The role of sample height in the stacking diagram of colloidal mixtures under gravity

Thomas Geigenfeind and Daniel de las Heras

Theoretische Physik II, Physikalisches Institut, Universität Bayreuth, D-95440 Bayreuth, Germany

E-mail: delasheras.daniel@gmail.com

Received 29 September 2016, revised 4 November 2016
Accepted for publication 16 November 2016
Published 21 December 2016

Abstract

Bulk phase separation is responsible for the occurrence of stacks of different layers in sedimentation of colloidal mixtures. A recently proposed theory (de las Heras and Schmidt 2013 Soft Matter 9 8636) establishes a unique connection between the bulk phase behaviour and sedimentation-diffusion-equilibrium. The theory constructs a stacking diagram of all possible sequences of stacks under gravity in the limit of very high (infinite) sample heights. Here, we study the stacking diagrams of colloidal mixtures at finite sample height, h. We demonstrate that h plays a vital role in sedimentation-diffusion-equilibrium of colloidal mixtures. The region of the stacking diagram occupied by a given sequence of stacks depends on h. Hence, two samples with different heights but identical colloidal concentrations can develop different stacking sequences. In addition, the stacking diagrams for different heights can be qualitatively different since some stacking sequences occur only in a given interval of sample heights. We use the theory to investigate the stacking diagrams of both model bulk systems and mixtures of patchy particles that differ either by the number or by the types of patches.

Keywords: sedimentation, colloidal mixtures, gravity, patchy colloids, stacking diagram

(Some figures may appear in colour only in the online journal)
phase diagram of a given binary system. The theory is based on the concept of sedimentation paths. Each sedimentation path, which is a line in the plane of chemical potentials representation of the bulk phase diagram, describes the state of the mixture under gravity, in sedimentation-diffusion-equilibrium. Using this theory the stacking diagrams of mixtures of spheres and platelets [18] and mixtures of platelets and non-adsorbing polymers [19] were obtained. Also, very recently, van Roij and coworkers have obtained the stacking diagrams of mixtures of thin and thick colloidal rods [9]. Although in all these cases the bulk phase diagrams of the colloidal systems are relatively simple, the resulting stacking diagrams are extremely rich and show complex stacking sequences. These works are focused on the limit of very high (infinite) sample heights. This idealized limit is very relevant in experimental work since the height of the test tube is typically larger than the gravitational lengths of the colloids.

The interplay between micro confinement and colloidal sedimentation has been experimentally and theoretically investigated [20]. However, little attention has been paid to the influence of the total (macroscopic) sample height in colloidal sedimentation. A remarkable exception is the experimental work of Jamie et al [21], in which the properties of the gas-fluid interface of a polymer-colloid mixture are analyzed as a function of the overall height of the container. By systematically changing the total sample height while keeping the polymer-colloid concentrations fixed, the interfacial properties were found to move towards the critical point. Theoretically, it has been shown that varying the sample height might lead to a change in the stacking sequence in mixtures of colloids and nonadsorbing polymers [16, 19].

Here, we use the theory of [18, 19] to study sedimentation-diffusion-equilibrium of colloidal mixtures with finite height samples. We first give a full account of the theory for the case of finite height samples. Next, we systematically investigate the role of sample height in the stacking diagrams of colloidal mixtures. To this end, we apply the theory to model binary systems. That is, systems with generic bulk phase diagrams typical of model Hamiltonian which we however do not explicitly specify. Finally, and as an application of current interest, we study sedimentation in patchy colloidal mixtures. Patchy colloids are functionalized colloids that interact via a directional and valence-limited potential [22, 23]. In our systems the two species of the mixture differ in either the number or in the types of patches. The bulk phase diagrams of these mixtures have been previously analyzed [24, 25] using Wertheim’s theory [26]. Here, we obtain the stacking diagrams at different heights using only the bulk phase diagrams as input. The stacking diagrams are much richer than the corresponding bulk phase diagrams. Moreover, we show that the sample height is a crucial variable in sedimentation-diffusion-equilibrium of colloidal mixtures. The stacking diagrams for the same mixture but for different sample heights differ not only quantitatively but also qualitatively. For example, some stacking sequences occur only in a given range of sample heights.

