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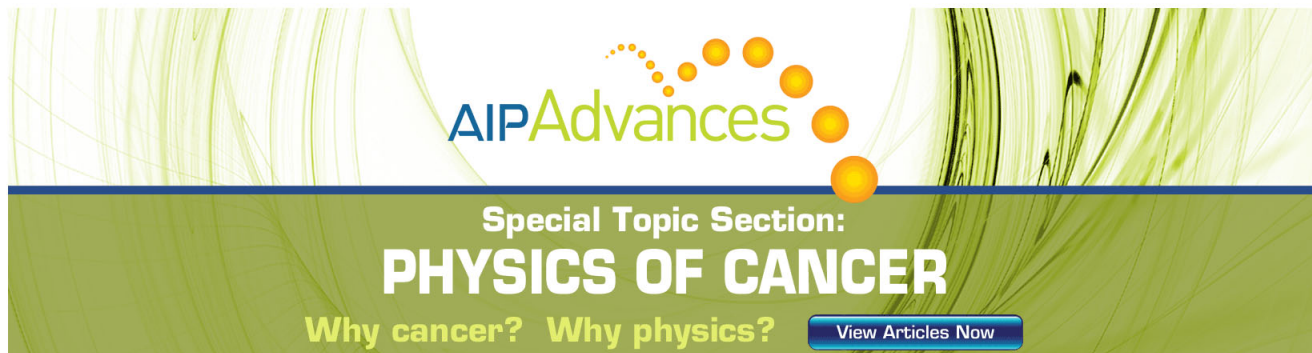
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Tuning effective interactions close to the critical point in colloidal suspensions

Nicoletta Gnan, Emanuela Zaccarelli, and Francesco Sciortino

Dipartimento di Fisica and CNR-ISC, Università di Roma "La Sapienza," Piazzale A. Moro 2, 00185 Roma, Italy

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We report a numerical investigation of two colloids immersed in a critical solvent, with the aim of quantifying the effective colloid-colloid interaction potential. By turning on an attraction between the colloid and the solvent particles we follow the evolution from the case in which the solvent density close to the colloids changes from values smaller than the bulk to values larger than the bulk. We thus effectively implement the so-called (+, +) and (−, −) boundary conditions defined in field theoretical approaches focused on the description of critical Casimir forces. We find that the effective potential at large distances decays exponentially, with a characteristic decay length compatible with the bulk critical correlation length, in full agreement with theoretical predictions. We also investigate the case of (+, −) boundary condition, where the effective potential becomes repulsive. Our study provides a guidance for a design of the interaction potential which can be exploited to control the stability of colloidal systems. © 2012 American Institute of Physics. [<http://dx.doi.org/10.1063/1.4745479>]

I. INTRODUCTION

Interactions between colloidal particles in dispersions depend, beside the colloid coordinates, on the degrees of freedom of the solvent and of the co-solutes. When a clear separation in time and space scales between solvent molecules and colloidal particles exists, it is possible to integrate out the solvent (and co-solute) degrees of freedom and derive an effective potential that describes the interaction between colloids.¹ A well known case is the depletion interaction derived long time ago by Asakura and Oosawa² which have shed light on the role of entropic forces on the phase diagrams of colloidal suspensions. Depletion interactions arise whenever small additives, such as polymers or surfactants, are added to colloidal dispersions; when two colloids are closer than the typical size of the co-solute, the latter is excluded by a “depletion” region in between the two colloids. As a result, a pressure gradient originates, giving rise to a short-range entropy-driven attractive effective potential – the depletion potential. Today we know that depletion forces can be strong enough to induce colloidal phase separation,^{3–5} or to enhance the stability of the crystalline phase, as in the case of proteins.^{6,7}

An interesting effective potential is the one arising when two colloids are dispersed in a critical fluid. Close to the critical point, thermal fluctuations of the order parameter are correlated over length-scales that are much larger than the solvent size and their properties become independent of the microscopic behavior of the system, i.e. they are *universal*. The confinement of the order-parameter fluctuations in between the two colloids give rise to long-range effective forces, named critical Casimir forces.⁸

In the last decade direct experimental evidence of such forces has been found.⁹ Theoretical investigation of critical Casimir forces, which started with the pioneering work of Fisher and de Gennes¹⁰ back in 1978, has provided accurate

predictions for the radial dependence of the effective potential and the associated scaling properties. Such studies, capitalizing on the universality aspect of the problem, have been mostly focusing on Ising model calculations^{11–14} and, more recently, on a field theoretical approach for a classical binary mixtures.^{15–19} The main result of these studies is the explicit derivation of the critical Casimir potential for a solvent confined between the surfaces of two large colloids:

$$\beta\Phi(z) = \frac{\sigma_c}{z} \Theta(z/\xi), \quad (1)$$

where σ_c is the colloid diameter and z is the distance between the surfaces of the two colloids. The scaling function $\Theta(z/\xi)$ depends on the distance from the critical point (via the correlation length ξ), on the bulk universality class of the solvent and on the boundary conditions (BC) imposed by the colloidal surface properties. It has been demonstrated¹⁵ that for $z \gg \xi \gg \sigma_s$ (the latter being a measure of the solvent size), the behavior of the scaling function in Eq. (1) is

$$\Theta(z/\xi)_{(\pm,\pm)}(z/\xi \gg 1) = \pi A_{(\pm,\pm)}(z/\xi) e^{-z/\xi}, \quad (2)$$

where the (\pm, \pm) signs are related to different BCs, i.e., to the different absorption preferences of the confining surfaces with respect to the solvent: (+, +) or (−, −) corresponds to symmetric BC and (+, −) (or equivalently (−, +)) to asymmetric BC. Experimental results have shown that it is possible to generate repulsive and attractive critical Casimir forces between a colloid and a substrate by modifying the BC of the substrate.⁹ Moreover it has been shown that it is possible to continuously tune the force from attractive to repulsive by producing a gradient in the physico-chemical properties of the substrate.²⁰ This can be exploited for inducing orientation-dependent effective forces in colloids close to chemically patterned surfaces.^{21,22} Critical Casimir forces have also been observed when the critical behavior of the host medium refers not to the solvent itself but to the presence of small

interacting co-solutes added in solution. If the inter-cosolutes interaction becomes strong enough, a phase separation (the analog of the gas-liquid) takes place in which the co-solutes partition themselves in two phases of different concentration. Close to the corresponding critical point, the critical fluctuations in co-solute concentration generate critical Casimir forces. Such mechanism has been exploited in the experimental study of Buzzaccaro *et al.*,¹⁸ in which PMMA colloidal particles are dispersed in an interacting micellar solution. In the same work, the authors presented an interesting connection between depletion forces generated by the presence of the micelles far from the critical point and Casimir forces generated by the critical fluctuations close to the micellar critical point.

