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Parasitic crystallization of colloidal electrolytes: growing a metastable crystal from the nucleus of a stable phase

Out-of-equilibrium growth of a metastable parasitic crystal (green) from the critical nucleus of the stable phase (red) occurs within the supersaturated fluid (grey). Despite its lower stability and higher nucleation barrier, a charged-disordered colloidal phase manages to crystallize from nuclei of the stable phase due to its enhanced kinetic crystal growth. By means of molecular simulations, we elucidate the different aspects behind this novel mechanism of crystallization.







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## growing a metastable crystal from the nucleus of a stable phase Ignacio Sanchez-Burgos, Da Adiran Garaizar, Da Carlos Vega, Db

Parasitic crystallization of colloidal electrolytes:

Eduardo Sanz b and Jorge R. Espinosa \*\*

Colloidal particles have been extensively used to comprehend the main principles governing liquid-crystal nucleation. Multiple mechanisms and frameworks have been proposed, through either experiments or computational approaches, to rationalise the ubiquitous formation of colloidal crystals. In this work, we elucidate the nucleation scenario behind the crystallization of oppositely charged colloids. By performing molecular dynamics simulations of colloidal electrolytes in combination with the Seeding technique, we evaluate the fundamental factors, such as the nucleation rate, free energy barrier, surface tension and kinetic pre-factor, that determine the liquid-to-solid transition of several crystalline polymorphs. Our results show that at a high packing fraction, there is a cross-over between the nucleation of the CsCl structure and that of a substitutionally disordered fcc phase, despite the CuAu crystal being the most stable phase. We demonstrate that the crucial factor in determining which phase nucleates the fastest is the free energy cost of the cluster formation rather than their kinetic ability to grow from the liquid. While at a low packing fraction, the stable phase, CsCl, is the one that nucleates and subsequently grows, we show how at moderate and high packing fractions, a disordered fcc phase subsequently grows regardless of the nature of the nucleating phase, termed parasitic crystallization. Taken together, our results provide a panoramic perspective of the complex nucleation scenario of oppositely charged colloids at moderate temperature and rationalise the different thermodynamic and kinetic aspects behind it.

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#### I. Introduction

Liquids can be often cooled down below their freezing temperature without undergoing spontaneous crystallization. The reason behind this common behaviour is that metastable liquids need to overcome a free energy barrier to transform into a stable solid phase. 1,2 The surmounting of such a barrier involves the formation of a critical nucleus of the solid phase, which, depending on the system and conditions, may range from a few to an arbitrarily large number of particles when close to coexistence (although within the range where experimentalists are able to determine nucleation rates, the critical nucleus ranges from tens to a few thousand particles). Nucleus formation is a stochastic process, which implies that foreseeing where and when the cluster will appear is not feasible. Besides,

critical nuclei are usually rather small (of a few nm) and exhibit fast growth after their emergence, making their observation extremely challenging.4-7

One of the most employed systems to gain comprehension about nucleation and crystallization processes is colloidal particles.<sup>8–13</sup> Besides being highly interesting systems per se for their many different technological applications, 14,15 they represent one of the few cases in which the formation of the nucleus can be directly tracked. <sup>4,7</sup> Due to their large size ( $\sim 0.5 \mu m$ ) and slow motion, colloids can be considered as big atoms, where the observation of the rising nucleus, and its size, shape, and post-critical growth can be experimentally accessible. 4,7,16,17

In complement, molecular simulations have been widely used to perform nucleation studies of many diverse systems, such as hard-spheres, <sup>18–25</sup> NaCl, <sup>26–32</sup> argon, <sup>20,33</sup> alloys, <sup>34,35</sup> methane hydrates<sup>36-38</sup> and water.<sup>39-48</sup> On one hand, computational methods are highly suitable to tackle liquid-crystal transitions since the typical size of the nuclei ideally fits within the feasible simulation system size range. On the other hand, the emergence of the critical nucleus and its subsequent growth also match the affordable timescales in computer

<sup>&</sup>lt;sup>a</sup> Maxwell Centre, Cavendish Laboratory, Department of Physics, University of Cambridge, J J Thomson Avenue, Cambridge CB3 0HE, UK. E-mail: jr752@cam.ac.uk

<sup>&</sup>lt;sup>b</sup> Departamento de Quimica Fisica, Facultad de Ciencias Quimicas, Universidad Complutense de Madrid, 28040 Madrid, Spain

simulations (from hundreds to thousands of nanoseconds). Nonetheless, since nucleation entails the surmounting of a free energy barrier, spontaneous crystallization rarely occurs. <sup>49</sup> Hence, to bias the formation of the critical nucleus, rare event techniques need to be employed. During the last 20 years, a huge effort in developing rare event methods to track nuclea-

tion by molecular simulations has been made. 50-54 However, a good example of how challenging nucleation studies are is the several discrepancies still present in the literature regarding the nucleation rate. The nucleation rate, J, is the number of nuclei per unit of volume and time that appear in a system under the given conditions. J is a key magnitude in characterising the nucleation phenomenon, since (1) it accounts for the likelihood of the system to undergo the liquid-crystal phase transition and (2) it can bridge predictions from experiments and computer simulations. Nonetheless, discrepancies in the nucleation rate of common substances, such as water, are currently under debate, not only between different simulation techniques<sup>39,55,56</sup> but also among different experimental setups.<sup>5,6,57</sup> Likewise, in hard-sphere colloidal crystal nucleation, a long-standing discrepancy among crystal nucleation rates from experiments and computer simulations still persists, 11,58,59 although several possible explanations have been recently discussed such as heterogeneous nucleation, <sup>60</sup> incomplete shear melting,<sup>61</sup> hydrodynamic effects on the crystal growth<sup>62</sup> and the nucleation rate<sup>63,64</sup> or sedimentation.<sup>65,66</sup>

In addition to these methodological, and experimental  $\nu$ s. numerical discrepancies in the nucleation rates of these systems, there is also a general debate in the nucleation community about the validity of Classical Nucleation Theory (CNT)<sup>67,68</sup> to successfully describe nucleation in widely diverse scenarios. A considerable body of literature has shown that CNT may lead to quantitative inaccuracies when predicting nucleation rates<sup>69-73</sup> (an excellent review on the assumptions, and therefore the limitations, of CNT is given in ref. 74). Some of these assumptions, such as the capillary approximation, 75 non-Markovian dynamics<sup>76,77</sup> or an inappropriate choice of the reaction coordinate<sup>78,79</sup> can lead to a breakdown of the CNT predictions. Nonetheless, CNT has also been shown to provide remarkably good predictions for many different systems such as for the liquid-crystal nucleation of the 3-D Ising model,80 hard-spheres,19,20 NaCl,28 aluminum alloys,81 supercooled high-pressure silica, 82 water, 20,56 and bubble nucleation of argon.83 These several open debates clearly highlight the extreme complexity of the nucleation phenomenon.

Within colloidal systems, a particularly interesting case is the one of oppositely charged colloids, <sup>84-86</sup> where the electrostatic interactions between colloids can be tuned such that different crystal structures <sup>87,88</sup> may arise in the phase diagram (see Fig. 1). <sup>87</sup> Due to this fact, polymorphic competition among different crystal phases can occur depending on the specific conditions, <sup>72,85</sup> and an interplay between the free energy barrier of the cluster formation and its kinetic ability to grow from the metastable liquid can be crucial.

In this work, we investigate the crystal nucleation scenario of oppositely charged colloids by means of computer simulations.

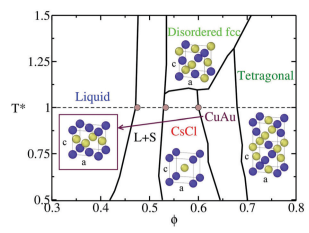


Fig. 1 Phase diagram in the  $T^*-\phi$  plane of the discontinuous Yukawa potential (HS-Y) parametrised for reproducing the behaviour of oppositely charged colloids, <sup>86</sup> taken from ref. 72 and 87. The unit cell of the different crystal structures is shown: CsCl (cubic symmetry), CuAu (tetragonal symmetry where  $c/a \sim 0.96$ ), tetragonal (tetragonal symmetry) and disordered fcc (cubic symmetry). The CuAu-tetragonal and CsCl-CuAu transitions are first order, but with a small density jump<sup>89</sup> (see also Fig. 2); for this reason, a line depicts these transitions. Note that the disordered fcc unit cell only illustrates one possible disordered distribution of the oppositely charged colloids. L + S stands for the liquid-solid coexistence region. The horizontal dashed line indicates the considered isotherm of this study ( $T^*=1$ ) and brown circles show the coexistence packing fractions predicted by our continuous model (PHS-Y). Note that the maximum packing fraction of the tetragonal phase is  $\sim 0.74$ , although to better visualise the tetragonal unit cell, this region has been expanded.