2. Theory

2.1. The sedimentation path

Consider a colloidal mixture under gravity in a sedimentation vessel of sample height $h$. The gravitational potential for each species $i = 1, 2$ is $m_i g z_i$, where $m_i$ is the buoyant mass of species $i$, $g$ is the acceleration due to gravity, and $z_i$ is the vertical coordinate (we set the origin of coordinates, $z = 0$, at the bottom of the sample). Using a local density approximation [8, 18, 19, 27], we define a height-dependent local chemical potential for $0 \leq z \leq h$ for each species

$$
\mu_i(z) = \mu_i^b - m_i g z_i,
$$

where $\mu_i^b$ is the bulk chemical potential, i.e. the chemical potential in absence of gravity. The local density approximation assumes that for each $z$ the state of the sample is analogous to a bulk system (no gravity) with chemical potentials given by (1). This constitutes a very good approximation provided that the correlation lengths are small compared to the gravitational lengths, $\xi_i = k_B T/ m_i g$ with $k_B$ the Boltzmann constant, and $T$ the absolute temperature. Combining the expressions for the local chemical potentials, see (1), and eliminating the height variable $z$ we find

$$
\mu_i(z) = a + s \mu_i^b,
$$

where both $a$ and $s$ are constants given by

$$
a = \mu_i^b - s \mu_i^b,
$$

$$
s = m_i/m_i^b = \xi_i/\xi_i^b.
$$

The finite size of the sample $0 \leq z \leq h$ is translated into a range for the local chemical potentials

$$
0 \leq \frac{\mu_i^b - \mu_i}{m_i g h} \leq 1, i = 1, 2.
$$

Equations (2) and (4) represent a line segment, which we refer to as the sedimentation path, in the plane of chemical potentials. The sedimentation path describes how the local chemical potentials vary along the height coordinate in the vessel. Each point in the sedimentation path corresponds to the state of the sample at a given $z$.

The sedimentation path is directly related to the stacking sequence, i.e. the sequence of stacks of different phases that appear under gravity. If a path crosses a boundary between two phases in the phase diagram, e.g. a binodal, an interface appears in the vessel. The sedimentation path provides a direct link between the bulk phase diagram of the mixture and the stacking sequence. An example of a sedimentation path and its corresponding stacking sequence is shown in figure 1.

A sedimentation path is fully described by its (i) slope, (ii) location in the $\mu_1 - \mu_2$ plane specified by a point on the path, (iii) direction and (iv) length. The slope is fixed by the ratio of the buoyant masses, see (3). The position is determined by the bulk chemical potentials in absence of gravity, and hence by the overall colloidal composition and concentration via a
change of variables using the equation of state of the mixture. The direction is given by the signs of the buoyant masses (note \( m_1 \) can be negative if the mass density of the solvent is higher than the mass density of the colloids). Finally, the length of the path is proportional to the height of the vessel since the difference in chemical potentials between the top \((z = h)\) and the bottom of the sample \((z = 0)\) is

\[
\Delta \mu = \mu_1(h) - \mu_1(0) = -m_1 gh.
\]  

(5)

2.2. The stacking diagram

We have shown how each sedimentation path is associated to a stacking sequence. The phase stacking diagram is the set of all possible stacking sequences for a given mixture.

**Infinite height.** For standard colloidal particles in typical sedimentation vessels the length of the sedimentation path is of several \( k_BT \) in the \( \mu_1 - \mu_2 \) plane. That is, the path extends over a big region of the bulk phase diagram of the mixture. Hence, a very relevant idealization is to consider the limit of very high (infinite) sample heights. Within this limit \([18, 19]\) a sedimentation path is a straight line of infinite length (not a line segment) in the plane of chemical potentials. Hence, a sedimentation path can be fully described by using only the slope of the path \( s \), and the crossing point between the path and the \( \mu_2 \) axis \( a \), see (3). The stacking diagram can then be represented in the \( s - a \) plane. There are three types of boundaries between different stacking sequences in the stacking diagram. Here we only describe each one briefly, see \([18, 19]\) for a full account:

(i) **Sedimentation binodals.** The set of all sedimentation paths tangent to a binodal in the bulk phase diagram is a boundary between two phases in the stacking diagram. The path labeled as (1) in figure 1 is an example. An infinitesimal small change in one or in both variables of the path, \( a \) and \( s \), can change the stacking sequence.

(ii) **Terminal lines.** The set of all paths crossing an ending point of a binodal in the bulk phase diagram is a boundary in the stacking diagram that we call the terminal line. The sedimentation path (2) in figure 1 is an example. An infinitesimal change of \( a \) changes the stacking sequence. An ending point can be e.g. a critical point, triple point, critical end point, etc.