Most of the theoretical and numerical investigations of critical Casimir forces have been based on studies of lattice models, exploiting the universality properties of the phenomenon. In a recent study,²³ we have reported a numerical evaluation of the effective interaction potential between two spherical hard-sphere colloidal particles immersed in a critical depletant, with the aim of exploring how the interaction potential changes from the depletion shape occurring far from the critical point, to the universal shape induced by critical fluctuations close to the depletant critical point. In the study we have modeled the depletant interaction as a short-range square-well attractive potential. We have confirmed the critical nature of the effective potential close to the critical point by showing that the quantity ξ entering in the effective potentials (see Eq. (2)) is consistent with the bulk correlation length of the critical co-solute extracted from the static structure factors. Interestingly enough, we have found that the strength of the effective potential between the two hard-sphere colloids (when the colloid-depletant interaction is only controlled by excluded volume, and hence in the $(-, -)$ BC class) is sufficient to drive bulk phase separation of the colloidal solution well before the critical region is approached.

In this article we explore the changes in the effective potential when the colloid-cosolute interaction is continuously modified from hard-core repulsion to strong attraction, continuously moving from the $(-, -)$ to the $(+, +)$ BC. We also explore the interesting case in which $(+, -)$ BC are present, i.e., the case of two colloids interacting in different ways with the co-solutes, again interpolating between the $(-, -)$ and the $(+, -)$ limits on changing (this time only for one of the two colloids) the colloid-cosolute interaction strength. We calculate numerically the effective potential both at a high T , where critical phenomena are absent as well as close to the co-solute critical point.

II. MODELS AND METHODS

The total colloid-colloid interaction potential ϕ_{CC} results from the sum of the bare colloid-colloid interaction V_{CC} and of the effective potential V_{eff} arising from the integration of the solvent and co-solute degrees of freedom,

$$\phi_{CC}(r_{12}) = V_{CC}(r_{12}) + V_{eff}(r_{12}). \quad (3)$$

We model V_{CC} as a excluded volume interaction between two colloids of size σ_c

$$V_{CC}(r_{12}) = \begin{cases} \infty, & r_{12} < \sigma_c \\ 0 & r_{12} \geq \sigma_c \end{cases}. \quad (4)$$

Inspired by the work of Buzzaccaro *et al.*,¹⁸ we model the critical medium as a fluid of interacting co-solutes dispersed in an implicit solvent. The co-solutes, of size σ_s , interact via a pairwise square-well potential (SW)

$$V_{SS}(r_{ij}) = \begin{cases} \infty, & r_{ij} < \sigma_s \\ -\varepsilon_s, & \sigma_s \leq r_{ij} < (1 + \delta)\sigma_s, \\ 0 & r_{ij} \geq (1 + \delta)\sigma_s \end{cases}, \quad (5)$$

where ε_s controls the strength of the interaction and δ the relative (respect to σ_s) width of the well. σ_s and ε_s are chosen as unit of length and energy. The temperature T is measured in units of ε_s . The co-solute is characterized by a gas-liquid critical point located at $(T_c = 0.478, \phi_c = 0.25)$,²⁴ where $\phi_c = (\pi/6)\rho_c\sigma_s^3$ is the critical packing fraction of the co-solute and ρ_c is its number density. For this model, it has been shown²³ that the correlation length and the susceptibility extracted from the static structure factors close to the critical point diverge with a power-law with the respective Ising critical exponents. The size ratio $q \equiv \sigma_s/\sigma_c$ between the co-solute and the colloids is fixed at $q = 0.1$.

To evaluate the effective potential, we perform Monte Carlo simulations of two colloids in a fluid of co-solute particles, in the canonical ensemble at fixed T and ρ_s in a parallelepiped-shaped box. The two colloids are constrained to move only along the x axis, sampling only a limited range of distances. Several overlapping relative distances windows are simulated, evaluating for each window $P(r)$, the probability of observing the two colloids at relative distance r . Splicing together the $P(r)$ evaluated in different windows provides an effective (and parallel) way for evaluating the entire $P(r)$. The logarithm of $P(r)$ is by definition the effective potential (apart from an overall constant which is fixed imposing $V_{eff}(\infty) = 0$). To minimize finite-size effects at the temperatures investigated, the dimensions of the box ($L_x; L_y; L_z$) are chosen in such a way that the surface-to-surface colloidal distance evaluated via the boundary conditions is more than twice the distance over which the effective potential goes to zero. Moreover, along all directions, the solvent density profile reaches a constant value on approaching the box boundaries. Close to the critical point and along the critical isochore, the size of the box is $[L_x = 52; L_y = 26; L_z = 26]$, requiring 16 000 co-solute particles. The bulk density is estimated *a posteriori* by calculating the local density far from the two colloids.

We also evaluate the co-solute density profile for different BC by selecting a volume centered along the x axis of transversal section equal to σ_s^2 and we average the local density with a mesh of the order of $0.1\sigma_s$.