By developing a continuous potential version of the Yukawa model parametrised for reproducing the phase behaviour of colloidal electrolytes, 86,87 we perform highly-optimized parallel Molecular Dynamics (MD) simulations within the Seeding framework 90-92 to unveil the kinetic and thermodynamic interplay during the crystallization of several polymorphs with different crystal structures, CsCl, CuAu and a substitutionally ion-disordered fcc phase. In what follows, we show how, depending on the phase diagram region, the nucleating crystal phase may differ from the stable one, and that out-ofequilibrium post-critical growth of a parasitic phase can also take place independently of either the nucleating or the most stable crystalline phase under those conditions. Overall, our results provide a detailed characterisation of the key factors that determine the polymorphic crystallization scenario of oppositely charged colloids.

# II. A continuous potential for oppositely charged colloidal particles

Monodisperse binary mixtures of oppositely charged colloids are standardly modelled by using a purely repulsive hard-sphere (HS) potential in combination with a screened Coulombic-Yukawa interaction term,  $u_{\rm Yukawa}$ , of the form:  $^{86,87}$ 

$$u_{\text{Yukawa}} = \pm \varepsilon \frac{\sigma}{r} e^{-\kappa(r-\sigma)}$$
 (1)

where the sign is positive for equally charged colloids and negative for oppositely charged ones,  $\varepsilon$  accounts for the temperature-dependent interaction strength,  $\sigma$  is the molecular diameter of the colloids and  $\kappa$  is the inverse of the Debye length. We refer to the combination of the HS and Yukawa potentials as the HS-Y potential.

However, the HS-Y potential is discontinuous and, hence, not convenient for MD simulations. To be able to use MD simulations that can be efficiently parallelized, we substitute the HS potential by the successful pseudo-HS (PHS) model of Jover et al. 93 The PHS potential quantitatively reproduces many properties of the HS phase behaviour such as the melting pressure, 94 equation of state, 93 diffusion coefficient, 95 surface tension<sup>96</sup> and nucleation rate among many others:<sup>97</sup>

$$u_{\text{PHS}} = \begin{cases} \lambda_{\text{r}} \left( \frac{\lambda_{\text{r}}}{\lambda_{\text{a}}} \right)^{\lambda_{\text{a}}} \varepsilon_{\text{R}} \left[ \left( \frac{\sigma}{r} \right)^{\lambda_{\text{r}}} - \left( \frac{\sigma}{r} \right)^{\lambda_{\text{a}}} \right] + \varepsilon_{\text{R}}; & \text{if } r < \left( \frac{\lambda_{\text{r}}}{\lambda_{\text{a}}} \right) \sigma \\ 0; & \text{if } r \geq \left( \frac{\lambda_{\text{r}}}{\lambda_{\text{a}}} \right) \sigma \end{cases}$$
(2)

In this model, the diameter of the colloids is represented by  $\sigma$ ,  $\lambda_a$  = 49 and  $\lambda_r$  = 50 are the exponents of the attractive and repulsive terms, respectively,  $\varepsilon_R$  is the energy shift of the pseudo hard-sphere interaction and r is the distance between two PHS particles.

We refer to the combination of the PHS and the Yukawa potentials as the PHS-Y model. To reproduce with this model the phase behaviour of 50:50 binary mixtures of oppositely charged colloids, <sup>15</sup> we set  $\sigma = 3.405$  Å,  $\varepsilon_R/k_B = 119.87$  K and  $\kappa \sigma$  = 6, as described in ref. 72 and 87. Since the continuous version of the HS potential,  $u_{PHS}$ , can only capture the original HS behaviour at  $k_BT/\varepsilon_R = 1.5$  (which implies T = 179.81 K), we fixed the temperature in our simulations at T = 179.81 K and defined the temperature as  $T^* = k_B T / \varepsilon$ , modulating  $\varepsilon$  of the Yukawa term to vary the effective temperature of the system. Given that our study has been performed at  $T^* = 1$ , the value of  $\varepsilon/k_{\rm B}$  is set to 179.81 K. The chosen cut-off radius for the colloidal interactions is  $3\sigma$ . In what follows, we shall use reduced units: the reduced temperature  $T^* = k_B T / \varepsilon$ , the reduced pressure  $p^* = p\sigma^3/k_BT$ , the reduced density  $\rho^* = (N/V)\sigma^3$  and the volume fraction as  $\phi = (\pi/6)\rho^*$ . The unit of time is defined as  $\tau =$  $\sigma^2/6D_L$ , with  $D_L$  being the liquid diffusion coefficient at each corresponding state.

To implement the potential and carry out our simulations, we employed the MD LAMMPS package<sup>98</sup> with the following numerical details; the chosen timestep for the Verlet integration of the equations of motion was 0.0016  $\sigma \sqrt{m/k_BT}$  (m is the mass of the colloidal particles and is 20 u.a.m), for NVT simulations we used the Nosé-Hoover thermostat99,100 with a typical relaxation time of 0.16  $\sigma \sqrt{m/k_BT}$ , and for NpT simulations, we additionally applied the Nosé-Hoover barostat<sup>101</sup> with a typical relaxation time of 0.805  $\sigma \sqrt{m/k_BT}$ . The continuous PHS-Y potential described in eqn (1) and (2) is introduced in tabular form.

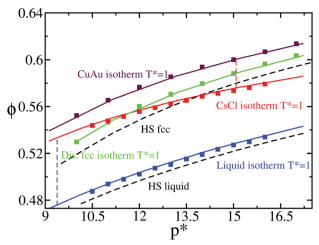


Fig. 2 Equation of state (EOS) of the liquid (blue) and the different crystal phases CsCl (red), CuAu (maroon) and disordered fcc (green) at  $T^* = 1$ . Square symbols represent values from the HS-Y model using Monte Carlo simulations, while lines represent the values of our new continuous potential, PHS-Y. The disordered fcc crystals were generated using MC swap moves at  $T^* = 1.4$ . A quadratic fit of the liquid EOS of our continuous model is given by  $p^* = 99.087 - 455.61\phi_1 + 561.24\phi_1^2$ . Note that this fit is only valid from a liquid volume fraction of  $\phi_1$  = 0.47 to  $\phi_1$  = 0.55. Black dashed curves indicate the equation of state of the liquid and the fcc crystal phase of hard spheres<sup>93</sup> for later comparison. The coexistence pressure of the liquid-CsCl and CsCl-CuAu transitions is represented by blue and red vertical dashed lines, respectively.

We tested the accuracy of the new continuous potential (PHS-Y) against the original Yukawa model (HS-Y)86,87 by evaluating the equation of state (EOS) of the liquid and the different crystalline structures that may compete in nucleating at  $T^* = 1$ . In Fig. 2, we compare Monte Carlo (MC) results of the HS-Y model (squares) with our continuous potential (lines) for the EOS of the liquid (blue), CsCl (red), CuAu (maroon) and substitutionally ion disordered face-centered-cubic (fcc) crystal phase (green). As can be seen, the agreement between the continuous and the discontinuous potential is excellent. Moreover, we computed the liquid and CsCl packing fraction at the liquid-CsCl coexistence pressure at  $T^* = 1$  as well as the CuAu  $\phi$  at the CsCl-CuAU coexistence line for our model (further details are provided in Section IV.A). As shown in Fig. 1 (brown circles), the phase diagram of the PHS-Y potential employed in this work is very similar to that of the HS-Y model (at least at  $T^* = 1$ , although it is expected to work for other  $T^*$ values as well). In Fig. 2, MC data obtained by following the numerical details given in ref. 86 and 87 are shown. For generating the substitutionally disordered fcc crystal configurations used in this work, we employed MC swap moves as detailed in ref. 85 and 87, adjusting the disorder degree of the lattice to the one corresponding to  $T^* = 1.4$ , just above the disordered-to-ordered (CuAu) transition temperature observed from  $T^* \sim 1.3$  to 1.15 depending on  $\phi$ . 89 Interestingly, despite having a similar arrangement of atoms (when c/a = 1), the density of the CuAu crystal phase is significantly higher than that of the disordered fcc phase. This is due to the fact that in the CuAu phase, every colloid is surrounded by other colloids of the opposite charge, so the lattice can reach a higher density without being destabilised by unfavourable interactions of the same sign as in the disordered fcc crystal.

