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Introduction

The addition of a second component to a colloidal dispersion generates a wealth of new phenomena, such as the induction of effective interactions of the primary colloidal component by adding a depletion agent,¹ the formation of complex ionic colloidal crystals by adding an oppositely charged component,² and the self-assembly of patchy colloidal mixtures into two interpenetrating networks.3,4 Binary colloidal crystals are promising candidates *e.g.* for photonic applications.⁵ Gravity can have a strong effect on colloidal mixtures because on typical length scales in the lab the gravitational energy and the thermal energy are comparable. The resulting observations are often very counter-intuitive, including denser particles floating on top of a fluid of lighter colloidal spheres,6 or the emergence of a floating nematic phase sandwiched between two isotropic phases in mixtures of colloidal platelets and spheres.7 The number of distinct layers in a sample in the sedimentation-diffusion equilibrium can be considerably large: e.g. a vertical stack of six different phases was observed in mixtures containing charged colloidal platelets.8 Much attention was devoted to sedimentation in charged colloidal systems.9 Although the investigation of the bulk phase behaviour of colloidal mixtures is often based on sedimentation experiments,10 the interpretation of the experimental findings, and in particular drawing quantitative conclusions about the bulk phase diagram, can be a very subtle issue. Understanding and controlling the stacking sequence of colloidal mixtures is of potential interest in industrial applications, such as for preventing phase separation in commercial

The phase stacking diagram of colloidal mixtures under gravity

Daniel de las Heras* and Matthias Schmidt

The observation of stacks of distinct layers in a colloidal or liquid mixture in the sedimentation-diffusion equilibrium is a striking consequence of bulk phase separation. Drawing quantitative conclusions about the phase diagram is, however, very delicate. Here we introduce the Legendre transform of the chemical potential representation of the bulk phase diagram to obtain a unique stacking diagram of all possible stacks under gravity. Simple bulk phase diagrams generically lead to complex stacking diagrams. We apply the theory to a binary hard core platelet mixture with only two-phase bulk coexistence, and find that the stacking diagram contains six types of stacks with up to four distinct layers. These results can be tested experimentally in colloidal platelet mixtures. In general, an extended Gibbs phase rule determines the maximum number of sedimented layers as a function of the number of binodals and their inflection points.

colloidal products and inducing demixing in order to sort colloidal particles by size or density.¹¹ In molecular mixtures rather than colloidal mixtures corresponding effects due to gravity arise on larger length scales, *e.g.* relevant for species segregation with depth in oil reservoirs.^{12,13}

The connection between sedimentation equilibria and thermodynamics in one-component systems has been previously analysed theoretically, see *e.g.* ref. 14–16, and investigated experimentally, see *e.g.* ref. 17. There is, however, a lack of systematic understanding of how to relate the bulk properties of a binary mixture to its sedimentation equilibria.

Here we show for the first time that the phenomenology of all possible stacking sequences of a colloidal or liquid mixture is directly related to the bulk phase diagram of the system, without requiring further information about the equation of state. A systematic stacking diagram follows from the bulk phase diagram by Legendre transform in a unique way. Here the Legendre transform acts on the features of the bulk phase diagram, such as the binodal(s), critical point(s) etc., rather than on the thermodynamic potential(s). The Legendre transform introduces the ratio of the buoyant masses as a control parameter. In experiments this could be varied e.g. by changing the solvent of a colloidal mixture. We apply the theory to predict the stacking diagram of a mixture of colloidal platelets, using density functional theory to first obtain the bulk phase diagram. The mixture models a binary smectite-gibbsite platelet system, and hence the results can be tested experimentally.

Theory

We base our theory on the chemical potential as the central quantity. In the presence of gravity, one can define a height- and species-dependent local chemical potential:^{7,18,19}

Theoretische Physik II, Physikalisches Institut, Universität Bayreuth, D-95440 Bayreuth, Germany. E-mail: delasheras.daniel@gmail.com

$$\psi_i(z) = \mu_i^{\rm b} - m_i g z, \tag{1}$$

where z is the vertical coordinate, $\mu_i^{\rm b}$ is the chemical potential of species *i*, *m_i* is its buoyant mass and *g* is the acceleration due to gravity. Eqn (1) immediately implies that the difference in the local chemical potentials between two different heights is experimentally accessible by measuring the difference in heights between the two points. Eliminating *gz* from (1) yields

$$\psi_2(\psi_1) = a + s\psi_1,\tag{2}$$

where the constants are $a = \mu_2^b - s\mu_1^b$ and $s = m_2/m_1$. Eqn (2) describes a straight line in the plane of local chemical potentials ψ_1, ψ_2 . The slope *s* of this "sedimentation path" is given by the ratio of the buoyant masses or, equivalently, by the inverse ratio of the gravitational heights, $s = \xi_1/\xi_2$, where $\xi_i = k_B T/(m_ig)$ is the gravitational height of species *i*; here k_B is the Boltzmann constant and *T* is the absolute temperature.