(iii) **Asymptotic terminal lines.** The third type of boundaries in the stacking diagram is formed by those paths that are parallel to the asymptotic behaviour of a binodal. See the path (3) in figure 1. In this case, an infinitesimal change of the slope \( s \) alters the stacking sequence. Both the binodal and the path do not terminate at finite chemical potentials.

A binodal is not the only possible boundary between two regions present in the bulk phase diagram. For example, a percolating line dividing the bulk phase diagram into percolated and nonpercolated states is another type of a boundary between phases. Any boundary present in the bulk phase diagram generates boundaries in the stacking diagram (sedimentation binodals, terminal lines and asymptotic terminal lines).

For convenience we speak always of binodal lines but one should bear in mind that other lines also give rise to boundaries in the stacking diagram. The patchy colloid mixtures studied below feature such percolation lines.

**Finite height.** In this paper we focus on the stacking diagrams for finite height samples. There exist several possibilities to represent the stacking diagram for finite heights. In an experimental work one typically varies the concentration and composition of the mixture, while keeping the solvent and the mass density of the colloids unchanged. The sample height is, in principle, easy to adjust\(^3\) and hence forms a useful control parameter. Under these circumstances the slope and the length of the path in the \( \mu_1 - \mu_2 \) plane are fixed, see (3) and (5), and its position in the \( \mu_1 - \mu_2 \) plane varies. A sensible choice of variables for the stacking diagram is the plane of average local chemical potentials along the path \( \bar{\mu}_1 - \bar{\mu}_2 \). As the sedimentation paths are straight lines, the average local chemical potentials are just the local chemical potential evaluated at the middle of the sample \( \bar{\mu}_1 = \mu_1(z = h/2) \).

The stacking diagram for finite height samples in the \( \bar{\mu}_1 - \bar{\mu}_2 \) plane contains three possible types of boundaries between different stacking sequences. Two of them are sedimentation binodals originated from coexisting lines in the bulk phase diagrams and one boundary is due to the ending

\(^3\) Solvent evaporation might occur, changing the effective sample height and hence the concentration of colloids.
be the parameterization of the chemical potentials \( \mu_1 - \mu_2 \). Two phases A and B coexist at the binodal (solid black line). The binodal ends at a critical point (empty circle). The solid red lines are selected sedimentation paths (finite height). The arrow indicates the direction of the path from the bottom to the top of the sample. The coordinates of the middle point of the path are the average local chemical potentials \( \bar{\mu}_1, \bar{\mu}_2 \) as indicated in one of the paths. The top (bottom) path marked with (1) (2) (3) starts (ends) at the binodal. The path labeled as (2) is tangent to the binodal. The path (3) crosses the critical point.

points of the binodals. We describe each of them in detail in the following.

(i) Sedimentation binodal type I (SBI). The set of all sedimentation paths that either start or end at a binodal form a sedimentation binodal of type I. A path starts (ends) at a binodal if the bottom (top) of the sample is located at the binodal. The path labeled as (1) in figure 2 is an example. For each binodal in the bulk phase diagram there are two corresponding SBI in the stacking diagram. One SBI for those paths that end at the binodal and the other SBI for those paths that start at the binodal. Both SBI lines have the same shape as the bulk binodal. This type of boundary is not present in the case of infinite height because the paths do not have starting and ending points.

Let \( \mu_{2,C}(\mu_1) \) be the parameterization of the chemical potential of species 2 at a bulk coexistence line, such as a binodal, as a function of \( \mu_1 \). Then, the two corresponding sedimentation binodals of type I are given by

\[
\bar{\mu}_2(\bar{\mu}_1) = \mu_{2,C}(\mu_1) + m g \frac{h}{2},
\]

\[
\bar{\mu}_2(\bar{\mu}_1) = \mu_{2,C}(\mu_1) - m g \frac{h}{2},
\]

where

\[
\mu_1^\pm = \bar{\mu}_1 \pm m g \frac{h}{2}.
\]

Here, \( \mu_1^\pm \) (\( \mu_1^- \)) is the local chemical potential of species 1 at the bottom (top) of the sample.

(ii) Sedimentation binodal type II (SBII). The set of all paths tangent to a bulk binodal is also a boundary (sedimentation binodal type II) in the stacking diagram. See the path (2) in figure 2. This boundary is analogous to the sedimentation binodals in the case of infinite height. The SBII boundaries are straight lines in the stacking diagram. A SBII line is present if and only if the slope of the path is the same as the slope of the binodal at some point(s). Each point of a binodal sharing the same slope as the path generates a SBII line.