# III. Seeding framework for crystal nucleation

In the last few years, among the different available rare event techniques used to study crystal nucleation, 50-52,54 a simple and computationally efficient approach has emerged, the Seeding technique. 90-92 Even though the success of Seeding in predicting accurate nucleation rates and free energy barriers relies on the validity of Classical Nucleation Theory<sup>67,68</sup> (extensively discussed in ref. 74) and on the assumption that a local order parameter can adequately distinguish between particles belonging to the stable and metastable phases, the popularity of Seeding for nucleation studies is sharply increasing. 102-107 Despite its approximations, Seeding has been shown to be consistent with independent calculations at moderate metastability using techniques like Umbrella sampling50 or Forward flux sampling<sup>51</sup> as well as with non-biased calculations (i.e. brute force simulations) in the deep supercooled regime. 20,28,108 Thus, we employ Seeding to investigate the crystal nucleation scenario among the different crystal polymorphs of oppositely charged colloids. Within the CNT-Seeding framework, the nucleation rate J is defined as 18,68

$$J = \rho_1 Z f^+ \exp(-\Delta G/(k_{\rm B}T)) \tag{3}$$

where  $(\rho_1 Z f^*)$  represents the kinetic pre-factor  $(K_{\rm kinetic})$  and  $\Delta G/(k_{\rm B}T)$  is the free energy barrier of the system to nucleate. In the kinetic pre-factor,  $\rho_1$  is the number density of the fluid (which for our model will be expressed in reduced units,  $\rho_1^*$ ), Z is the Zeldovich factor, which accounts for the curvature of the top of the free energy barrier, and  $f^+$  is the frequency of ion attachment/detachment to the critical cluster. The expression for Z is

$$Z = \sqrt{|\Delta\mu|/(6\pi k_{\rm B} T N_{\rm c})} \tag{4}$$

where  $\Delta\mu$  is the chemical potential difference between the bulk solid and bulk liquid phases under the conditions where the cluster is critical, and  $N_{\rm c}$  is the number of particles in the critical cluster. To compute  $f^+$ , we adopt the expression proposed by Auer and Frenkel:<sup>18</sup>

$$f^{+} = \left\langle \frac{(N(t) - N_{\rm c})^2}{2t} \right\rangle \tag{5}$$

with N(t) being the number of particles of a cluster around the top of the free energy barrier at a given time t. Finally, the free energy barrier term of eqn (3) can be calculated through the following expression:

$$\Delta G = \frac{1}{2} N_{\rm c} |\Delta \mu|. \tag{6}$$

Additionally, by assuming a spherical shape of the critical cluster, the CNT framework also provides an expression for the

interfacial free energy:

$$\gamma = \left(\frac{3N_{\rm c}\rho_{\rm s}^2|\Delta\mu|^3}{32\pi}\right)^{1/3} \tag{7}$$

where  $\rho_s$  is the number density of the solid phase (also given in what follows in reduced units,  $\rho_s^*$ ).

As previously mentioned, the success of Seeding predictions relies on how accurate a local order parameter is when distinguishing the number of particles that belong to the critical cluster. In earlier work,  $^{20}$  it has been demonstrated that when the number of mislabelled particles by the order parameter is the same for the bulk solid phase and for the bulk liquid one, the order parameter can provide reliable estimates of J and, hence, free energy barriers and interfacial free energies.

Here, we use the  $\bar{q}_4$ - $\bar{q}_6$  local order parameters proposed by Lechner and Dellago<sup>109</sup> within the mislabelling scheme<sup>20</sup> to identify the number of particles belonging to clusters of each crystal polymorph. The cut-off distance for both computing the  $\bar{q}_4$  and  $\bar{q}_6$  parameters and identifying the biggest cluster was 1.41 $\sigma$ . In Fig. 3(a), we show a plot of the  $\bar{q}_4$ - $\bar{q}_6$  points for colloidal particles in the bulk liquid (blue) and in the different bulk crystal phases, CsCl (red), CuAu (maroon) and disordered fcc (green), at  $T^* = 1$  and  $p^* = 13$  (liquid packing fraction  $\phi_1$  = 0.514). As can be observed, the  $\bar{q}_6$  parameter successfully discriminates between liquid-like and the different crystal-like particles. Likewise, the  $\bar{q}_4$  parameter distinguishes between the CsCl (body centered cubic) crystal structure and the ion-disordered fcc and CuAu crystal phases. However, the  $\bar{q}_4$ parameter cannot reliably discriminate between CuAu-like particles and disordered fcc-like ones, since the main difference between these two crystal phases is the charge identity of the ions rather than their position within the lattice. A criterion to distinguish between CuAu-like particles and disordered fcc-like ones is described in Section IV.B (Fig. 7). Additionally, in Fig. 3(b), we plot the mislabelling  $\bar{q}_6$  optimal threshold for discriminating between liquid-like and crystal-like colloids as a function of the liquid packing fraction at  $T^* = 1$ . The  $\bar{q}_4$  optimal threshold for identifying CsCl-like particles versus CuAu-like or disordered fcc ones as a function of  $\phi_1$  is shown in the inset of Fig. 3(b). Within the mislabelling scheme, we find that the percentage of mislabelled particles in the bulk phases at the selected  $\bar{q}_6$  and  $\bar{q}_4$  optimal thresholds is below 0.001% and 0.1%, respectively.

# IV. Nucleation scenario of oppositely charged colloids

## A. On the relative stability between the liquid and the different crystal phases

The phase diagram of 50:50 binary mixtures of colloidal electrolytes contains several different crystal phases<sup>87</sup> that may compete in nucleating from the liquid depending on the  $T^*$ - $\phi$  conditions (see Fig. 1). Whereas at moderately high temperatures ( $T^* > 1.15$ ), the liquid coexists with a substitutionally disordered fcc colloidal phase, at low temperatures, the

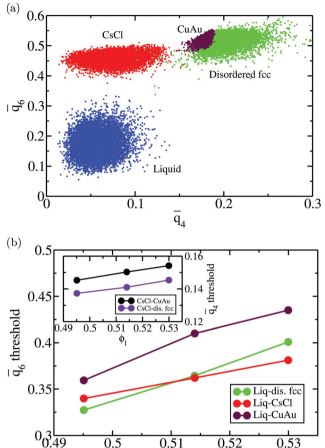


Fig. 3 (a)  $\bar{q}_4 - \bar{q}_6$  plot for 10 000 particles of the bulk liquid (blue) and the different crystal bulk phases, CsCl (red), CuAu (maroon) and disordered fcc (green), at  $T^* = 1$  and  $p^* = 13$ , which corresponds to a liquid packing fraction of  $\phi_1$  = 0.514. (b) Dependence of the  $\bar{q}_6$  optimal threshold within the mislabelling criterion between the liquid and the different crystalline structures as a function of the liquid packing fraction at  $T^* = 1$ . The liquid and solid phases studied are compared at the same pressure, which corresponds to that of the liquid at the liquid packing fraction indicated in the figure. The inset shows the  $ar{q}_4$  optimal threshold for distinguishing between the CsCl crystal phase (body centered cubic) and the CuAu and disordered fcc phases as a function of  $\phi_l$  at  $T^* = 1$ .

stable solid coexisting with the liquid is the CsCl crystal phase. Additionally, at a low temperature and a higher packing fraction, the CuAu crystal becomes more stable than the CsCl phase. However, the stability limit of the CuAu phase only extends up to a reasonably high packing fraction, where upon reaching that limit, the most stable phase becomes the tetragonal crystal phase<sup>87,89</sup> (see Fig. 1).

Since we shall focus on the nucleation scenario at  $T^* = 1$ , we start determining the relative stability between the liquid and the different crystal phases that may nucleate at  $T^* = 1$  using direct coexistence (DC) simulations. 94,110 By placing in the same simulation box a liquid slab of  $\sim 3000$  colloidal particles in contact with a slab of  $\sim 3000$  colloidal particles of each crystal phase, we evaluate the coexistence pressure at which the crystals neither grow nor melt. The coexistence pressures at  $T^* = 1$  for each solid phase with the liquid are 9.38(3), 11.19(3) and 11.62(3) (in reduced units) for the CsCl, CuAu and disordered fcc phases, respectively.

Furthermore, we perform thermodynamic integration<sup>111</sup> to (1) evaluate the chemical potential difference ( $\Delta \mu$ ) between the liquid and each crystal phase along  $\phi_1$  and (2) study the relative stability among the different crystal phases. In Fig. 4, we plot  $\Delta \mu$  as a function of the liquid packing fraction for the three different crystal phases. As can be seen, at a low packing fraction, the CsCl crystal (red) is the most stable phase, while at a moderate packing fraction,  $\phi_1 > 0.53$ , the CuAu crystal (maroon) becomes the most stable phase. We also observe that, independently of  $\phi_1$ , the disordered fcc phase (green) always remains metastable, with respect to the CsCl phase (at a low packing fraction) or to the CuAu crystal (at any  $\phi_1$ ). However, at  $\phi_1 \sim 0.54$ , the chemical potentials of the CsCl and disordered fcc phases become similar.