Eqn (2) attains great significance when combined with a local density approximation (LDA),¹⁸ which applies when all relevant correlation lengths in the system are small compared to all ξ_i ; this is analogous to dividing the system in small horizontal slabs, which are treated as individual equilibrium systems.^{12,13} Then one can assume that the state of the system at height *z* is analogous to a bulk state with chemical potential μ_i and that

$$\mu_i = \psi_i(z) \tag{3}$$

As a consequence, the sedimentation path (2) is directly related to the experimentally observed stacking sequence in the vessel.^{7,19} The local phase coexistence between A and B occurs at height z_{AB} , provided that $\psi_i(z_{AB}) = \mu_{i,AB}$ for both species; here $\mu_{i,AB}$ is the chemical potential of species *i* at bulk coexistence between phases A and B. As a result a (horizontal) AB interface at height z_{AB} is observed in the sample.

In order to illustrate these effects, we plot in Fig. 1 a schematic bulk phase diagram of a mixture with stable A and B bulk phases. We show a sedimentation path that starts in the region where B is stable, crosses the binodal, and ends in the region where A is stable. The corresponding stacking sequence consists of bottom B and top A, which we write as BA. Clearly, the thickness of each sedimentation layer is proportional to the difference in chemical potentials between its upper and lower interface, cf. eqn (1). The difference in chemical potentials between the crossing of the sedimentation path and the binodal and the bottom of the sample in Fig. 1 is simply $\Delta \mu_i = -m_i g h_{\rm B}$, with $h_{\rm B}$ being the macroscopic thickness of the bottom sedimentation layer. By neglecting effects due to the finite sample height (to which we will turn below), the conditions for a sedimented sample are fully determined by the parameters *a* and *s*, which we now treat as new (state) variables. Each pair a, s determines uniquely a sedimentation path (2). We will in the following identify three types of boundaries in the *a*, *s* plane, where the sedimentation behaviour changes qualitatively upon infinitesimal deviation of the parameters. We refer to these boundaries as the sedimentation binodal, the terminal line, and the asymptotic terminal line.



Fig. 1 Schematic representation of the bulk phase diagram of a colloidal mixture in the plane of chemical potentials μ_2 , μ_1 . The black-solid line represents the binodal where the phases A and B coexist. The red-dashed line represents a sedimentation path. Its direction (from bottom to top of the sample) is indicated by an arrow. The inset shows the stacking sequence and the relative thickness of each sedimentation layer corresponding to the sedimentation path. The dotted red line represents a sedimentation path tangent to the bulk binodal.

(i) *Sedimentation binodal.* As T = const, the Gibbs phase rule leaves one remaining free thermodynamic variable to parameterize $\mu_{i,AB}$. Choosing this parameter as μ_1 results in $\mu_{1,AB} = \psi_1$ and $\mu_{2,AB}(\mu_1) = \psi_2(\psi_1)$, from which one obtains at coexistence

$$\mu_{2,AB}(\psi_1) = \psi_2(\psi_1). \tag{4}$$

The marginal case is obtained when the slope of the sedimentation path equals the slope of the binodal, which implies that

$$\mu'_{2,AB}(\psi_1) = \psi'_2(\psi_1), \tag{5}$$

is simultaneously satisfied with (4); in (5) the prime denotes the derivative with respect to the argument. Multiplying (5) by ψ_1 , subtracting it from (4), and observing the structure in (2) yields

$$a_{AB}(s) = \mu_{2,AB}(\psi_1) - \psi_1 \mu'_{2,AB}(\psi_1), \tag{6}$$

where $a_{AB}(s)$ is the intercept of the marginal case. Eqn (6) establishes $a_{AB}(s)$ as the Legendre transform of the chemical potential representation of the bulk binodal. Hence any path (2), where the parameters are linked *via* $a = a_{AB}(s)$, is special in that it divides the *a*, *s* parameter space into distinct regions. These regions differ in that a qualitative change in the stacking sequence occur, namely the emergence of a floating phase (*e.g.* from A to ABA). All paths that satisfy (6) are tangent to the binodal (see Fig. 1).