In this paper, when discussing the critical behavior, we will show results for effective potentials and density profiles evaluated at the critical packing fraction and at the reduced

temperature $T/T_c = 1.0251$, corresponding to a critical correlation length $\xi = 2.5\sigma_s$.²³

III. RESULTS

A. (−, −) BC

In our previous work²³ we have discussed the evolution of the effective potential when the interaction between the colloids and the solvent V_{CS} are treated as hard-spheres. In such condition, close to the colloid surface, the density of the solution is smaller than the average, effectively generating a (−, −) BC. We have shown that close to the critical point, V_{eff} is long-ranged, signaling the onset of criticality. Its radial dependence is well described by the exponential decay of Eq. (1), with the same characteristic length ξ of the bulk critical fluctuations.

To get more insight into the mechanism which drives attraction between colloids close to the critical point in the (−, −) BC case, we show in Fig. 1 the co-solute density profile along the horizontal x axis for two different relative colloid-colloid distances. Notice that the two colloids are always located symmetrically with respect to the origin. Thus the density profile is symmetric with respect to $x = 0$. For this reason in Fig. 1 we show only the positive x region. In between the two colloids, the density is significantly lower than the bulk density, a typical depletion effect. Outside, the density relaxes toward the bulk value with an exponential decay, again controlled by the bulk critical correlation length. It is interesting to discuss the physical origin of the net attractive force between the two colloids. Since the co-solute colloid in-

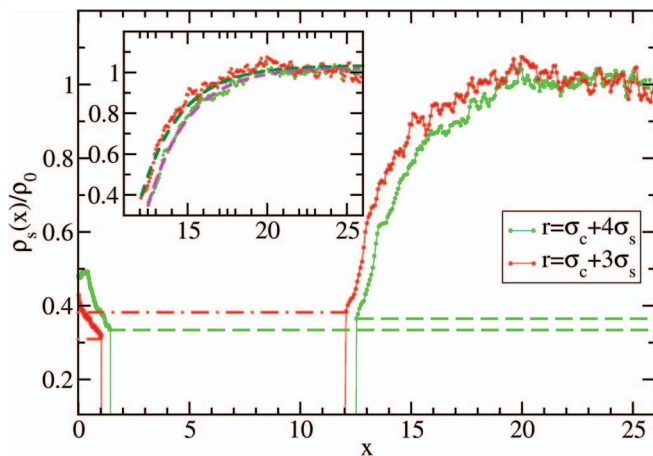


FIG. 1. Normalized co-solute density profiles with respect to the bulk density ρ_0 along the horizontal x axis at $T/T_c = 1.0251$ and at the critical density $\phi_c = 0.2501$. Co-solute particles interact with a short-range SW attraction while colloid-cosolute interaction is modeled with a HS repulsion. The two profiles are evaluated for two different colloid-colloid distances, namely for $r = \sigma_c + 3\sigma_s$ and $r = \sigma_c + 4\sigma_s$, with $\sigma_c = 10\sigma_s$. The box is centered at $x = 0$ and the two colloids are centered at $\pm r/2$ along the x axis. Being the profiles symmetric with respect to $x = 0$ only positive x are shown here. Note that each colloid excludes the solvent in a region equal to $\sigma_c + \sigma_s$. The dashed lines highlight the mismatch in the density at contact between the inner and outer side of the colloid. The inset shows that the outer part of the density profile is well fitted by an exponential $\rho = \rho_0 + A\exp(-r/\xi)$, where ξ has been fixed to $\xi = 2.5\sigma_s$, i.e., to the bulk critical correlation length value for $T/T_c = 1.0251$ at ϕ_c . In this case the fit parameter $A < 0$.

teraction is modeled via an hard-sphere potential, it is possible to prove that the effective force results from the mismatch in the contact density along the two sides of each colloid. Figure 1 shows that indeed, at contact, the density outside is slightly larger than inside. The figure also shows that the mismatch decreases on increasing the relative distance between the two colloids in parallel with the decrease of the effective force.

B. From the (−, −) to the (+, +) BC

To drive the transition from (−, −) toward (+, +) BC we tune the strength of a short-range attraction in the colloid-cosolute interaction V_{CS} . The attraction gives rise to an enhanced accumulation close to the surface of the colloids, at first compensating and then inverting the depletion characteristic effect of the excluded volume interaction.^{25,26}

We model the attraction via a pairwise SW potential

$$V_{CS}(r_{ij}) = \begin{cases} \infty, & r < \sigma_{cs} \\ -\varepsilon, & \sigma_{cs} \leq r < \sigma_{cs} + \Delta\sigma_s \\ 0 & r \geq \sigma_{cs} + \Delta\sigma_s \end{cases}, \quad (6)$$

where $\sigma_{cs} = (\sigma_s + \sigma_c)/2$ and $\Delta = 0.35$. The width $\Delta\sigma_s$ has been chosen as a compromise between limiting the colloid-cosolute interaction to the nearest-neighbor shell and maximizing the volume over which co-solute bind to the colloid. The parameter ε is used as control parameter to drive the cross-over from the hard-sphere like behavior ($\varepsilon \rightarrow 0$) to the wetting case $\varepsilon/kT \gtrsim 1$).

We start by discussing the behavior of the effective potential at high T , where critical phenomena are not present. The evolution of the effective potential upon changing ε is shown in Fig. 2. When $\varepsilon = 0$, the well known depletion interaction is observed. On increasing ε , the depletion attractive interaction is progressively weakened, and the potential at contact becomes repulsive. For even large ε values, the colloid becomes surrounded by a persistent layer of co-solutes which extent the effective radius of the colloid, making it impossible to attract a neighboring colloid for distances closer than $\sigma_c + \sigma_s$.²⁷ Under such strong coupling conditions, the effective potential acquires an oscillatory character, with minima originating from the preferential distances allowing for an integer number of co-solute layers between the colloids. The configuration associated to the first minimum, called bridging, is the most energetically favorable since one co-solute particle is bonded with both colloids. The other minima are related to particular configurations in which bridging is obtained by particle chains. A sketch of such situation is shown in Fig. 2. The intermediate maxima occur when the co-solute particles (or particle chains) are not bonded to both colloids. For example the first maximum corresponds to a situation, also illustrated in Fig. 2, in which a single co-solute particle cannot be bonded to both colloids, since $r > \sigma_s$. The evolution of the effective potential reported in Fig. 2 clearly show how the minimum at contact progressively turns into a maximum, and simultaneously a new minimum develops at the bridging distance. Intermediate values of ε thus provide a viable mechanism for contrasting depletion interaction and favoring colloidal stability. The case $\varepsilon = 0.5$ is emblematic, since the