#### Crystal nucleation rate of the different polymorphs

We exploit the Seeding suitability to investigate the different crystalline structures emerging from the metastable liquid 112 by performing Seeding simulations of the CsCl, CuAu and disordered fcc solid phases. In Table 1, we report the results for the number of particles in the critical cluster  $(N_c)$ , the pressure,  $p_c^*$ , and the packing fraction,  $\phi_l$ , at which the cluster was found critical, as well as the density of the liquid and solid phases  $(\rho_1^*$  and  $\rho_s^*$ ), the chemical potential difference between the solid and the liquid  $(\Delta \mu)$ , the liquid-crystal interfacial free energy  $(\gamma)$ , the free energy barrier height  $(\Delta G)$ , the attachment

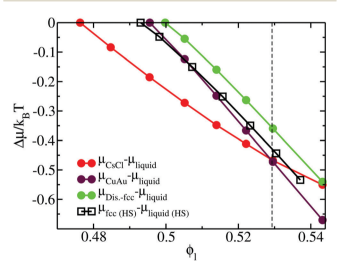


Fig. 4 Chemical potential difference (per particle) ( $\mu_{\rm solid} - \mu_{\rm liquid}$ ) between the different colloidal crystal phases and the liquid (filled circles) as a function of the liquid packing fraction at  $T^* = 1$ . The chemical potential of the solid is evaluated at the same pressure that has to be applied in the liquid to obtain  $\phi_{\mathrm{l}}$ . Results of the chemical potential difference between the fcc crystal phase and the bulk liquid phase (empty squares) of purely repulsive hard-sphere colloids<sup>93</sup> as a function of  $\phi_1$  are also included. The black vertical dashed line indicates the packing fraction at which the CsCl-CuAu transition takes place in the HS-Y model from ref. 72 and 87.

Table 1 Results of the Seeding technique at  $T^* = 1$  for the CsCl and disordered fcc crystal phases. See the main text for the meaning of each variable

| $p_{\mathrm{c}}^{*}$ | $\phi_1$ | $N_{ m c}$ | $ ho_1^*$ | $ ho_{ m s}^*$ | $\Delta\mu/k_{\rm B}T$ | $\gamma/\left(k_{\rm B}T/\sigma^2\right)$ | $\Delta G/k_{\rm B}T$ | $f^{\dagger}/(6D_{\rm L}/\sigma^2)$ | Z      | $\log_{10} (J/(6D_{\rm L}/\sigma^5))$ |
|----------------------|----------|------------|-----------|----------------|------------------------|---|-----------------------|-------------------------------------|--------|---------------------------------------|
| CsCl                 |          |            |           |                |                        |   |                       |                                     |        |                                       |
| 12.931               | 0.5119   | 212        | 0.9777    | 1.0760         | -0.331                 | 0.643                                     | 35.07                 | 525.5                               | 0.0091 | -14.56                                |
| 12.564               | 0.5086   | 276        | 0.9715    | 1.0711         | -0.304                 | 0.642                                     | 41.93                 | 1143.5                              | 0.0076 | -17.28                                |
| 12.022               | 0.5036   | 399        | 0.9619    | 1.0634         | -0.261                 | 0.621                                     | 52.10                 | 1084.0                              | 0.0059 | -21.84                                |
| 11.350               | 0.4971   | 801        | 0.9495    | 1.0533         | -0.203                 | 0.606                                     | 81.35                 | 1164.5                              | 0.0037 | -34.72                                |
| Disordered fcc       |          |            |           |                |                        |   |                       |                                     |        |                                       |
| 15.140               | 0.5295   | 193        | 1.0114    | 1.1264         | -0.357                 | 0.692                                     | 34.43                 | 2537.0                              | 0.0099 | -13.55                                |
| 14.700               | 0.5263   | 334        | 1.0052    | 1.1195         | -0.315                 | 0.731                                     | 52.61                 | 4128.0                              | 0.0071 | -21.38                                |
| 14.260               | 0.5229   | 460        | 0.9988    | 1.1121         | -0.272                 | 0.699                                     | 62.59                 | 4935.5                              | 0.0056 | -25.74                                |
| 13.904               | 0.5201   | 681        | 0.9934    | 1.1057         | -0.237                 | 0.691                                     | 80.59                 | 3082.5                              | 0.0043 | -33.88                                |
|                      |          |            |           |                |                        |   |                       |                                     |        |                                       |

rate  $(f^+)$ , the Zeldovich factor (Z) and the decimal logarithm of the nucleation rate  $(\log_{10} I)$  at such conditions.

Even though crystalline clusters of the three polymorphs were inserted, we were only able to measure the nucleation rate for the CsCl and disordered fcc phases. CuAu crystal clusters rapidly show a martensitic transformation 113 into the stable CsCl phase at any packing fraction below  $\phi_1$  < 0.528, and beyond that, the fluid spontaneously crystallizes (so that Seeding is not needed). The mechanism by which the CuAu nuclei transform into CsCl ones in the stability region of the CsCl crystal (below a packing fraction of 0.528) is a diffusionless phenomenon (i.e., particle diffusion is not necessary to produce it) in which the solid interconversion (martensitic transformation) occurs via deformation of the unit cell. In this case, the tetragonal unit cell of the CuAu crystal nuclei transforms into the cubic unit cell of the CsCl crystal. This is possible because the cubic CsCl crystal (c = a) can also be considered as a tetragonal CuAu-like structure with  $c/a = 1/\sqrt{2}$  (see Fig. 1 for more details on these two unit cells).89

We show the nucleation scenario between the CsCl and the disordered fcc crystal phases at  $T^* = 1$  by plotting in Fig. 5 the nucleation rate of both polymorphs as a function of  $\phi_1$ . Our results clearly evidence that, from low to moderate packing fraction ( $\phi_1 < 0.53$ ), the CsCl structure possesses the fastest nucleation rate (red curve). At a higher packing fraction  $(\phi_1 > 0.535)$ , the disordered fcc I (green curve) becomes equivalently competitive with respect to the CsCl rate. Nonetheless, for  $\phi_1 > 0.517$ , we observe post-critical crystal growth of the disordered fcc phase from the CsCl critical nuclei, termed parasitic crystallization (red dashed dotted vertical line). We also show an estimation of the nucleation rate for the CuAu structure in Fig. 5 (dashed dotted maroon curve). By assuming that  $\gamma$  does not depend sensibly on the crystal structure (as will be shown in Section IV.C) and using the previously calculated values of  $\Delta\mu$  and  $\rho_s^*$  for the CuAu crystal, we calculate  $N_c$  as a function of  $\phi_1$  via eqn (7). Then, using eqn (8), we evaluate  $\Delta G$ , and assuming that the attachment rate must be of the order of magnitude of the CsCl one, we estimate J (using eqn (3))

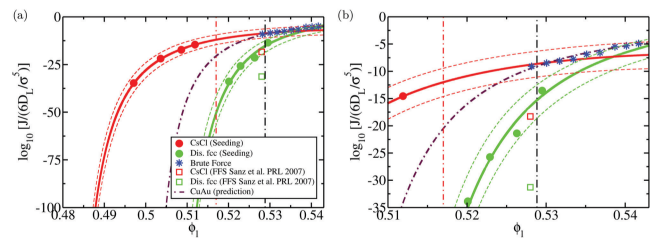


Fig. 5 (a) Logarithm of the nucleation rate as a function of the liquid packing fraction for the CsCl (red) and disordered fcc (green) crystal phases. Circles indicate the Seeding estimates of J while continuous lines are the CNT-like fits<sup>20</sup> to the Seeding results. Dashed curves indicate the upper and lower bounds of uncertainty of the Seeding results. Empty squares represent the nucleation rate obtained via Forward-Flux Sampling calculations from ref. 72 for the CsCl crystal phase (red) and for a disordered fcc phase (green). Blue asterisks account for the nucleation rates obtained using brute force calculations. The dashed dotted maroon curve shows a prediction of the nucleation rate for the CuAu crystal phase assuming that its y is the same as those of the CsCl and disordered fcc phases. The vertical black dashed dotted line indicates the coexistence line between the CsCl and CuAu crystal phases according to the crossover in  $\Delta\mu$  shown in Fig. 4, while the red dashed dotted line represents the lowest  $\phi_1$  at which parasitic crystallization from the CsCl phase is observed. (b) Inset of (a) to better visualise the spontaneous nucleation regime.

for the CuAu crystal phase. As expected, the nucleation rate of the CuAu structure only becomes competitive with respect to that of the CsCl phase at pressures where  $\phi_1 > 0.53$  (*i.e.* at a moderately high packing fraction), in the stability region of the CuAu crystal. Since  $\gamma$  for CuAu is assumed to be the same as those for the CsCl and disordered fcc crystals, the key factors in determining the nucleation rate are the solid density and the chemical potential difference  $\Delta\mu$ . Note that the predicted CuAu J can also be regarded as the approximate upper bound limit of very small disordered fcc crystals (with a positional ion-disorder degree lower than that at  $T^* = 1.4$ ).