(ii) *Terminal line.* The systems reach a critical point, when locally for both components

$$\psi_{i,\text{crit}}(z_{\text{crit}}) = \mu_{i,\text{crit}},\tag{7}$$

from which follows, upon using the general expression (2), that

$$a_{\rm crit}(s) = \mu_{2,\rm crit} - s\mu_{1,\rm crit},\tag{8}$$

which is linear in *s*, implying that a critical point corresponds to a straight (terminal) line in the *a*, *s*-space of sedimentation paths. Indeed the same reasoning can be applied to the triple point and other "special" points in the phase diagram, at which a binodal ends. In each of these cases the result is analogous to (8). The sedimentation paths that correspond to (8), *i.e.* that lie on the terminal line, are those that cross the special point.

(iii) Asymptotic terminal line. A bulk binodal that does not terminate at finite values of μ_i can either be connected to a phase transition of one of the pure subsystems or it can represent a large demixing region at very high chemical potentials. In both cases the bulk binodal tends to an asymptote with a well-defined slope,

$$\mu_{2,AB}/\mu_{1,AB} \to s_{\infty}, \tag{9}$$

and a corresponding (asymptotic terminal) line emerges in the stacking diagram, which is described by $s(a) = s_{\infty} = \text{const}$, for all values of *a*. The paths that are described by (9) are those that are parallel to the asymptote of the binodal.

Results

We show in Fig. 2 three schematic bulk phase diagrams in the μ_1, μ_2 plane (a, c, e) and the corresponding stacking diagrams in the a, s plane (b, d, f). The simplest example (a) consists of a binodal that ends at two critical points. This topology corresponds to a closed immiscibility loop in the pressure-composition or density-density plane, as predicted, e.g., for mixtures of patchy colloids.²⁰ Two sedimentation paths with the same slope but opposite directions correspond to stacking sequences with inverse order (e.g., AB and BA); in order to avoid this ambiguity we only consider the case $m_1 > 0$. The corresponding stacking diagram (b) contains one sedimentation binodal and one terminal line for each of the two critical points. Each terminal line is tangent to the end of the sedimentation binodal. The crossing point, where the two terminal lines intersect, represents the sedimentation path that connects the two critical points in the bulk phase diagram. The sedimentation binodal and the terminal lines divide the a, s plane into five different regions with differing stacking sequences. Those paths that cross the bulk binodal twice generate the stacking sequence BAB. This triple stacks occur although there is no triple point in the bulk phase diagram nor are there three different phases. A similar stacking sequence, a nematic phase sandwiched between two isotropic phases, has been theoretically predicted and experimentally observed very recently7 in mixtures of silica spheres and gibbsite platelets. In general, two consecutive phases in the stacking sequence coexist in bulk, but two nonconsecutive phases may or may not coexist in bulk and, as the example demonstrates, the same phase (B) can reenter the sequence. An obvious consequence is that the Gibbs phase rule does not apply to the maximum number of layers in the sedimentation equilibrium. Instead, the maximum number of sedimented layers in a mixture under gravity is $3 + 2(n_{\rm b} - 1) + n_i$, where n_b is the number of bulk binodals and n_i is the total number of inflection points in all binodals. This maximal



Fig. 2 Schematic bulk phase diagrams of colloidal mixtures in the plane of chemical potentials μ_1, μ_2 (a, c, e), and the corresponding stacking diagrams in the plane of slope *s* and intercept *a* of the sedimentation path (b, d, f); here we set $m_1 > 0$. Chemical potentials are given in arbitrary units. (g) Sedimentation phase diagram of the mixture of panel (e) in the α , r_{min} plane. In panels (a, c, e) solid lines represent binodals (modelled as Bézier curves), empty circles represent critical points, dashed lines indicate selected sedimentation paths (the arrows give the direction from bottom to top of the sample, labels indicate the corresponding stacking sequences). The triangle in (e) represents the triple point. In panels (b, d, f, and g) the solid lines indicate sedimentation binodals, dashed lines represent terminal lines, and dotted lines indicate asymptotic terminal lines. Different stacking regions are coloured and are labelled by their respective stacking sequence.

number of layers occurs if a sedimentation path crosses each binodal $2 + n_{b,i}$ times, with $n_{b,i}$ being the number of inflection points of each binodal. The occurrence of an inflection point in a binodal has been predicted in colloidal platelet–sphere mixtures.⁷ It is also a natural consequence of a binodal that connects two phase transitions in the pure components of the mixture. An example is the isotropic–nematic transition in binary platelet mixtures, to which we turn below.