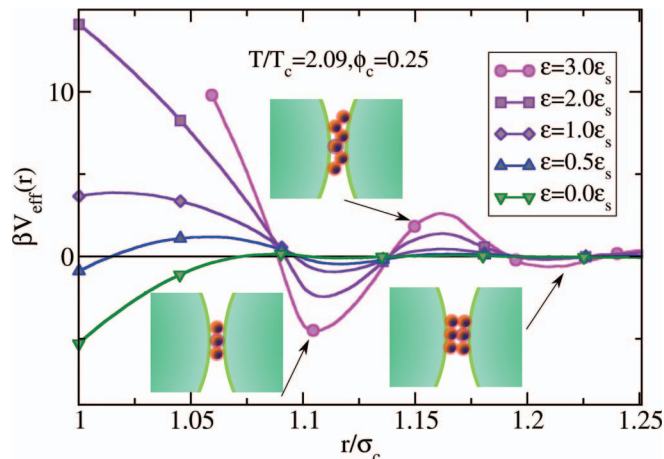


FIG. 2. Effective potential for two large colloids interacting through a hard-core repulsion at $T/T_c = 2.09$ and $\phi = \phi_c = 0.25$. Colloids are in solution with co-solute particles interacting (among themselves) via a SW attraction of width $\delta\sigma_s = 0.1\sigma_s$ and depth ε_s . The two colloids interact with co-solute particles via a SW attraction of width $\Delta\sigma_s = 0.35\sigma_s$ and depth ε . Sketches in the figure represent particular configurations adopted by the co-solute for different colloid-colloid distances. Such configurations give rise to maxima and minima in the potential.

minimum at contact has essentially disappeared while the new minimum at distance $1.1\sigma_c$ has not yet developed, such that the effective potential is never (in absolute value) significantly larger than $k_B T$. For very large ε values, V_{eff} becomes again sufficiently intense to drive a colloid phase separation.

Close to the critical point, critical Casimir forces add to the standard effects described before. Differently from the pure $(-, -)$ case, the presence of an attractive interaction between the colloid and the solvent brings in two new features that are distinguishable for medium and large values of ε : a repulsive contribution to the effective potential at short distances and the presence of oscillations induced by the granularity of the co-solutes. The theory on critical Casimir forces predicts that at distances larger than ξ the effective potential develops an attractive exponential tail, so that the effective potential behaves the same for the $(-, -)$ BC and the $(+, +)$ BC cases. Figure 3 shows how the effective potential evolves on increasing the value of ε . As expected the contact value becomes repulsive but the long tail behavior remains always attractive and has the same exponential character of the $\varepsilon = 0$ case.²³ Even for large values of ε , where the same oscillations characterizing the high T effective potentials modulate the shape, the overall behavior can be represented by an exponential function. The density profile for the $(+, +)$ case is shown in Fig. 4. Close to the colloidal surfaces a significant layering of the co-solutes is observed, consistent with the presence of oscillations in $V_{eff}(r)$. On the external sides, beyond the oscillations, the density decays again with an exponential shape controlled once more by ξ . In comparison with the $(-, -)$ case, the resulting interaction between the two particles is more difficult to visualize, since it arises from the competition between the density at contact (in which the solvent pushes the colloid) and the density at the well boundary (where the solvent attracts the colloid). Indeed, in the case of a colloid of diameter σ_c interacting via square-well interaction with solvent parti-

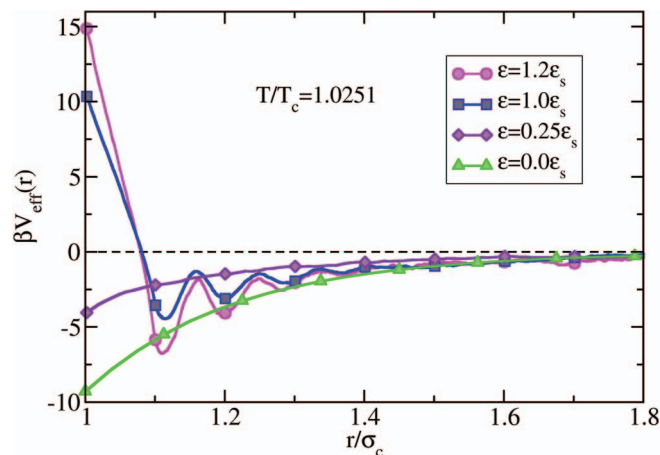


FIG. 3. Effective potential of two large colloids close to the critical point of the co-solute ($T/T_c = 1.0251$ and $\phi = \phi_c = 0.25$). The two colloids interact with co-solute particles via a SW attraction of width $\Delta\sigma_s = 0.35\sigma_s$ and depth ε . The two colloids have therefore $(+, +)$ BC. The increase of the attraction strength gives rise to large oscillations associated to bridging effects as discussed in Fig. 2.

cles, the pressure originates from the two points of discontinuity of the potential, the colloid-cosolute hard-wall distance (HWD) and the square-well distance (SWD) $\sigma_{cs} + \Delta\sigma_s$, according to the expression²⁸

$$\frac{\beta P^{ex}}{\rho} = \frac{2}{3}\pi\rho[\sigma_{cs}^3 g(\sigma_{cs}^+) - (\sigma_{cs} + \Delta\sigma_s)^3 g(\sigma_{cs} + \Delta\sigma_s^-)(1 - e^{\beta u_0})], \quad (7)$$