To further check which crystal phase nucleates the fastest at a moderately high  $\phi_1$  ( $p^*=15$ ), we insert "frozen" clusters (where the particles belonging to the crystal structure are kept rigid to avoid the martensitic transformation from CuAu to CsCl) of the three different polymorphs into the liquid phase. By inserting clusters of N=30, we observe that the average time (of five independent trajectories) that it takes for each solid cluster to irreversibly grow under such conditions is 25, 40 and  $2200\sigma^2/6D_{\rm L}$  for the CsCl, CuAu and disordered fcc phases, respectively. Although these results cannot be directly related to the nucleation rate, they support the predicted scenario of J shown in Fig. 5.

Moreover, to validate our Seeding estimations of the nucleation rate, we perform brute force calculations at a high packing fraction (from  $\phi_1 = 0.528$  to  $\phi_1 = 0.542$ , see Table 2). As can be seen in Fig. 5, the CsCl curve of J converges at  $\phi_1 > 0.528$ with the brute force nucleation rates (blue asterisks), whereas the disordered fcc values of J coincide with the brute force nucleation rates at  $\phi_1 > 0.538$ . We obtain the structure of the crystalline clusters near the top of the free energy barrier (just before irreversible crystal growth) by comparing their radial distribution function, g(r), to those of spherical clusters of similar size cut from the different bulk crystal polymorphs under such conditions (only the particles belonging to the cluster are considered when computing g(r), thus their convergence to zero occurs at long distances). Our simulations reveal that at  $p^* = 15$ , the structure of the crystalline clusters is predominantly CsCl-like (see snapshots a, b and c of Fig. 6). The radial distribution function of these clusters is plotted in

**Table 2** Results of the logarithm of the nucleation rate using brute force simulations in the NpT ensemble. The total number of colloidal particles,  $N_T$ , the average volume of the metastable liquid before freezing occurs,  $\langle V \rangle$ , and the average time to observe nucleation  $\langle t \rangle$  are given. Five independent trajectories were performed at each pressure

| p*    | $N_{ m T}$ | $\langle V \rangle / \sigma^3$ | $\langle t \rangle / \! \left( \sigma^2 / 6 D_{ m L}  ight)$ | $\log_{10} (J/(6D_{\rm L}/\sigma^5))$ |
|-------|------------|--------------------------------|--|---------------------------------------|
| 17.25 | 1000       | 964                            | 19.9   | -4.28                                 |
| 17    | 2000       | 1932                           | 41.4   | -4.90                                 |
| 16.75 | 2000       | 1938                           | 109.0  | -5.32                                 |
| 16.5  | 2000       | 1945                           | 162.6  | -5.50                                 |
| 16.25 | 54000      | 52 670                         | 73.0   | -6.58                                 |
| 16    | 54000      | 52 806                         | 140.4  | -6.87                                 |
| 15.75 | 54000      | 52941                          | 1123   | -7 <b>.</b> 77                        |
| 15.5  | 54000      | 53 177                         | 2401   | -8.11                                 |
| 15.25 | 54000      | 53 347                         | 5461   | -8.46                                 |
| 15    | 54000      | 53 530                         | 22 459   | -9.08                                 |

Fig. 6(j) against those of the CsCl, CuAu and disordered fcc spherical clusters of 50 colloidal particles cut from equilibrated bulk configurations. As indicated by black arrows, the g(r)functions of the different brute force clusters exhibit the characteristic peaks of a CsCl-like radial distribution function. Nevertheless, at  $p^* = 16.75$ , the structure of the clusters turns out to be more disordered, although, given their small size, it is difficult to discern. In the inset of Fig. 6j, we show how their g(r)functions compare to those of spherical clusters of 15 colloids for each polymorph cut from bulk simulations at  $p^* = 16.75$ . Obviously, it is difficult (and maybe useless) to assign a solid structure to a cluster having only 15 particles. We additionally determine the nature of the resulting crystallized systems from brute force simulations by comparing their radial distribution function with those of the different bulk polymorphs. As shown in Fig. 6k, the resulting crystals (blue thin curves) predominantly exhibit fcc coordination with ion charge disorder as depicted in the snapshots shown in (d) and (h) for  $p^* = 15$  and  $p^* = 16.75$ , respectively.

We also compare the potential energy U (configurational contribution to the internal energy) of the different crystallized systems with those of the different bulk phases as a function of pressure (Fig. 6i). As can be seen, the obtained crystals from brute force simulations show slightly higher values of U, but of the same order, than that of the disordered fcc bulk phase under such conditions. The higher values of U can be explained by the presence of lattice defects caused by fast crystal growth, defects near the boundary conditions due to a mismatch between the crystal orientation and the simulation box, or just by a higher degree of ion disorder in the crystal lattice. For similar reasons, the potential energy of a crystallized CsCl-seeded trajectory into a CsCl crystal (according to the  $\bar{q}_4$  local order parameter, as shown in Fig. 7) also exceeds the bulk CsCl U under the same conditions (empty red square in Fig. 6i).

In Fig. 5, we also include the values of J evaluated via Forward flux sampling (FFS) calculations at  $p^* = 15$  for the CsCl crystal phase (red square) and a disordered fcc structure (green square) reported in ref. 72. Our Seeding results predict a nucleation rate of both phases about 10 orders of magnitude higher than those reported in ref. 72. In contrast, our Seeding results for the CsCl phase are in very good agreement with the brute force calculations under such conditions (and also up to a higher packing fraction). Since brute force results contain no approximation, and they can be regarded as a benchmark for methodologies aiming to estimate J for nucleation, these results support the validity of CNT in describing the nucleation behaviour of colloidal electrolytes (at least at  $T^* = 1$ ). The discrepancy between the Seeding and FFS nucleation rates is somewhat not surprising since, on the one hand, it has been previously found that FFS calculations may underestimate the nucleation rate by a few orders of magnitude when the sampling is not extremely large, 26,44,114,115 and on the other, because Seeding results may also have an uncertainty (systematic + statistical) of about 3-5 orders of magnitude in the nucleation rate (see Fig. 5). However, our estimations of J qualitatively agree with those from ref. 72 in arguing that the