We next consider case (c) where the AB phase transition persists in the pure system of species 2. The bulk binodal tends asymptotically to the value of μ_2 at the phase transition of the pure species 2 when $\mu_1 \rightarrow -\infty$. In the stacking diagram (d) an asymptotic terminal line appears: as laid out above, this vertical line is at a position *s* that gives the slope of the asymptote of the bulk binodal.

The complexity of the stacking diagram increases very significantly with the number of stable phases in the bulk phase diagram. In Fig. 2(e) we show a still rather simple bulk phase diagram with three bulk phases A, B and C that coexist at a triple point. The pure species 1 undergoes a BC phase transition that persists in the mixture and ends at the triple point. The BC bulk binodal tends asymptotically to the value of μ_1 at the phase transition of the monodisperse system of species 1 when $\mu_2 \rightarrow -\infty$ (vertical asymptote). The component 2 of the mixture undergoes an AB phase transition. The binodal has a horizontal asymptote and ends at the triple point. At chemical potentials above the triple point there is strong demixing between A and C. The AC binodal has asymptotically a well-defined slope, for which the general expression holds: $d\mu_2/d\mu_1|_{coex} = \Delta \rho_1/\Delta \rho_2$, with $\Delta \rho_i$ being the jump in density of species *i* at the phase transition. At very high chemical potentials both species approach the close packing density and the slope of the binodal will be constant. The corresponding stacking diagram, Fig. 2(f), has three sedimentation binodals (one for each bulk binodal), one terminal line due to the triple point, and two asymptotic terminal lines. The asymptotic terminal line of the BC bulk binodal is not visible because its slope is infinite. To overcome this problem, we show in panel (g) the stacking diagram in the α , r_{\min} plane, where r_{\min} is the distance of the sedimentation path to the origin in the μ_1 , μ_2 plane and tan $\alpha = s$, cf. Fig. 1. Here we use the sign convention that r_{\min} is positive (negative) for a clockwise (counterclockwise) path in order to discriminate between paths with identical values of α , but with negative values of s. We find six sedimentation binodals (two for each bulk binodal because in this plane we plot all possible paths, $0 \le \alpha \le 2\pi$, lifting any sign restriction on m_1), one terminal line, and six asymptotic terminal lines at $\alpha = 0, 0.35\pi, \pi/2, \pi, 1.35\pi$, and $3\pi/2$. Each pair α and $\alpha + \pi$ represents the asymptotic behaviour (and direction) of a bulk binodal in the stacking diagram. All intersections between sedimentation binodals and terminal lines define the boundaries of the sedimented phases. There are 22 different stacking sequences, which vary in complexity from the simple AC to the exotic CACBAB. Sedimentation monophases do not occur, as each path crosses at least one binodal. These examples demonstrate the extreme richness of the stacking diagram.

We next apply the theory to a mixture of infinitely thin circular hard platelets with the aspect ratio $R_2/R_1 = 1.4$, where R_i is the radius of species *i*. We use a microscopic geometry-based density functional to calculate the bulk phase diagram. The functional goes beyond the Onsager limit and has been used previously to analyze the bulk phase diagram of platelet–platelet mixtures with different aspect ratios.²¹ The pure fluid of hard platelets undergoes an isotropic–nematic (IN) phase transition as a function of the chemical potential. Further phases with

positional order, such as columnar or crystal phases, are not present in the model due to the vanishing thickness of the particles.²² The bulk phase diagram of the binary mixture is depicted in the μ_1 , μ_2 plane in Fig. 3(a). The IN bulk binodal connects the phase transitions of the pure components. The chemical potential of the small platelets, μ_1 , decreases when a small fraction of big platelets is added to the pure fluid of small platelets (big platelets favour the orientational order of the small platelets). The behaviour of μ_2 is the opposite: its coexistence value increases if a tiny fraction of small platelets is added to the pure system of big platelets. As a result, the bulk binodal, that connects both limits, has a curvature change and a maximum at intermediate values of composition.