Let \( (\mu_1, \mu_2) \) be the chemical potentials of a point on a bulk binodal. Let its local slope be that of the sedimentation path. That is

\[
\frac{d\mu_{2,C}}{d\mu_1} \bigg|_{\mu_{1,s}} = s.
\]

Then, the associated SBI line is given by

\[
\bar{\mu}_2(\bar{\mu}_1) = \mu_{2,a} + (\bar{\mu}_1 - \mu_{1,a}) s.
\]

The finite size of the path limits the range of \( \bar{\mu}_1 \) to

\[
|\bar{\mu}_1 - \mu_{1,a}| \leq m g \frac{h}{2}.
\]

(iii) Terminal lines (TL). The terminal lines are, as in the infinite height case, the set of all paths that cross an ending point of a binodal. See path (3) in figure 2. For each ending point in the bulk phase diagram there is one and only one TL in the stacking diagram. The TL is always a straight line.

Let \( (\mu_{1,e}, \mu_{2,e}) \) be the chemical potentials of an ending point in bulk, such as a critical point, a triple point, etc. The corresponding terminal line is

\[
\bar{\mu}_2(\bar{\mu}_1) = \mu_{2,e} + (\bar{\mu}_1 - \mu_{1,e}) s,
\]

for

\[
|\bar{\mu}_1 - \mu_{1,e}| \leq m g \frac{h}{2}.
\]

In the three cases (i)–(iii) any infinitesimal displacement of the path changes the stacking sequence (except for the special case in which the displacement is such that the path moves along the boundary of the stacking diagram). The asymptotic terminal lines that occur in the case of infinite sample height do not appear at finite height since the slope of the sedimentation path is fixed and the paths are of finite length.

The three boundaries SBI, SBII, and TL divide the stacking diagram in different regions. Each region corresponds to a different stacking sequence. In order to identify each sequence we first select one point inside of the desired region. Next we plot the corresponding path in the bulk phase diagram such that we can determine the sequence by inspecting the crossings between the path and the binodals.

Once the stacking diagram has been calculated in the \( \bar{\mu}_1 - \bar{\mu}_2 \) plane, we can easily transform to any other set of variables provided that the equation of state of the mixture is known.
In order to ease comparison to experimental work, a sensible choice of variables for the stacking diagram is the \( \bar{\eta}_{\bar{i}} \) plane, where \( \bar{\eta}_{\bar{i}} \) is the average packing fraction of species \( i \).

\[
\bar{\eta}_{\bar{i}} = \frac{1}{h} \int_0^h dz \eta_i(z).
\]

Here, \( \eta_i(z) \) is the local packing fraction of species \( i \) at a distance \( z \) from the bottom of the sample. To obtain \( \eta_i(z) \) we first compute the local chemical potentials at \( z \) using equation (1), and then use the equation of state of the system \( \eta_i = \eta_i(\mu_1, \mu_2) \). The phase diagram in the \( \bar{\eta}_{\bar{i}} \) plane is then obtained by transforming the coordinates of the boundaries in the stacking diagram from \( (\mu_1, \mu_2) \) to \( (\bar{\eta}_{\bar{i}}, \bar{\eta}_{\bar{i}}) \). Other representations of the stacking diagram such as for example average osmotic pressure versus average composition are also possible, following a similar transformation procedure.

### 3. Results

We first apply our theory to obtain the stacking diagrams at finite height of different bulk model phase diagrams (section 3.1). Although the bulk phase diagrams do not correspond to real microscopic models, they are representative of the behaviour of typical colloidal mixtures. The model bulk phase diagrams provide relevant examples of possible topologies of the stacking diagrams. In section 3.2 we apply the theory to model binary mixtures of patchy colloids for which we use Wertheim’s theory to obtain the bulk phase diagram. Finally, in section 3.3 we compare the stacking diagrams at finite and infinite sample heights.

#### 3.1. Model bulk phase diagrams

One of the simplest possible bulk phase diagrams of a binary mixture is schematically represented in figure 3(a). There is a single binodal at which two phases A and B coexist. The binodal ends at a critical point (empty circle). The solid red lines are representative sedimentation paths (finite height) corresponding to samples with two different heights, \( h_1 \) and \( h_2 > h_1 \), as indicated. The slope of the path is \( s = 1 \) in both cases. The arrow indicates the direction of the path from the bottom to the top of the sample. The red circle on the path \( h_2 \) is located at the center of the path. Its coordinates are the average local chemical potentials along the path (from bottom to top). The sedimentation path corresponding to the point highlighted with a red circle in panel (b) is represented in the bulk phase diagram (a). The lower panels (c) and (d) show the same diagrams for the case of sedimentation paths with slope \( s = -1 \). The sedimentation binodals of type II are indicated by black dashed lines. The inset on the left of panel (d) is a close view of a region of the stacking diagram.