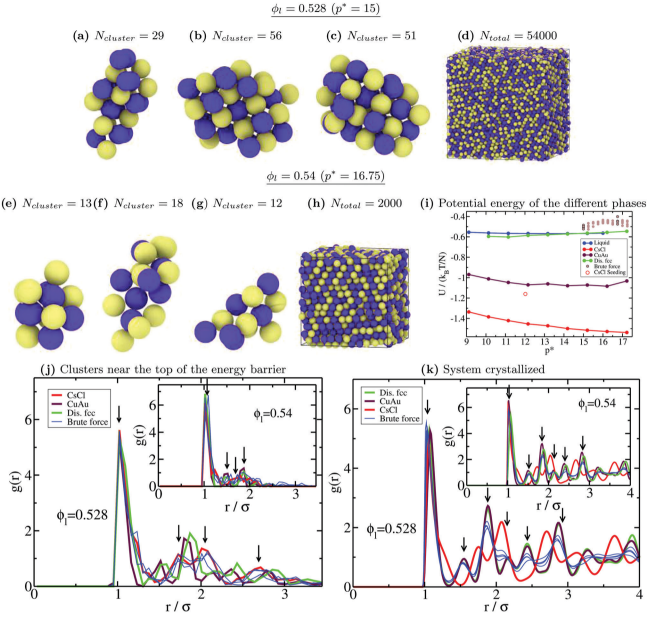


Fig. 6 (a)-(c) Snapshots of three different crystalline clusters from independent trajectories identified just before irreversible crystal growth in brute force calculations at  $\phi_l = 0.528$  and  $T^* = 1$ . Yellow particles indicate positively charged colloids and blue ones indicate negatively charged colloids.  $N_{\text{cluster}}$  indicates the total number of colloidal particles determined by the  $\bar{q}_6$  local order parameter within the mislabelling criterion. (d) Snapshot of the crystallized system (disordered fcc) of a brute force trajectory at  $\phi_1$  = 0.528. Similar results in all the different trajectories were found. (e)–(g) Snapshots of three different crystalline clusters from independent runs identified just before irreversible crystal growth in brute force simulations at  $\phi_1 = 0.54$  and  $T^* = 1$ . (h) Snapshot of the crystallized system (disordered fcc) of a brute force simulation at  $\phi_1 = 0.54$ . (i) Potential energy U of the bulk liquid (blue), and CsCl (red), CuAu (maroon) and disordered fcc (green) crystal phases as a function of pressure. Brown circles indicate the potential energy of the different brute force trajectories upon crystallization (thick contours indicate the two pressures considered in this figure). The red circle represents U for the crystallized system of a Seeding trajectory with a CsCl seed at  $\phi_{\rm l}$  = 0.512 (see the snapshot in Fig. 7). (j) All-particle radial distribution function of spherical 50-colloid clusters cut from bulk configurations of the different crystal polymorphs at  $\phi_1 = 0.528$  and  $T^* = 1$  (inset: 15-colloid clusters cut from bulk crystal configurations at  $\phi_1 = 0.54$  and  $T^* = 1$ ). Blue thin curves represent the radial distribution function of the clusters shown above. Black down arrows indicate the most representative peaks shown by the cluster q(r) functions. (k) Allparticle radial distribution function of the different bulk crystal structures at  $\phi_1 = 0.528$  and  $T^* = 1$  (inset:  $\phi_1 = 0.54$  and  $T^* = 1$ ) along with those obtained for the crystallized system (blue curves) resulting from several independent trajectories of brute force simulations under those conditions.

nucleation rate of CsCl is sensibly faster than that of the disordered fcc phase at this packing fraction.

The Seeding uncertainty in the nucleation rate (and also in  $\Delta G$  and  $\gamma$ ) mainly comes from (1) the systematic error in the calculation of the number of particles in the critical cluster  $(N_c)$ 

and (2) the statistical uncertainty in the determination of the conditions under which the cluster becomes critical. It is difficult to accurately evaluate the impact of the systematic error in  $N_c$  since different (reasonable) order parameters may provide moderately different values of the critical cluster size.

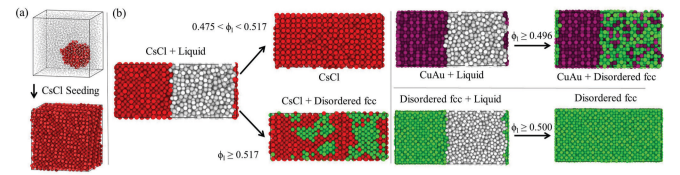


Fig. 7 (a) Initial and final configurations of a CsCl-seed Seeding trajectory at  $\phi_1 = 0.512$  and  $T^* = 1$ . CsCl-like particles are labelled in red and liquid-like particles are labelled in grey according to the  $\bar{q}_6$  mislabelling criterion. To provide better visualisation of the nucleus, liquid-like colloids are resized to a lower  $\sigma$ . The internal energy of the final configuration is given in Fig. 6(i) (empty red circle). (b) Direct coexistence simulations showing the crystal growth behaviour of the different polymorphs at different packing fractions at  $T^* = 1$ . CsCl-like particles are labelled in red, CuAu-like ones are labelled in maroon, disordered fcc-like ones are labelled in green and liquid-like ones are labelled in grey. To discriminate between liquid-like and crystal-like particles, we use the  $\bar{q}_6$  local order parameter, while for differentiating between CsCl-like (bcc) particles and either CuAu or disordered fcc-like ones, we use the  $\bar{q}_4$  parameter, as shown in Fig. 3. To distinguish between CuAu (ordered) and disordered fcc crystal-like particles, we employ a criterion based on labelling CuAu-like particles those that have at least 12 CuAu-like neighbours, considering as a CuAu neighbour a particle that possesses at least 8 or more neighbours of the opposite charge at a cut-off distance of  $1.25\sigma$  (the first minimum of an all-particle radial distribution function of a bulk fcc crystal including the first two coordination shells).

For that reason, we consider that our estimates of  $N_c$  might be affected by an uncertainty of  $\pm N_c^{2/3}$  (the outer layer particles in the cluster, which is where different order parameters may provide different estimations). Regarding the statistical uncertainty in the determination of the critical pressure of each cluster (trajectory stochasticity + the use of a given grid of simulated pressures), we ascribe to the associated error of the critical pressure  $\pm 1/2$  of the pressure grid employed to determine the critical pressure of each cluster (we typically use ten independent trajectories every  $0.01p^*$ ). Other minor sources of error, such as the uncertainty in the chemical potential difference between the liquid and the crystal phases and their density dependence on pressure, have also been considered, although they contribute very modestly. By taking into account these different sources of error, we have evaluated the upper and lower bounds of the nucleation rate uncertainty, as shown in Fig. 5 (the uncertainty in  $\Delta G$  and  $\gamma$  will be shown in the following section).

We observe in our Seeding simulations that independent of the nature of the inserted crystalline seeds, the post-critical cluster growth is highly enriched in disordered fcc crystal domains at a moderately high packing fraction. To quantitatively characterise this parasitic crystal growth behaviour of a metastable phase upon the formation of the critical nucleus, we perform direct coexistence simulations of a liquid slab in coexistence with each different crystalline structure from low to high packing fraction.

As depicted in Fig. 7, CsCl crystals (the spherical cluster in (a) or the solid slab with a flat interface in (b) left panel) only grow in equilibrium up to packing fractions of  $\phi_1$  < 0.517, well below their stability limit at  $\phi_1$  = 0.528. Above  $\phi_1$  > 0.517, the crystal growth occurs so fast that it impedes the correct positional order of the colloids into the stable CsCl lattice, and thus, stacking mixtures of CsCl and disordered fcc layers arise. Conversely, CuAu and disordered fcc crystals (right panel)

exhibit positional-charge disorder growth either at a low or high packing fraction. These results clearly indicate that parasitic growth of the metastable disordered fcc phase (which likely contains stacking faults of hexagonal-close-packing due to the high resemblance in free energy that these two crystal lattices present for hard-spheres116) occurs regardless of the nature of the stable crystalline phase, at moderate  $\phi_1$  from the CsCl crystal, and at high packing fraction from the CuAu phase. This behaviour may explain why in ref. 87 CuAu crystals in the presence of disordered close-packed ones were found within 18 days after the sample preparation of an oppositely charged colloidal suspension at  $\phi_1 = 0.58$ . According to our simulations, positionally disordered close packing crystal domains (most likely containing stacking faults) may have quickly grown from the critical nuclei along with stable CuAu domains, or even the CuAu domains may have appeared later after recrystallization from the disordered close packing domains.

We also evaluate the crystal growth rate (u) as a function of  $\phi_1$  for the different polymorphs using direct coexistence simulations (Fig. 8A). To measure u, we monitor the position of the crystal front as a function of time for each interface during the DC simulation. We find that the growth rate of the disordered fcc phase is between 4 to 10 times faster than that of CsCl depending on the considered  $\phi_l$ . We also observe that the disordered fcc growth rate may depend on the initial crystal phase in contact with the liquid (at least when measured at short distances from the precursor crystal phase as in our simulations). When the disordered fcc phase grows from the CsCl crystal (bcc lattice), it requires more time to grow the same distance than when starting from its own lattice or the CuAu phase (fcc lattice). This is likely happening (1) because of the initial time spent by the system to accommodate the fcc lattice (and its density) from the precursor bcc lattice, (2) because the disordered fcc phase may grow under stress conditions due to the mismatch between its crystal lattice and the simulation box