The stacking diagram of the mixture is plotted in Fig. 3(b) in the α , r_{\min} plane. There are two sedimentation binodals and four asymptotic terminal lines at $\alpha = 0$, $\pi/2$, π and $3\pi/2$. We can identify six different sedimentation sequences, including a floating isotropic phase NIN, a double floating isotropic NINI and nematic ININ. Such states with four layers arise from the curvature change of the bulk binodal. The NINI four-stack can be understood as a fractionation between two double NI stacks, the lower one being rich in heavier platelets, and the upper one being rich in the lighter platelets. This interpretation also applies to the inverted ININ sequence, which appears in the region of negative buoyant masses.



Fig. 3 Phase diagram and stacking diagram of a binary mixture of colloidal platelets. (a) Bulk phase diagram as a function of the chemical potentials μ_1 and μ_2 of small platelets (species 1) and big platelets (species 2) with the size ratio $R_2/R_1 = 1.4$. The solid line represents the IN bulk binodal. The dashed line indicates a selected sedimentation path for a vessel with height h = 10 mm. The arrow gives the direction from bottom to top of this sample. The inset illustrates the corresponding stacking sequence. (b) Sedimentation phase diagram for infinite sedimentation paths in the α , r_{min} plane. The solid lines represent the sedimentation binodals, and the vertical dotted lines indicate the asymptotic terminal lines. (c) Sedimentation phase diagram in the plane of (scaled) reference chemical potentials $\beta \mu_D^h$, $\beta \mu_D^b$, *cf.* eqn (1), for sample height h = 10 mm, *i.e.* sedimentation paths of finite length. The solid lines represent the sedimentation binodals. Each colour in (b) and (c) represents a different sedimentation state, labeled by its corresponding stacking sequence upon increasing the height.

Real samples possess a finite height *h*, such that there is only an interval of the corresponding infinite sedimentation path accessible. In order to characterize such a state, four variables are required, which we choose to be the slope *s* of the path, the two reference chemical potentials μ_1^b and μ_2^b , cf. eqn (1), and the sample height *h*. We adopt the convention that the origin of the coordinate system is in the middle of the sample, such that -h/2 <z < h/2. As a consequence at z = 0 the local chemical potential $\psi_i(z) = \mu_i^{\rm b}$. We choose $\xi_1 = 0.805$ mm and $\xi_2 = 3.89$ mm, which is compatible with, e.g., a colloidal mixture of gibbsite platelets (species 1: average diameter 214 nm and thickness 10 nm) and smectite (species 2: average diameter 300 nm and thickness 1 nm) dispersed in water at room temperature (see e.g. ref. 7 and references therein). We obtain $s = \xi_1/\xi_2 = 0.207$ or $\alpha = 192^\circ$, and we choose the vessel height as h = 10 mm. An example of a sedimentation path and the corresponding sedimentation sequence for these conditions are shown in Fig. 3(a). The stacking diagram for the finite height is shown in Fig. 3(c). Due to the finite sample height, additional sedimentation binodals arise when a sedimentation path starts or ends at the bulk binodal. While for infinite sedimentation paths there are only two sedimentation phases, NI and NINI, for finite height we find six different stacks I, N, NI, NIN, INI, and NINI. Hence the main effect of the finite length of the sedimentation path is that new sedimentation stacks appear. These are formed by removing layer(s) on top or bottom of the infinite stacking sequences. Note that the μ_1^b, μ_2^b plane can be converted to the plane of average densities by calculating the density profiles and average density of those paths that form the phase boundaries of the stacking diagram.

Discussion

A colloidal mixture under gravity can be represented by a straight line in the plane of chemical potentials, which we refer to as a sedimentation path. A crossing between the sedimentation path and a binodal corresponds to an interface in the sample, which establishes a direct relationship between the path and the observed phase stacking sequence under gravity. We have developed a general theory that relates all possible stacking sequences of a colloidal mixture to its bulk phase diagram. We have shown how to group the stacking sequences in a stacking diagram that follows in a unique way from the bulk phase diagram. The binodals, their ending points, and their asymptotic behaviour are the three distinct elements that determine the boundaries between the different sedimentation states in the stacking diagram. The stacking and the bulk phase diagrams are linked by a well defined mathematical mapping based on the Legendre transform.