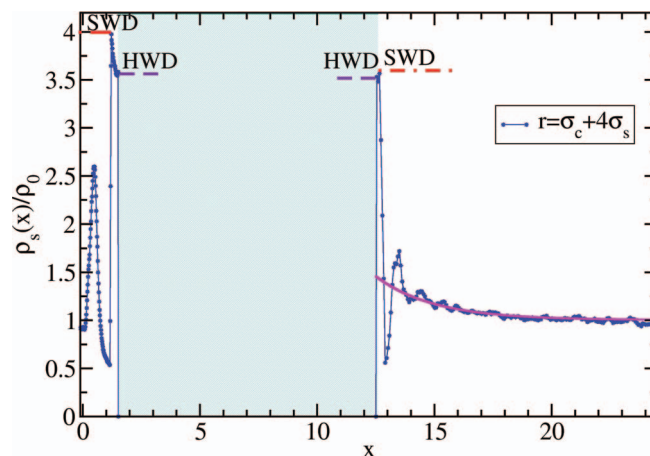


FIG. 4. Normalized co-solute density profile with respect to the bulk density ρ_0 along the horizontal x axis at $T/T_c = 1.0251$ and at the critical density $\phi_c = 0.2501$. Co-solute particles interact with a short-range SW attraction of width $\delta\sigma_s = 0.1\sigma_s$ and depth ε_s while colloid-cosolute interaction is modeled with a SW interaction of width $\Delta\sigma_s = 0.35\sigma_s$ and depth $\varepsilon = 1\varepsilon_s$. The box is centered at $x = 0$ and the two colloids are centered at $\pm r/2$ along the x axis. Being the profile symmetric with respect to $x = 0$ only positive x are shown here. Note that each colloid excludes the solvent in a region equal to $\sigma_c + \sigma_s$. The dashed segments highlight the mismatch in the density at contact (HWD) and at the well boundary (SWD) between the inner and outer side of the colloid. The profile is associated to the effective potential in Fig. 3 (squares). As for Fig. 1, the criticality of the co-solute can be caught from the exponential decay of the density outside the colloids towards $\rho_s(x)/\rho_0 = 1$. The exponential fit shown in the figure (solid line), has been performed by fixing the exponential decay with the value of the bulk critical correlation length for the temperature and density investigated, i.e., $\xi = 2.5\sigma_s$.

where $g(\sigma_{cs})$ is the pair correlation function evaluated at the HWD and $g(\sigma_{cs} + \Delta\sigma_s)$ at the SWD. From Eq. (7) one can notice that the density at the hard-wall generates a positive contribution to the pressure, while the density at the well boundary provides a negative contribution. Figure 4 shows that indeed the density at the SWD inner side (left side of Fig. 4) is significantly higher than the density at the SWD outer side (right side of Fig. 4), and it is responsible for the resulting net attractive force between the two colloids. Indeed, the excluded volume contribution in this case would tend to separate the two colloids, being the contact density outside slightly larger than the contact density inside.

C. From the $(-, -)$ to the $(+, -)$ BC

We now discuss the situation in which one of the two colloids (C_1) interacts with the co-solute through a square-well attraction ($V_{C_1S}(r_{1i})$) while the other colloid (C_2) experiences only excluded volume interactions $V_{C_2S}(r_{2j})$. More precisely

$$V_{C_1S}(r_{1i}) = \begin{cases} \infty, & r_{1i} < \sigma_{cs} \\ -\varepsilon, & \sigma_{cs} \leq r_{1i} < \sigma_{cs} + \Delta\sigma_s \\ 0 & r_{1i} \geq \sigma_{cs} + \Delta\sigma_s \end{cases} \quad (8)$$

and

$$V_{C_2S}(r_{2j}) = \begin{cases} \infty, & r_{2j} < \sigma_{cs} \\ 0 & r_{2j} \geq \sigma_{cs}. \end{cases} \quad (9)$$

Also in this case we start investigating a T significantly larger than the critical one for different interaction strengths ε . The results are shown in Fig. 5.

The trend is similar to the one observed going from $(-, -)$ to the $(+, +)$ case; (i) the contact value of the effective potential grows continuously on increasing ε , progressively offsetting the original $\varepsilon = 0$ depletion interaction. Still, the strength of the repulsion is significantly smaller than the one observed in the $(+, +)$ case. (ii) only weak oscillations (with

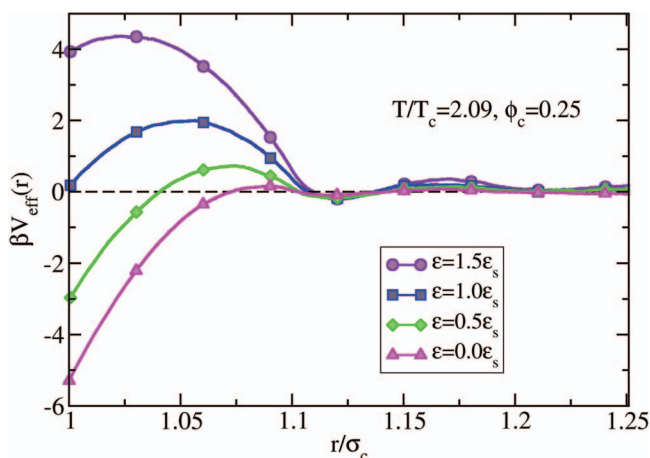


FIG. 5. Effective potentials for two large colloids interacting through a hard-core repulsion at $T/T_c = 2.09$ and $\phi = \phi_c = 0.25$. Co-solute particles, in solution with the two colloids, interacts (among themselves) through a SW attraction of width $\delta = 0.1\sigma_s$ and depth ε_s . One of the two colloids (C_1) interact with co-solute particles via a SW attraction of width $\Delta = 0.35\sigma_s$ and depth ε . The other colloid (C_2) has instead a HS repulsion with the co-solute.