Figure 3. (a) Schematic bulk phase diagram of a binary mixture in the plane of chemical potentials \( \mu_1 - \mu_2 \). Two phases A and B coexist at the binodal (solid black line). The binodal ends at a critical point (empty circle). The solid red lines are representative sedimentation paths (finite height) corresponding to samples with two different heights, \( h_1 \) and \( h_2 > h_1 \), as indicated. The slope of the path is \( s = 1 \) in both cases. The arrow indicates the direction of the path from the bottom to the top of the sample. The red circle on the path \( h_2 \) is located at the center of the path. Its coordinates are the average local chemical potentials along the path (from bottom to top). The sedimentation path corresponding to the point highlighted with a red circle in panel (b) is represented in the bulk phase diagram (a). The lower panels (c) and (d) show the same diagrams for the case of sedimentation paths with slope \( s = -1 \). The sedimentation binodals of type II are indicated by black dashed lines. The inset on the left of panel (d) is a close view of a region of the stacking diagram.

4 This ignores phase coexistence phenomena in polydisperse systems.
The stacking diagram of the chosen model bulk phase diagram for the case of infinite height is shown in figure 5 of [19]. Here we focus on the case of finite height. In figure 3(b) we represent the stacking diagram ($\mu_1 - \mu_2$ plane) for two different heights $h_1$ and $h_2$ with $h_2 > h_1$. In both cases the slope of the paths are the same, $s = m_2/m_1 = 1$, and both buoyant masses are positive such that both local chemical potentials decrease from the bottom to the top of the sample. We show representative sedimentation paths in figure 3(a). Each sedimentation path in the $\mu_1 - \mu_2$ plane in the bulk diagram is a point in the $\mu_2 - \mu_1$ plane of the stacking diagram (the coordinates of the middle point of the path). The stacking diagrams contain two sedimentation binodals of type I (generated by paths starting and ending at the bulk binodal) and one terminal line (paths crossing the critical point). There are three possible stacking sequences, namely A, B, and AB. We label the sequences according to the order of different stacks from bottom to top.

Next, we study the same model bulk phase diagram but for sedimentation paths with a different slope, $s = -1$. See representative paths in figure 3(c). Here $m_1 > 0$ and $m_2 < 0$ such that $\mu_1$ ($\mu_2$) decreases (increases) from the bottom to the top of the sample. The slope of the path is, in this case, compatible with the slope of the binodal in the sense that there is one point at the binodal whose derivative equals the slope of the path, see (8). Hence, the stacking diagram contains a sedimentation binodal of type II which is formed by the set of paths that are tangent to the binodal in bulk. The boundaries of the stacking diagrams (figure 3(d)) are: two SDI lines, one SDII line, and one TL. These boundaries split the stacking diagram into five regions. The possible stacking sequences are A, AB, ABA, and BA. The ABA sequence appears when a path crosses the bulk binodal twice [8, 27].

This very simple example already shows the richness of the stacking diagram. It also suggests that the sample height plays a major role. The size of the area of the stacking diagrams occupied by each stacking sequence depends strongly on the height of the sample. For example, the AB region substantially increases with $h$, see figure 3(b). Two samples of different height and different stacking sequences might have the same composition and concentration of colloids (we will see examples in the next section). The height of the sample might have an even stronger influence on the stacking diagram, as we will demonstrate in the following.

In figure 4(a) we show a further model bulk phase diagram. There are three different phases: A, B and C. Three binodals for A-B, A-C, and B-C coexistence meet at a triple point. A phase diagram like this might correspond to a mixture in which the species 1 represents spherical colloids and the species 2 consists of e.g. elongated colloidal particles. The elongated particles can undergo isotropic-nematic and nematic-smectic phase transitions.

