp} = -\rho_c \rho_p X/k_B T \) where the prefactor \( X \) is unknown at this point. Then the fluxes are given by

\[
\begin{pmatrix}
\mathbf{J}_c \\
\mathbf{J}_p
\end{pmatrix} =
\begin{pmatrix}
\rho_c D_c & \rho_c \rho_p X \\
\rho_c \rho_p X & \rho_p D_p
\end{pmatrix}
\begin{pmatrix}
1/\rho_c & V_{cp} \\
V_{cp} & 1/\rho_p
\end{pmatrix}
\begin{pmatrix}
\nabla \rho_c \\
\nabla \rho_p
\end{pmatrix}.
\]

(D4)

(cancelling \( \beta = 1/k_B T \) throughout). On multiplying through

\[
\begin{pmatrix}
\mathbf{J}_c \\
\mathbf{J}_p
\end{pmatrix} \approx - \begin{pmatrix}
D_c & \rho_c (V_{cp} D_c + X) \\
\cdots & D_p
\end{pmatrix}
\begin{pmatrix}
\nabla \rho_c \\
\nabla \rho_p
\end{pmatrix}.
\]

(D5)

In this we have dropped terms in the diagonal elements which are \( O(\rho_c) \) since we suppose we only have tracer amounts of colloid. Similarly, the lower left off-diagonal term is irrelevant since it multiplies \( \nabla \rho_c \) and makes a negligible contribution to colloid. Thus we arrive at \( \mathbf{J}_c = -D_c \nabla \rho_c - \rho_c (V_{cp} D_c + X) \nabla \rho_p \) and \( \mathbf{J}_p = -D_p \nabla \rho_p \). The term proportional to \( \nabla \rho_p \) in \( \mathbf{J}_c \) is also proportional to \( \rho_c \). We therefore identify it as a drift term, corresponding to a colloid drift velocity \( \mathbf{U} = -(V_{cp} D_c + X) \nabla \rho_p \). Note that although \( X \neq 0 \) corresponds to a 2nd order term in the Onsager matrix, it has been promoted to a 1st order correction in the drift velocity, essentially because \( \rho_p \) in the Onsager coefficient cancels \( 1/\rho_p \) in the chemical potential gradient.

If \( X = 0 \) the drift velocity can be written \( \mathbf{U} = -(D_c/k_B T) V_{cp} \nabla (\rho_c k_B T) \), with the interpretation (reading right to left) that the osmotic pressure gradient in the polymer solution generates a buoyancy force as in the generalised Archimedes principle, which when multiplied by the Stokes mobility gives the drift velocity. This is the approximation made by Zhou et al. [20], and the low density limit of Howard et al. [19]’s model, which now clearly corresponds to the argument made by Fortini et al. [16]. But our central claim is that this neglects solvent backflow, and overestimates the true diffusiophoretic drift velocity. Therefore \( X = 0 \) is inadmissible.

To recover what we claim to be the correct diffusiophoretic drift, we therefore expect that the prefactor \( X \) should be negative, and large enough to cancel the leading size dependence in the bare Stokes result. One could therefore envisage rather crudely ‘patching up’ the DDFT by including an off-diagonal Onsager coefficient consistent with the above arguments. We leave this for future investigation, though we note that the much more sophisticated analysis by Brady [30] identifies the exact way that hydrodynamic interactions conspire to cancel (most of) the bare Stokes result.

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