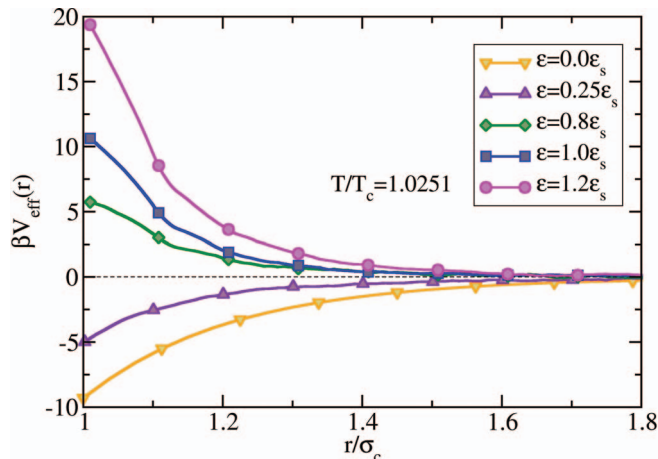


FIG. 6. Evolution of the effective potential of two colloids close to the critical point of the co-solute ($T/T_c = 1.0251$ and $\phi = \phi_c = 0.2501$). One of the two colloids (C_1) interacts with co-solute particles via a SW attraction of width $\Delta = 0.35\sigma_s$ and depth ε . Different curves refer to different values of ε . The other colloid (C_2) interacts via a HS repulsion with the co-solutes. This setting provides a realization of $(+, -)$ BC.

amplitude smaller than $k_B T$) characterize the radial dependence of the effective potential, signaling the absence of a strong layering of the co-solute between the colloids. We notice that similar features have also been observed in the case of non-additive HS mixtures.^{29,30}

Close to the critical point, the theory predicts that, on increasing ε , V_{eff} changes from an attractive to a repulsive exponential decay, on changing the boundary conditions from $(-, -)$ to $(-, +)$. The effective potentials for different ε close to the critical point are plotted in Fig. 6. While for $r < 1.3\sigma_c$ the echo of the layering effects is still visible, for larger r values the decay of all curves is compatible with the same exponential function decay, again supporting the identification of the interaction potential in this spatial region as arising from the universal behavior imposed by the critical fluctuations. As in the case of $(-, -)$ BC, for $(+, -)$ BC the density profile provides useful informations on the mechanism that gives rise to the repulsion. Figure 7 shows the co-solute particles density profile along the x axis. The different colloid-co-solute interaction results in a different density profile around the two colloids. Around C_2 ($-$ BC), the solvent density is lower than the average, while the opposite behavior is observed for C_1 ($+$ BC). The repulsive force on C_2 originates from the slight mismatch of the contact density (larger inside than outside), while the repulsive force on C_1 originates from larger density at the SWD on the outside, compared to the inside. Contrary to what observed in the $(-, -)$ to $(+, +)$ BC case, here tuning the attraction strength allows us to modify the sign of the effective force.⁹ In fact we observe that for small ε the effect of the colloid-co-solute attraction is not sufficiently strong to substantially change the shape of the effective potential, which remains completely attractive. For higher values of ε the potential turns into a completely repulsive one and no sign of oscillations driven by co-solute structures is visible. It is interesting to note that under these conditions it is in principle possible to tune finely ε in order to obtain a flat co-solute density profile

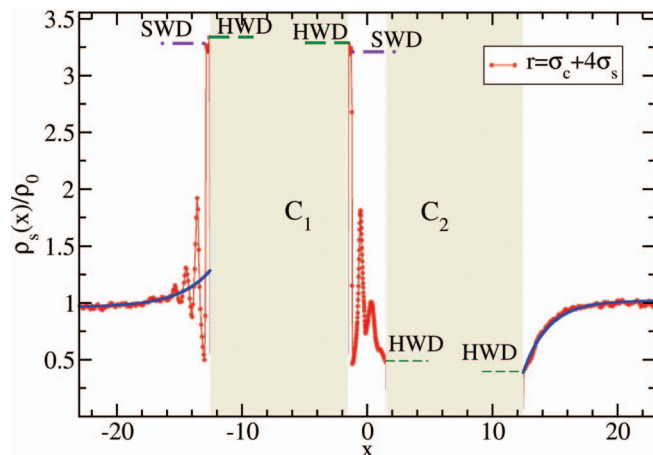


FIG. 7. Normalized co-solute density profile with respect to the bulk density ρ_0 along the horizontal x axis at $T/T_c = 1.0251$ and at the critical density $\phi_c = 0.2501$. Co-solute particles interact with a short-range SW attraction of width $\delta = 0.1\sigma_s$ and depth ε_s . One of the two colloids (C_1) interacts with co-solute particles via a SW attraction of width $\Delta = 0.35\sigma_s$ and depth $\varepsilon = 1\varepsilon_s$. The other colloid (C_2) interacts via a HS repulsion with the co-solute. The profile is associated to the effective potential in Fig. 6 (squares). The different colloid-cosolute interaction results in a different density profile around the two colloids. Around C_2 (colloid on the right), the solvent density is lower than the average, while the opposite behavior is observed on C_1 (colloid on the left). The repulsive force on C_2 colloid originates from the slight mismatch of the contact density (larger inside than outside), while the repulsive force on C_1 originates from larger density at the square-well location on the outside, compared to the inside. The exponential fits (solid lines) have been done by setting the exponential decay to the value of the bulk critical correlation length $\xi = 2.5\sigma_s$.

(apart from the layering at contact). This corresponds to impose Dirichlet BC¹² and from a conceptual point of view to identify a sort of Θ condition, in analogy with polymer solutions,¹ where the effective interaction potential is close to zero.