The stacking diagrams for this mixture are depicted in figure 4(b) for two different heights, $h_1$ and $h_2$, with $h_1 < h_2$. In both cases the slope of the path is $s = 1$ and both buoyant masses are positive. The boundaries in the stacking diagram are: six SDI lines (two for each of the three binodals), one SDII line (the slope of the path matches the slope of the B-C binodal at one point), and one TL line (originating from the triple point). The stacking diagrams for heights $h_1$ and $h_2$ differ substantially from each other, see left and right panels of figure 4(b), respectively. We observe two main differences between the diagrams for short and long samples:

First, the sedimentation paths for the small system ($h_1$) fit in the space between the A-B and B-C binodals of the bulk phase diagram, see an example in figure 4(a). Consequently the stacking sequence B occurs in the stacking diagram, figure 4(b) (left). In contrast, the stacking sequence B does not occur in the large samples ($h_2$). The B sequence is replaced by an ABC state, figure 4(b) (right). The sedimentation paths in this case are long enough such that they do not fit in the region

---

1 Note that sequences with a single stack, such as A, are actually one phase-systems and not proper sequences made of different stacks.
between the A-B and B-C binodals in bulk. Instead, the path must cross at least one of the binodals.

Second, the sequence CABC is generated by paths crossing the three binodals in bulk. This sequence is present only in the long samples, figure 4(b) (right). The paths corresponding to the short samples ($h_1$) are not long enough to cross the three binodals, and hence the CABC sequence is absent.

These examples illustrate how the stacking diagrams for different heights might differ qualitatively. By changing the overall height of the sample some stacking sequences are replaced by others (e.g. the B sequence for $h_1$ is replaced by ABC for $h_2$) and one also observes the occurrence of new sequences, such as the CABC sequence for $h_2$.

3.2. Mixtures of patchy colloids

We next apply our theory to patchy colloidal binary mixtures. We study two cases in which the species differ either by the number or by the type of patches.

3.2.1. Different number of patches. We model the colloids by hard spheres of diameter $\sigma$ with identical patches (spheres of size $\delta$) on the surface, see figure 5(a). If two patches overlap the internal energy of the system decreases by $\epsilon$. We use Wertheim’s first order perturbation theory [26] and a generalization of the Flory-Stockmayer theory of polymerization [28, 29] to compute the bulk phase diagram of the mixture. We follow exactly the same implementation of the theory as in [24]. Theory and Monte Carlo simulations for the bulk phase behaviour are in semi-quantitative agreement with each other [30, 31].

The species 1 has two patches, and the species 2 has three patches. The colloids with three patches undergo a phase transition between two fluids with different densities. With only two patches present the particles of species 1 can form only chains. The absence of branching prevents phase separation and there is no fluid–fluid phase transition in the pure system of species 1. In the mixture the transition between high and low density fluids ends at a critical point. See the binodal in the bulk phase diagram of the mixture shown in figure 6(a) for scaled temperature $k_BT/\epsilon = 0.09$. In addition to the binodal, the phase diagram contains a percolation line that divides percolated and non-percolated states. The system is percolated if the probability that a patch is bonded, $p_c$, is higher than the percolation threshold $p_T$. The percolation line intersects the binodal close to the critical point on the low density side. The high density phase (G) is an equilibrium gel or network fluid which is always percolated. The low density phase does not percolate (N) except for a very narrow region close to the critical point (G’). We refer the reader to [24, 30] for further details about the bulk phase behaviour of this mixture.

To proceed and to obtain the stacking diagrams we need to set the slope of the sedimentation paths and the height of the sample. We fix the gravitational lengths of the colloids to $\xi_1 = 5\,\text{mm}$ and $\xi_2 = 2\,\text{mm}$ (typical values for colloidal particles). Hence, the slope of the path is fixed to $s = \xi_1/\xi_2 = 2.5$. The stacking diagrams in the $\bar{\mu}_1 - \bar{\mu}_2$ plane for three different heights $h = 1\,\text{mm}$, 10\,mm, and 25\,mm are shown in figure 6(b). Each of them contains four SDI lines (two for the binodal and two for the percolation line) and two terminal lines (one for the critical point and one for the ending point of the percolation line). Six different stacking sequences are possible for this value of the slope: N, G, GN, G-N, G–‘G’, and G–‘G’N. We use a dash between two stacks in the stacking sequence, like in the G-N sequence, to indicate that the sedimentation path crosses the binodal. The absence of a dash, e.g. in the GN sequence, indicates that the path crosses the percolation line.

Once the stacking diagrams in the plane of average chemical potentials have been computed, we can transform the variables using the procedure described at the end of section 2. In figure 6(c) we show the bulk phase diagram in the plane of packing fractions. The bulk phase diagram can be interpreted as the stacking diagram in the limit of zero sample height for which the sedimentation path is just a point. In figure 6(d) we present the stacking diagrams for $h = 1\,\text{mm}$ and $h = 10\,\text{mm}$ in the plane of average packing fractions.