In order to characterize the sedimentation states, a suitable order parameter is the array of thicknesses, h_m , of the individual layers m = 1, 2, ... in a sedimented stack. When crossing a boundary in the sedimentation diagram, at least one of the $h_m \rightarrow 0$, which implies a continuous phase transition in the inhomogeneous systems. Finite-size and surface effects, that are beyond our LDA treatment, will modify and enrich this scenario *via* wetting at the upper and lower boundaries and capillary evaporation effects of very thin floating layers.¹⁹

The individual stacks are always in the order of decreasing total mass density, $\rho_m(z) = \sum_i m_i \rho_i(z)$. The condition $0 \ge d\rho_m(z)/dz \equiv \sum_{ij} m_i(\partial \rho_i/\partial \mu_j) d\psi_j(z)/dz$ follows by imposing the local thermodynamic stability along the sedimentation path, *i.e.* the matrix of partial derivatives $\partial \rho_i/\partial \mu_j|_{T,V,\mu k}$, where *V* is the system volume, is positive definite.²³ This is consistent with the findings in^{12,13} and is not in conflict with the possibility of denser particles floating on top of lighter ones, as recently reported by Piazza *et al.*,⁶ provided that the floating phase has lower mass density than the supporting phase.

Gravity induces very rich phenomenology even for the simplest of mixtures. For example, the binary hard core platelet mixture that we investigated has only isotropic and nematic phases stable in bulk. We found that the stacking diagram contains six different types of sequences, with up to four distinct layers. The maximum number of layers in a stacking sequence is not limited by the Gibbs phase rule for bulk phases. Instead, an extended Gibbs phase rule holds, which accounts for the maximum number of layers that can appear in the sedimentation-diffusion equilibrium. The relevant parameters are the number of binodals and the number of their inflection points, rather than only the number of components as is relevant in bulk. As a consequence, extremely rich stacking sequences can be found even for very simple mixtures. An example is the NINI stack that we predict to occur in the binary platelet mixture. Mixtures with three binodals can lead to stacking sequences with seven distinct layers in the simplest case with no inflection points.

Our results demonstrate that the ratio between the buoyant masses of both species, s, is a key parameter that controls the stacking sequence of a mixture. A change in the strength of gravity, i.e. by using centrifugal forces as in analytic centrifugation, leaves the slope *s* invariant. However, if the species are made of different materials, one can vary s experimentally by changing the density of the solvent, because $m_2/m_1 = v_2(\rho_{m,2} - \rho_s)/(v_1(\rho_{m,1} - \rho_s))$, where v_i is the particle volume, $\rho_{m,i}$ is the mass density of species *i*, and ρ_s is the solvent density. Alternatively, one could vary s by designing colloids with cores made of different materials.²⁴ While clearly the average density of colloids in the sample (related to a cf. eqn (2)) plays a major role in determining the stacking sequence, our consideration for the binary platelet mixture demonstrates that the total height of the sample is a further relevant parameter because it controls the accessible region of the corresponding infinite sedimentation path. As different intervals of an infinite sedimentation path can lead to distinct stacking sequences, the height of the sample should be carefully controlled in sedimentation experiments. See the discussion of the effect of finite sample height in the binary platelet mixture, Fig. 3.

Our approach is relevant for computer simulation work, as the sedimentation–diffusion equilibrium of colloidal mixtures can be predicted by simulating only the bulk phase behaviour, which can be less computationally demanding than performing simulations of inhomogeneous systems under gravity. Although we have restricted our study to the case of binary mixtures, the theory is valid for multicomponent systems, as sedimentation paths remain as lines in the space spanned by all chemical potentials, *cf.* eqn (1). Our theory can also be used to describe the sedimentation of mixtures with non-colloidal components. An example is mixtures of platelets and non-adsorbing polymers where the gravitational length of the polymers is much higher than that of the colloids. We have checked that the results from our approach, using the free-volume theory in ref. 25 to describe the bulk phase diagram of the mixture, match with those of the effective one-component approach by Wensink and Lekkerkerker²⁶ and the experiments by van der Kooij *et al.*²⁷ Our theory also applies to molecular mixtures, provided that the bare mass rather than the buoyant mass is used.

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