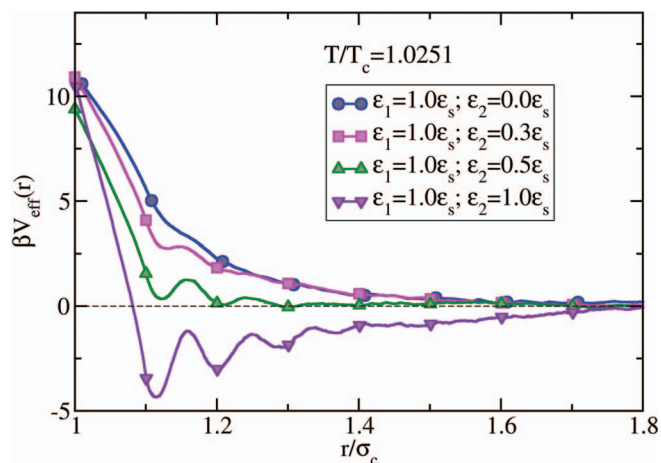


FIG. 8. Evolution of the effective potential for two colloids close to the critical point of the co-solute ($T/T_c = 1.0251$ and $\phi = \phi_c = 0.2501$). One of the two colloids (C_1) interacts with co-solute particles via a SW attraction of width $\Delta = 0.35\sigma_s$ and depth $\varepsilon_1 = 1.0\varepsilon_s$. The other colloid (C_2) interacts with the co-solute through a SW attraction as well with the same width Δ but depth ε_2 . Starting from $\varepsilon_2 = \varepsilon_1$, i.e. (+, +) BC, the interaction strength ε_2 is decreased down to the limit case of $\varepsilon_2 = 0$ corresponding to (+, -) BC. Contrary to the case illustrated in Fig. 6, here only the long distance tail can be tuned from attractive to repulsive, while the short range part of the potential remains repulsive.

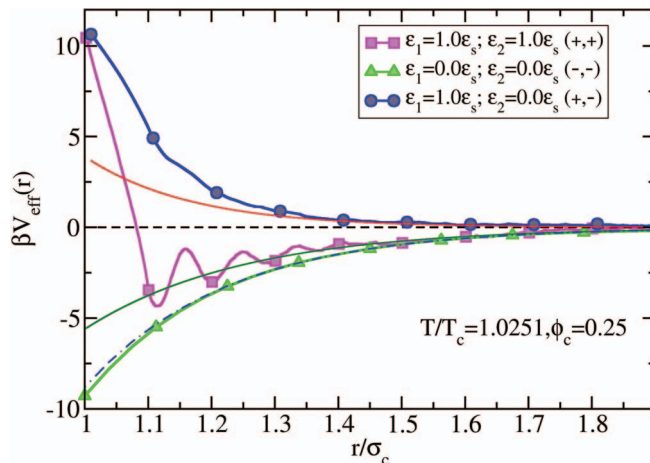


FIG. 9. Effective potentials for different BC, evaluated at the co-solute critical packing fraction and at $T/T_c = 1.0251$. In the critical region, independently on the BC, the long range part of the effective potential decays exponentially. According to Eq. (1) it is possible to extract the characteristic length that controls the decay of the exponential. To extract this value the three curves have been interpolated in the same range with Eq. (1) starting from $r = 1.3\sigma_c$. For the three cases we find $\xi_{(-,-)} = 2.24\sigma_s$, $\xi_{(+,+)} = 2.49\sigma_s$, $\xi_{(+,-)} = 2.20\sigma_s$ which is consistent for the bulk correlation length $\xi = 2.5\sigma_s$ of the co-solute for this T/T_c .²³ The non universal amplitudes found from the exponential fits are $A_{(-,-)} = -0.63$, $A_{(+,+)} = -0.45$, $A_{(+,-)} = 0.48$.

D. From (+, +) to (+, -) BC

According to theoretical predictions¹⁵ the effective potential is expected to change in the transition from (+, +) to (+, -) exactly as in the case from (-, -) to (+, -). These predictions refer to the scaling region of the potential. In the present numerical study, we also access the short distances, where the effective potential probes the non-universal aspects of the solvent-colloid interaction. To highlight the difference at short distances between the two cases, Fig. 8 shows the evolution of the effective potential when the attraction between only one of the two colloids and the co-solute is progressively reduced. Differently from the results of Fig. 6, in this case only the long distance behavior changes sign, while the short distance part of the potential remains always repulsive. Finally in Fig. 9 we plot cases differing in their BCs, but all at the same temperature and co-solute critical packing fraction. In all cases, the long-distance behavior of V_{eff} can be described, as expected theoretically, by an exponential decay, with a correlation length consistent with the bulk critical correlation length ξ .

IV. CONCLUSIONS

In this article we have discussed how the effective potential, resulting from confining critical co-solute particles between the surfaces of two large colloids, depends on the colloid-cosolute interaction. In a previous study²³ we have already investigated the case of (-, -) BC, i.e. when only colloid-cosolute hard-core repulsion is present. In such case we have shown that the resulting potential for $T/T_c \rightarrow 1$ at the co-solute critical packing fraction is monotonic, attractive, and long range. The co-solute density profile along the x axis of the simulation box shows that for (-, -) BC the

density close to colloids is lower than the bulk density and it relaxes exponentially to the bulk value far from the two colloids. Upon switching on the colloid-cosolute attraction ε , depletion effects are progressively weakened and the cosolute density close to the colloid varies from values smaller than the bulk to values larger than the bulk. Correspondingly, the contact value of the potential varies from negative to positive while the long-distance part of $V_{\text{eff}}(r)$, dominated by critical fluctuations, remains always attractive as theoretically predicted in the (+, +) BC case. The increase in ε has instead a profound effect on the non-universal short-distance part of the potential, which progressively develops large oscillations. Such oscillations are related to specific geometries at characteristic lengths associated to integer number of cosolutes between the two colloids. At these specific distances, the local energy is minimized. For intermediate values of ε the colloid-cosolute attraction compensates the depletion interaction, effectively reducing the short-distances interaction potential. We have also addressed the case of two colloids interacting differently with the co-solutes, a realization of the so-called (+, -) BC. In this case, the local density close to the two colloids is respectively lower and higher than the bulk value and the critical Casimir forces are expected to be repulsive. We have shown that indeed, when the asymmetry in the interaction potential is sufficiently intense to drive different BC, the potential becomes repulsive at all length scales. We have shown that, independently on the BCs, the critical long range part of the effective potential is always described by an exponential whose decay is controlled by the critical correlation length ξ describing the thermal correlation length of the bulk co-solute close to the critical point, in full agreement with theoretical predictions.¹⁵ We have investigated the behavior of the density profile in all examined BC cases. Building on the fact that all interactions are modeled as excluded volume or as square-well attraction, the net pressure over the colloids can be estimated simply by the density of co-solute at contact and at the square-well distance (in the case of attraction between the colloid and the co-solute). The density profiles confirm that in the (-, -) case, the contact density inside is smaller than outside, determining the net attraction. In the (+, +) case a different mechanism for attraction is observed: the leading contribution arises from the mismatch of the density at the well distance, larger inside than outside, determining a net attraction. Finally, in the (+, -) case, repulsion is driven by two different mechanisms for the two colloids. The hard-sphere colloid is pushed out by the larger contact density inside. The attractive colloid is dragged out by a larger density at the square-well distance.