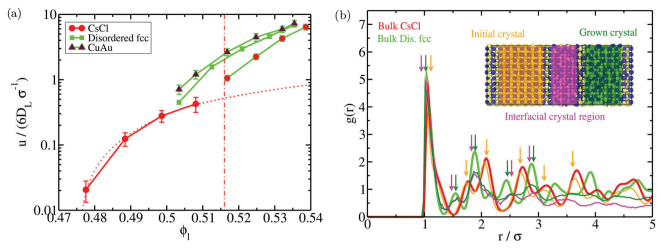


Fig. 8 (a) Crystal growth rate (u) as a function of the liquid packing fraction computed by means of direct coexistence simulations for different initial crystalline slabs in contact with the liquid. Red circles indicate the obtained growth rates when the initial crystal phase was CsCl, maroon diamonds indicate the case when the initial crystal phase was CuAu, and green squares indicate the case when the starting crystal phase was disordered fcc. The color of the continuous lines connecting the different symbols indicates whether the emerging crystal phase is CsCl (red) or disordered fcc (green). The red dotted line shows a linear fit to the CsCl growth rates extrapolated into the parasitic crystallization regime for visual comparison with the disordered fcc u. The vertical red dashed dotted line indicates the lower  $\phi_1$  at which parasitic crystallization from CsCl is observed. (b) All-particle radial distribution function (g(r)) of the initial CsCl crystal (orange), the grown crystal (dark green), and the interfacial region from which the emerging crystal grew (pink) of a direct coexistence simulation at  $\phi_1 = 0.518$  and  $T^* = 1$ . Bulk CsCl (red) and disordered fcc (light green) all-particle radial distribution functions at  $\phi_1 = 0.518$ and  $T^* = 1$  are also included for comparison. Note that the q(r) functions of the different crystal regions of the DC simulation converge to zero at long distances since the radial distribution functions have only been evaluated for those particles of the considered region. Bulk CsCl and disordered fcc g(r) functions converge to 1 at long distances. Different coloured down arrows indicate the most representative peaks of each q(r). A snapshot of the different considered regions of the direct coexistence simulation for which the radial distribution function has been computed is shown.

size that matches the precursor crystal lattice, and (3) because the growing crystal orientation that best matches the precursor CsCl lattice could have a lower growth rate than those emerging from the disordered fcc or CuAu crystals. Nonetheless, even under these unfavourable circumstances, the disordered fcc crystal manages to grow much faster from the CsCl crystal than the CsCl itself. To further confirm the nature of the emerging crystals without the use of a local order parameter (in Fig. 7, that was determined using the  $\bar{q}_4$  and  $\bar{q}_6$  parameters), we compute the all-particle radial distribution function of the initial CsCl slab (orange curve), the grown crystal (dark green), and the interfacial region connecting the initial crystal and the emerged one (pink) of a DC trajectory at  $\phi = 0.518$  and  $T^* = 1$ (Fig. 8B). When comparing these three radial distribution functions to those of the bulk solid CsCl and disordered fcc phases under the same conditions (red and light green curves, respectively), we find that the structure of the new grown solid and the interfacial crystal mostly corresponds to a disordered fcc phase (although it probably contains a certain degree of hexagonal-close packing and CsCl stacking faults), as predicted by the order parameter in Fig. 7. We expect that the degree of CsCl-like vs disordered fcc-like particles of the emerging crystal will depend on  $\phi_1$ , being higher for fcc-like particles at a higher packing fraction. These results evidence how parasitic crystallization is not an artifact of the employed order parameter.

Even though parasitic crystallization entails the nucleation of one phase and the subsequent post-critical crystal growth of a different phase from the critical cluster, it should not be regarded as the well-known Ostwald step rule 117 proposed in

1897. In the Ostwald step rule, the crystal phase that nucleates from the melt need not be the one that is thermodynamically most stable, but the one that is closest in free energy to the fluid phase. Next, from the nucleated phase, the thermodynamically most stable phase is expected to form. 118 This rule was later reexamined by Stranski and Totomanow, 119 who argued that the nucleating phase will be the phase with the lowest freeenergy barrier of formation, rather than the phase that is globally most stable under the prevailing conditions. However, what we term as parasitic crystallization here is a phenomenon where, even when the nucleating phase is the thermodynamically most stable one, the post-critical crystal growth occurs through an out-of-equilibrium process by which a different (metastable) phase from the nucleating one and/or the most stable one grows from the critical nucleus.

#### C. Key factors behind the polymorphic crystal competition

To elucidate the thermodynamic and kinetic factors leading to the crystal nucleation scenario shown in the previous section, we analyse the magnitudes that play a role in the polymorphic competition among the different crystal phases. To that end, we firstly focus on the kinetic pre-factor,  $\rho_1 Z f^+$ , of the CNT expression for the nucleation rate (eqn (3)). We compute the kinetic pre-factors of the CsCl and disordered fcc crystals (both polymorphs for which Seeding calculations were successfully performed) by multiplying  $\rho_1 Z f^+$  from the given values in Table 1. We find that the kinetic term of the disordered fcc clusters is about 5 times higher than that of CsCl at any  $\phi_1$  from 0.50 to 0.53. Considering that J varies by tens of orders of

magnitude in small  $\phi_1$  intervals, we conclude that the kinetic

pre-factor cannot play a crucial role in ultimately determining which phase nucleates the fastest.

Hence, the significant differences in the nucleation rate between the two polymorphs must come from the free energy barrier. In Fig. 9, we show the free energy barrier,  $\Delta G$ , for each solid as a function of the liquid packing fraction. Up to  $\phi_1 = 0.53$ ,  $\Delta G$  is sensibly lower for CsCl than those measured for disordered fcc clusters, thus explaining the observed behaviour in the nucleation rate (Fig. 5). Only at a high packing fraction,  $\phi_1 > 0.535$ , the free energy barrier of both crystals become comparable. Our results point to the fact that, at  $\phi_1 > 0.54$ , disordered fcc clusters will nucleate faster than those of CsCl. In contrast to the nucleation rate, for the free energy barriers, we find excellent agreement between our Seeding calculations and the umbrella sampling results reported in ref. 72 for the CsCl and disordered fcc  $\Delta G$  (red and green empty squares in Fig. 9, respectively).

Provided that the differences in I between the two polymorphs come from the free energy barrier, we now focus on the thermodynamic factors that are behind such a difference. According to the CNT expression, the free energy barrier of a spherical cluster is

$$\Delta G = \frac{16\pi\gamma^3}{3\rho_*^{*2}|\Delta\mu|^2}.$$
 (8)

Therefore, the chemical potential difference between the solid and bulk liquid phases  $(\Delta \mu)$ , the density of the crystal phase  $(\rho_s^*)$  and the liquid-crystal interfacial free energy  $(\gamma)$  are the factors that modulate the height of the free energy barrier and, thus, the nucleation rate. Since in Fig. 2 and 4 we have already evaluated the density of the solid phases  $(\rho_s^* = 6\phi_s/\pi)$ 

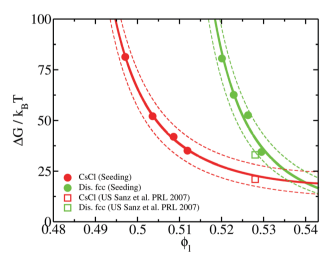


Fig. 9 Free energy barriers obtained from our Seeding calculations as a function of liquid packing fraction for CsCl (red circles) and disordered fcc clusters (green circles). Continuous lines represent CNT-like fits (as shown in ref. 20) to our Seeding results, while dashed curves account for their upper and lower uncertainty bounds. Umbrella sampling free-energy calculations from ref. 72 for CsCl (red square) and disordered fcc (green square) crystals are also included

and its  $\Delta \mu$  dependence on the liquid packing fraction, respectively, we now plot the interfacial free energy of the CsCl and disordered fcc phases as a function of  $\phi_1$  (Fig. 10).

Our results reveal that the interfacial free energy of both solids increases with  $\phi_1$ . Assuming a linear dependence of  $\gamma$  versus  $\phi_1$ , as our data suggest, and as has been successfully proposed for hard-sphere colloids,  $^{20,97}$  we find that  $\gamma(\phi_1)$  for both crystals is roughly similar. Therefore, we cannot ascribe the differences in the free energy barrier to significant variations in the liquid-crystal interfacial free energy among these two polymorphs. Based on these results, we hypothesize our prediction for the nucleation rate of the CuAu crystal phase given in Fig. 5. Moreover, our simulations indicate that the increasing similarity of  $\Delta G$  between the CsCl and disordered fcc clusters at high  $\phi_1$  comes from  $\Delta \mu$  and/or  $\rho_s^*$ .

As shown in Fig. 2 and 4, at a low packing fraction, the density of the disordered fcc and CsCl phases is comparable, while  $\Delta\mu$  is sensibly lower for CsCl. This contributes to the substantial decrease of the free energy barrier of CsCl clusters with respect to those of disordered fcc, as shown in Fig. 9. Nonetheless, at a high packing fraction,  $\phi_1 > 0.535$ ,  $\Delta \mu$  for both polymorphs becomes alike, whereas the density of the disordered fcc phase becomes larger than that of CsCl. This behaviour leads to the observed cross-over of the disordered fcc clusters in  $\Delta G$  and J at high  $\phi_1$ .