The number and types of stacking sequences remain the same for the sample heights investigated here. However, the region of the phase space occupied by each sequence significantly depends on the value of the sample height. We show a specific example in figure 7 in which we plot the density profiles of two samples with the same average packing fractions ($\eta_1 = 0.002$, and $\eta_2 = 0.35$), but different heights ($h = 25\,\text{mm}$ and 10\,mm). The corresponding state points are highlighted by green solid circles in the stacking diagrams of figure 6. Despite the average colloidal concentrations being the same, the stacking sequences differ: G-N for the sample with $h = 25\,\text{mm}$ and G for the case $h = 10\,\text{mm}$. Other values of the sample height and the gravitational lengths will result in identical phenomenology provided that the ratios $h/\xi_i$ with $i = 1, 2$, are unchanged.

3.2.2. Different types of patches. As a concluding example we study a binary mixture of patchy colloids with different types of patches. The species 1 (2) possesses three patches of type A (B), see figure 5(b) for an illustration. When two patches of type $\alpha$ and $\beta$ with $\alpha, \beta = \{A, B\}$ overlap, the energy
of the system decreases by $\epsilon_{0,3}$. The bulk phase diagram of this model has been studied theoretically [25] and by Monte Carlo simulations [31] for different values of the bonding energies $\epsilon_{0,3}$. The phenomenology that emerges is very rich as different types of gels can occur depending on the set of bonding energies. Here, we set $\epsilon_B = \epsilon$ (energy scale), $0.80\AA = \epsilon$, and $0.85\AB = \epsilon$. We fix the scaled temperature, as in the previous case, to $kT = \epsilon_{0,3}$.

The bulk phase diagrams in the planes of chemical potentials and of packing fractions are shown in figures 6(a) and (b), respectively. At the value of temperature considered only the species 2 (strongest bonds) undergoes a fluid–fluid phase transition. Hence, in the mixture there is only one binodal that ends at a critical point. In addition there are three percolation lines. One of these indicates whether the full mixture percolates, and the other two percolation lines indicate whether the individual species percolate. Although species 1 does not undergo a first order fluid–fluid phase transition at this temperature, it still undergoes a percolation transition. The percolation lines and the binodal divide the bulk phase diagram into five different regions. At low chemical potentials (and hence low densities) the system is non-percolated (N). The other four states are equilibrium percolated gels: (i) a mixed gel (M) in which the mixture percolates but none of the species percolates independently, (ii) a bicontinuous gel or bigel (B) in which the mixture and both species percolate, (iii) two gels ($G_i$, $i = 1, 2$) in which the mixture and the species $i$ percolate. See [25, 31] for further details about the bulk behaviour.

Figure 6. Bulk phase diagram of a binary mixture of patchy colloids with two (species 1) and three (species 2) identical patches in the plane of chemical potentials (a) and packing fractions (c). The temperature is $kT = \epsilon_{0,3}$. The black solid line indicates the binodal. The dashed line is the percolation line of the mixture. The empty circle indicates the critical point. The black solid circle is the ending point of the percolation line. The inset in (a) is a close view of the region near the critical point. (b) and (d) show the stacking diagrams of the mixture under gravity in the plane of average chemical potentials (b) and average packing fractions (d) for three different sample heights, as indicated. A dash between two letters, like in the sequence G-N, indicates that the sedimentation path crosses the binodal. The absence of a dash, such as e.g. in GN, indicates that the path crosses the percolation line.