The possibility of varying the potential from repulsive to attractive and to finely control its shape and intensity by tuning the BC, provides a guidance for controlling equilibrium properties of colloids dispersed in precritical suspensions. This is important for future applications, for instance in the case in which the confining surfaces are chemically patterned

colloidal particles, such as patchy³¹ or Janus particles.^{21,32} In this case, the introduction of a geometrical constraint in the colloid-cosolute interaction would give rise to a torque that can be used to control the orientation of colloids, as shown for chemically patterned substrates,²² and that could give rise to new unexplored phases. More in general, our results provide useful informations for designing interaction potentials which can be exploited to control the stability of colloidal systems.³³

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- ¹C. N. Likos, *Phys. Rep.* **348**, 267 (2001).
- ²S. Asakura and F. Oosawa, *J. Polym. Sci.* **33**, 183 (1958).
- ³V. J. Anderson and H. N. W. Lekkerkerker, *Nature (London)* **416**, 811 (2002).
- ⁴H. Lekkerkerker, W. C. K. Poon, P. N. Pusey, A. Stroobants, and P. Warren, *Europhys. Lett.* **20**, 559 (1992).
- ⁵M. Dijkstra, R. van Roij, and R. Evans, *Phys. Rev. E* **59**, 5744 (1999).
- ⁶P. R. ten Wolde and D. Frenkel, *Science* **26**, 1975 (1997).
- ⁷R. Piazza, *Curr. Opin. Colloid Interface Sci.* **5**, 38 (2000).
- ⁸M. Kerch, *The Casimir Effect in Critical Systems* (World Scientific, Singapore, 1994).
- ⁹C. Hertlein, L. Helden, A. Gambassi, S. Dietrich, and C. Bechinger, *Nature (London)* **451**, 172 (2007).
- ¹⁰M. E. Fisher and P. G. de Gennes, *C. R. Acad. Sci. Ser. B* **287**, 207 (1978).
- ¹¹O. Vasilyev, A. Gambassi, A. Maciolek, and S. Dietrich, *Europhys. Lett.* **80**, 60009 (2007).
- ¹²O. Vasilyev, A. Gambassi, A. Maciolek, and S. Dietrich, *Phys. Rev. E* **79**, 041142 (2009).
- ¹³T. F. Mohry, A. Maciolek, and S. Dietrich, *Phys. Rev. E* **81**, 061117 (2010).
- ¹⁴O. Vasilyev, A. Maciolek, and S. Dietrich, *Phys. Rev. E* **88**, 041605 (2011).
- ¹⁵A. Gambassi, A. Maciolek, C. Hertlein, U. Nellen, L. Helden, C. Bechinger, and S. Dietrich, *Phys. Rev. E* **80**, 061143 (2009).
- ¹⁶T. F. Mohry, A. Maciolek, and S. Dietrich, *J. Chem. Phys.* **136**, 224902 (2012).
- ¹⁷T. F. Mohry, A. Maciolek, and S. Dietrich, *J. Chem. Phys.* **136**, 224903 (2012).
- ¹⁸S. Buzzaccaro, J. Colombo, A. Parola, and R. Piazza, *Phys. Rev. Lett.* **105**, 198301 (2010).
- ¹⁹R. Piazza, S. Buzzaccaro, A. Parola, and J. Colombo, *J. Phys.: Condens. Matter* **23**, 194114 (2011).
- ²⁰U. Nellen, L. Helden, and C. Bechinger, *Europhys. Lett.* **88**, 26001 (2009).
- ²¹A. Gambassi and S. Dietrich, *Soft Matter* **7**, 1247 (2011).
- ²²F. Soyka, O. Zvyagolskaya, C. Hertlein, L. Helden, and C. Bechinger, *Phys. Rev. Lett.* **101**, 208301 (2008).
- ²³N. Gnan, E. Zaccarelli, P. Tartaglia, and F. Sciortino, *Soft Matter* **8**, 1991 (2012).
- ²⁴J. Largo, M. A. Miller, and F. Sciortino, *J. Chem. Phys.* **128**, 134513 (2008).
- ²⁵S. Karanikas, J. Dzubiella, A. Moncho-Jord, and A. A. Louis, *J. Chem. Phys.* **128**, 204704 (2008).
- ²⁶D. Fiocco, G. Pastore, and G. Foffi, *J. Phys. Chem. B* **114**, 12085 (2010).
- ²⁷G. S. Grest, Q. Wang, P. in't Veld, and D. J. Keffer, *J. Chem. Phys.* **104**, 144902 (2011).
- ²⁸J. P. Hansen and I. R. McDonald, *Theory of Simple Liquids*, 3rd ed. (Academic, Amsterdam, 2006).
- ²⁹A. A. Louis and R. Roth, *J. Phys.: Condens. Matter* **13**, L777 (2001).
- ³⁰R. Roth, R. Evans, and A. A. Louis, *Phys. Rev. E* **64**, 051202 (2001).
- ³¹F. Sciortino and E. Zaccarelli, *Curr. Opin. Solid State Mater. Sci.* **15**, 246 (2011).
- ³²A. Walther and H. Müller, *Soft Matter* **4**, 663 (2008).
- ³³F. Romano and F. Sciortino, *Nat. Commun.* **3**, 975 (2012).