#### D. Hard-spheres vs. oppositely charged colloids: a comparison between their nucleation scenarios

In this section, we focus on how the crystal nucleation landscape of purely repulsive HS compares with that of colloidal electrolytes. To that end, we examine the main factors that determine the nucleation rate in both colloidal systems. In the first place, we focus on the kinetic term of the nucleation rate of HS and colloidal electrolytes. Consistent with our previous observations for CsCl and disordered fcc clusters, the kinetic

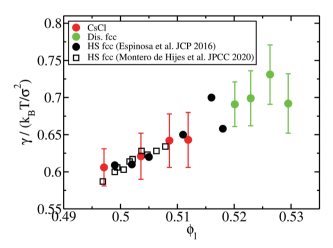


Fig. 10 Liquid-solid interfacial free energy as a function of the liquid packing fraction for CsCl (red circles) and disordered fcc crystals (green ones). Black filled circles and empty squares depict the interfacial free energy as a function of  $\phi_1$  evaluated in ref. 20 and 97, respectively, for fcc clusters of purely repulsive hard-spheres.

pre-factor of HS fcc clusters (see Table 3 of ref. 20) is of the same order of magnitude as those reported in Table 1 for both charged colloidal polymorphs. Hence, in what follows, we analyse the different factors  $(\gamma, \Delta \mu \text{ and } \rho_s^*)$  that determine the height of the free energy barrier.

In Fig. 10, we compare the interfacial free energy of HS fcc clusters<sup>20,97</sup> as a function of  $\phi_1$  (black symbols) with our results of  $\gamma$  for CsCl and disordered fcc clusters. For the three systems, we find very similar values of  $\gamma$  versus  $\phi_1$ . This is not surprising given the resemblance between the liquid and solid structures and the packing fraction in hard-spheres and oppositely charged colloids. On the other hand, we also analyse in Fig. 4 the chemical potential difference between the bulk liquid and the fcc crystal of HS as a function of  $\phi_1$ . Our results show that  $\Delta\mu$  for HS is almost identical to that of the CuAu crystal with the liquid in charged colloids. Additionally, we observe that  $\Delta\mu$  for HS is moderately lower than that of the disordered fcc phase, where unfavourable interactions among colloids of the same sign partially destabilize the fcc lattice. Finally, regarding the density of the HS fcc crystal phase, we observe in Fig. 2 that its packing fraction is slightly lower than that of the disordered fcc phase as a function of  $p^*$ . However, as the liquid  $\phi_1$  of HS is also lower than that of charged colloids as a function of  $p^*$ , when comparing both solid densities as a function of  $\phi_1$ , we find that HS fcc  $\rho_s^*(\phi_1)$  is roughly similar to that of the disordered fcc crystal.

Having discussed the relative differences among the factors behind the nucleation rate of HS and colloidal electrolytes, we compare J for the CsCl phase, predicted CuAu and disordered fcc crystals along with previous computational 18,20,21,120 and experimental nucleation studies<sup>8-10</sup> of HS. As shown in Fig. 11, the nucleation rate of hard-spheres (black curve) lies close to the CuAu prediction of I (maroon dashed line) and above the disordered fcc nucleation rate (green curve). In contrast, we observe that the nucleation rate of the CsCl crystal phase (red curve) is significantly higher than that of HS, especially at low  $\phi_1$ . This is somehow expected since (1)  $\gamma$  is very similar for all the systems and (2)  $\Delta\mu$  is roughly the same for fcc HS and for the CuAu phase, moderately lower than that of the iondisordered fcc phase, and sensibly higher than that of CsCl.

Simultaneously, in Fig. 11, considerable disagreement can be observed between the experimental (black triangles) and computational (black curve and empty symbols) homogeneous nucleation rates for HS at a moderate packing fraction  $(\phi_1 \sim 0.515)$ . 11,58,60,61,66 Several different possible explanations for the discrepancy between experiments and simulations have already been postulated, such as heterogeneous nucleation, sedimentation, incomplete shear melting or hydrodynamic effects. 11,60,61,65,66 Nonetheless, quite surprisingly, we observe that the nucleation rate curve of the CsCl crystal phase closely follows the experimental trend of J for HS. In this respect, our results may suggest that an alternative crystalline polymorph sustained by weak electrostatic attractive interactions among (the assumed) HS colloidal particles might lead to higher nucleation rates at low-moderate packing fractions, and that phase may post-critically grow through a parasitic crystal phase

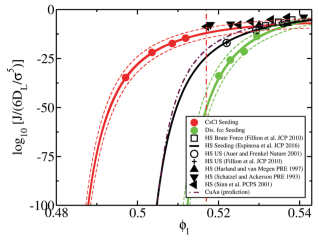


Fig. 11 Logarithm of the nucleation rate versus liquid packing fraction for several colloidal systems. Red, green and maroon (dashed dotted) curves represent the CsCl, disordered fcc and CuAu (predicted) nucleation rates of charged colloidal crystals, respectively, while the black curve indicates J for fcc crystals of purely repulsive HS from ref. 20 using the Seeding method. Dashed red and green curves indicate the upper and lower bounds of uncertainty of the Seeding results for the CsCl and disordered fcc phases, respectively. Empty circles represent the computational estimations of J for HS using Umbrella sampling calculations, <sup>18,120</sup> whereas cross symbols and empty squares represent those calculated via Forward flux sampling and brute force simulations, respectively (ref. 21). Black filled triangles depict different experimental measurements of the nucleation rate for hard-sphere colloids from ref. 8-10. The vertical red dashed dotted line represents the lowest liquid packing fraction at which CsCl nuclei postcritically grow as disordered fcc crystals (parasitic crystallization).

with a lower J but a faster growth rate similarly to the phenomenon that occurs in oppositely charged colloids with the disordered fcc phase.

### V. Concluding remarks

In this work, we study the crystal nucleation scenario of oppositely charged colloids by means of computer simulations. For that purpose, we develop a continuous potential that allows highlyefficient parallel MD simulations of unprecedentedly large systems and that accurately reproduces the original Yukawa model parametrized for describing the phase diagram of colloidal electrolytes. We focus on the crystal nucleation competition of three different solids at moderate temperature,  $T^* = 1$ ; the CsCl crystal, the CuAu crystal, and a substitutionally disordered fcc phase. On the one hand, our findings show that, from low to moderately-high packing fraction,  $\phi_1$  < 0.53, the CsCl crystal phase exhibits the fastest nucleation rate, while at higher  $\phi_1$ , a cross-over between the nucleation rate of the disordered fcc phase (and possibly of the CuAu phase as well) and that of the CsCl polymorph takes place, with nucleation of the two (three) phases becoming comparably quick. A martensitic transformation of the CuAu clusters into CsCl ones is also observed at any packing fraction below spontaneous crystallization.

Our results reveal that colloidal electrolytes only show standard nucleation behaviour at low packing fractions,  $\phi_1$  < 0.517, where

the stable phase, CsCl, is the one that nucleates and grows. In contrast, at higher packing fractions ( $\phi_1 > 0.53$ ), neither the nucleating nor the post-critical growing phase is the stable crystal phase. We observe that the post-critical crystal growth of the nuclei in the moderate/high  $\phi_1$  regime (i.e., between  $\phi_1 = 0.517$  and  $\phi_1 = 0.535$ ) occurs so fast that the growing crystals are predominantly composed of disordered fcc domains rather than constituted by crystals of the stable CsCl or CuAu phases. Furthermore, we analyse the thermodynamic and kinetic factors leading to this complex polymorphic scenario, finding that the free energy cost of the cluster formation

is the crucial factor in determining which phase nucleates the

fastest, whereas the crystal growth rate of the different poly-

morphs is what determines the post-critical crystallization.

We illustrate how a phase that is not thermodynamically stable (the disordered fcc phase) manages to grow at postcritical level using the critical nucleus of the stable phase (CsCl). We term this behaviour parasitic crystallization. Nonetheless, to understand the values of J, one needs to use the nucleation rate of the stable thermodynamic phase (CsCl). Thus, the mechanism of crystallisation, nucleation and growth, may entail some surprises since one phase can firstly nucleate and then grow another one at post-critical level. However, it seems that, to understand the value of *J* at low and moderately high  $\phi_1$  values, the nucleation rate of the stable phase (CsCl in this case) should be considered.

To conclude, we compare the oppositely charged colloidal nucleation scenario with that of purely repulsive hard-spheres. Our results show that the interfacial free energy of HS fcc clusters is very similar to those of CsCl and disordered fcc clusters. Moreover, we note that the CsCl nucleation rate surprisingly lies close to the experimental nucleation rates of hard-sphere colloids, and that the computational J of HS resembles our prediction for the CuAu phase. Taken together, our results provide a panoramic perspective of the nucleation landscape of oppositely charged colloids and rationalise the different kinetic and thermodynamic aspects behind it.

#### Conflicts of interest

There are no conflicts to declare.

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