Figure 7. Density profiles $\rho_i$ of species 1 (green solid line) and species 2 (orange dashed line), percolation threshold $\rho_T$ (dot-dashed blue line), and bonding probability $f_b$ (dotted black line) of a binary mixture of patchy particles with two (species 1) and three (species 2) patches under gravity. The height of the container is $h = 25$ mm (a) and 10 mm (b). The system is percolated if $f_b > \rho_T$. In both cases the average packing fractions of the colloids of each species are the same, $\bar{\eta}_1 = 0.0021$ and $\bar{\eta}_2 = 0.352$. The stacking sequences are G-N (a) and G (b), schematically represented in the upper right corner of the figure. The insets in (a) and (b) are close views of the $\rho_1$ profile. The red arrow in panel (a) indicates the position of the G-N interface.
Here, we study sedimentation-diffusion-equilibrium. As in the example above we chose the gravitational lengths to be $\xi_1 = 5 \text{ mm}$ and $\xi_2 = 2 \text{ mm}$, and study two different sample heights $h = 1 \text{ mm}$ and $10 \text{ mm}$. The resulting stacking diagrams are extremely rich, see figures 8(c) and (d), with more than 20 distinct stacking sequences. Again, the regions occupied for each stacking sequence depend on the sample height. In some cases the same stacking sequence occurs in a completely different range of average packing fractions when varying the sample height, see for example the sequence 11 (G2M) in figure 8(d). Even more important is the fact that the stacking diagrams for $h = 1 \text{ mm}$ and $h = 10 \text{ mm}$ are qualitatively different. There are several stacking sequences that are present only in one of the selected sample heights. For example, the sequence MG1 (number 10 in figure 8) is only present in samples with $h = 1 \text{ mm}$, and the sequence G2MG1M (number 25) occurs only for the case $h = 10 \text{ mm}$.

3.3. Infinite versus finite height stacking diagrams

We conclude the section with several comments regarding the connection between the stacking diagrams for infinite and finite samples. The main effect is the occurrence of new sequences in the case of finite height samples. The new sequences are formed by the removal of one or more stacks of the sequences for $h \to \infty$. In general, a sequence observed at finite height might be a truncated sequence of the infinite system. This observation has strong implications for the correct interpretation of observed stacking sequences in finite height samples.
The infinite height stacking diagram provides the set of possible sequences for different values of \( s \). Here, we have only analysed the value \( s = 2.5 \) of the slope for finite height samples. The infinite height stacking diagram shows that for other values of \( s \) further complex phenomenology occurs. For example, for negative values of the slope, i.e. \( m_2 < 0 \), it is possible to stabilize the sequence NGN-G which constitutes a reentrant percolation phenomenon. This sequence also occurs in two-dimensional binary mixtures of patchy colloids [10].

Experimentally one can change the slope of the path via the synthesis of colloids with cores of different materials [32, 33] or changing the mass density of the solvent. Hence, the full range of stacking sequences of a given colloidal mixture is, in principle, experimentally accessible.

4. Discussion and conclusions

Our theory is based on a local density approximation which assumes that for each \( z \) the state of the sample can be approximated by a bulk state. Non-local effects might modify the stacking diagrams. In particular, the theory neglects the surface tension of the interfaces between stacks in the stacking sequence. If one of the stacks is very narrow the surface tension of the upper and lower interfaces might be higher than the gain in free energy due to the formation of the stack, as observed in colloid-polymer model mixtures [27]. Under such circumstances, the final equilibrium stacking sequence might be different than that predicted by our local theory. This condition is analogue of capillary condensation/evaporation. Surface effects such as e.g. the occurrence of wetting and layering near the walls of the vessel might also modify the stacking diagrams.

Our theory can be easily extended to multicomponent systems since the sedimentation paths remain lines in the phase space of chemical potentials. Also, the theory is directly applicable to molecular systems. There, the gravitational lengths are orders of magnitude higher than in colloidal systems. Hence, to observe similar phenomenology one needs containers of considerable size, such as for example geological deposits.

We have obtained the stacking diagrams at constant sample height and fixed ratio of the buoyant masses. Other choices, such as for example keeping the colloidal concentrations fixed and varying the sample height, are also possible. A stacking diagram in which one of the variables is the height might be relevant to study the effects of slow solvent evaporation, which is a process that changes the total volume but keeps the particle number fixed.

We have shown that two samples with the same colloidal concentrations but placed in vessels of different heights might have different stacking sequences. We have also shown that the stacking diagrams might be qualitatively different for different heights. Therefore, the sample height plays a major role in sedimentation-diffusion-equilibrium experiments. This role is as important as for example the average colloidal concentrations. We conclude that the sample height should be carefully measured and specified in any sedimentation experiment.
We have focused on sedimentation-diffusion-equilibrium in colloidal mixtures. Future studies could consider the dynamics of sedimentation using dynamic density functional theory [34, 35] and power functional theory [36].

Acknowledgments

We thank Matthias Schmidt for very useful discussions and a careful reading of the manuscript. This work was supported in part by the Portuguese Foundation for Science and Technology (FCT) through project EXCL/FIS-NAN/0083/2012.

References

[18] de las Heras D and Schmidt M 2013 *Soft Matter* 9 8636
[34] Evans R 1979 *Adv. Phys.* 28 143