I. INTRODUCTION

Binary mixtures of polymers and colloidal particles in various solvents are the focus of sustained experimental and theoretical efforts, both because they constitute a challenging problem in Statistical Mechanics of “soft matter,” and because of their technological importance in many industrial processes. One of the most striking aspects of polymer–colloid mixtures, namely the depletion interaction between colloids induced by nonadsorbing polymer was recognized nearly 50 years ago. More recently, the importance of the polymer depletant in determining the phase behavior of the mixtures was realized, and much recent experimental work was devoted to the phase diagram, structure, interfaces, and to the direct measurement of the effective interactions. On the theoretical side, most efforts have concentrated on impenetrable spherical colloids, while various models and theoretical techniques have been investigated for the description of the nonadsorbing polymer coils. The models include noninteracting (ideal) polymers, polymers represented as penetrable-spheres, and interacting polymers coarse-grained to the level of “soft colloids.” Monomer level representations of polymer chains, like the self-avoiding walk (SAW) model, appropriate in good solvent, have been considered within polymer scaling approaches, renormalization group (RG) theory, and fluid integral equations.

While many effects for the simplest case of colloids mixed with noninteracting polymer are quantitatively understood, the behavior of the experimentally more relevant case of polymers with excluded volume interactions is at best understood on a qualitative basis; a quantitatively reliable theory is still lacking. Clearly, to construct such a theory for finite concentrations of colloidal particles, one must first understand how interacting polymer coils distribute themselves around a single spherical colloid of radius \(R_c\). This problem is addressed in the present paper using a combination of Monte Carlo (MC) simulations and scaling theories to determine the key quantities, which are the monomer or center-of-mass (cm) density profiles \(\rho(r)\) of SAW polymers around a single impenetrable sphere, as well as the resulting surface tension. If \(R_g\) denotes the radius of gyration of the polymers, these quantities clearly depend on the ratio \(q = R_g/R_c\), which controls the curvature effects. The limit \(q \to 0\), corresponding to a polymer solution near a planar wall, will be examined first, before considering the case of finite \(q\). The complete theory for the opposite limit, \(q \gg 1\), will be the subject of a future publication, although we show some preliminary results here. Throughout this work we focus on the dilute and semidilute regimes of the polymers, where the monomer density \(c\) is low enough for detailed monomer–monomer correlations to be unimportant; the melt regime, where \(c\) becomes appreciable, will not be treated here.

The surface tension of a polymer solution surrounding a sphere is macroscopically defined by considering the immersion of a single hard colloidal particle into a bath of nonadsorbing polymer. Because this immersion reduces the number of configurations available to the polymers, resulting in an entropically induced depletion layer around the colloid, there is a free energy cost \(F_1\) for adding a single sphere to the polymer solution which naturally splits into volume and surface terms

\[
F_1 = \Pi(\rho) \frac{4}{3} \pi R_c^3 + 4 \pi R_c^2 \gamma_s(\rho).
\]

The first term in Eq. (1), describes the reversible work needed to create a cavity of radius \(R_c\) in the polymer solution. Since the osmotic pressure \(\Pi(\rho)\) of a polymer solution

\[\text{Density profiles and surface tension of polymers near colloidal surfaces}

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The surface tension of interacting polymers in a good solvent is calculated theoretically and by computer simulations for a planar wall geometry and for the insertion of a single colloidal hard sphere. This is achieved for the planar wall and for the larger spheres by an adsorption method, and for smaller spheres by a direct insertion technique. Results for the dilute and semidilute regimes are compared to results for ideal polymers, the Asakura–Oosawa penetrable-sphere model, and to integral equations, scaling and renormalization group theories. The largest relative changes with density are found in the dilute regime, so that theories based on noninteracting polymers rapidly break down. A recently developed “soft colloid” approach to polymer–colloid mixtures is shown to correctly describe the one-body insertion free-energy and the related surface tension. © 2002 American Institute of Physics. [DOI: 10.1063/1.1473658]
in the dilute and semidilute regimes is quantitatively known as a function of polymer concentration $\rho$ from RG calculations.\textsuperscript{25} This volume term is well understood. The problem of a quantitative description of a single colloid in a polymer solution thus reduces to understanding the second term, which defines the surface tension $\gamma_s(\rho)$, i.e., the free-energy per unit area that is directly related to the creation of the depletion layer. It is customary to relate the surface tension $\gamma_s(\rho)$ around a sphere to the surface tension $\gamma_w(\rho)$ near a planar wall, by expanding in powers of the ratio $q = R_g/R_c$

$$
\gamma_s(\rho) = \gamma_w(\rho) + \kappa_1(\rho) q + \kappa_2(\rho) q^2 + \mathcal{O}(q^3),
$$

which is expected to be most useful when $q$ is not too large. The coefficients $\kappa_i(\rho)$ control the curvature corrections. They are analogous to the Tolman corrections in the macroscopic case.\textsuperscript{33,34}

The paper is organized as follows: The case of a single plate or hard wall immersed in a polymer solution is discussed in Sec. II, where we report results for density profiles $\rho(z)$ at various polymer concentrations. These density profiles define the reduced adsorption $\hat{\Gamma}(\rho)$, from which the surface tension $\gamma_w(\rho)$ may ultimately be extracted. These considerations are extended to spherical colloids in Sec. III, where simulation results for the density profiles are reported for size ratios $q = 0.67, 1.05,$ and $1.68$. These data are then used to compute $\gamma_s(\rho)$ and the $\kappa_i(\rho)$; limiting forms are extracted for the $\rho \rightarrow 0$ and the semidilute regimes. The results are compared to the theoretical predictions for ideal polymers, for the penetrable sphere model, and wherever applicable, to RG and integral equation predictions. The limit of large $q$, where the expansion (2) becomes less useful is also discussed. For this limit we also report on some preliminary direct simulation results for $F_1$ based on the Widom insertion technique.\textsuperscript{35} Finally we show that the soft colloid paradigm has the correct thermodynamics of the single colloid problem automatically built in.

II. DENSITY PROFILES AND SURFACE TENSION NEAR A SINGLE WALL

A single hard wall in a bath of nonadsorbing polymers creates an entropically induced depletion layer because the polymers have fewer possible configurations near the wall. To calculate these density profiles we performed Monte Carlo simulations of the popular self avoiding walk (SAW) model on a cubic lattice. Even though this model ignores all chemical details of a real polymer system except the excluded volume and polymer connectivity, it reproduces the scaling behavior and many other physical properties of athermal polymer solutions.\textsuperscript{22,32} For $N$ polymers of length $L$ on a lattice of $M$ sites, the polymer density is given by $\rho = N/M$, while the monomer density is $c = LN/M$. The polymers are characterized by the radius of gyration, which scales as $R_g \sim L^{\nu}$, where $\nu \approx 0.588$ is the Flory exponent.\textsuperscript{22,25,32} For densities $\rho$ less than the overlap density $\rho^* = 1/(2\pi R_g^2)$ the system is in the dilute regime, while for $\rho \gg \rho^*$, and $c \ll 1$, the system is in the semidilute regime. We use $L = 500$ SAW polymers in our simulations, which are expected to exhibit properties close to those corresponding to the scaling limit $L \rightarrow \infty$. Further details of the simulation method and the model can be found in Refs. 18–21. Note that a small correction to these results must be applied.\textsuperscript{36}

Since our models are all athermal, we set $\beta = 1/(k_B T) = 1$.

Examples of the depletion layer density profiles near a hard wall are depicted in Fig. 1 for a polymer center-of-mass (CM) representation, as well as for a monomer representation. Both profiles have, by definition, the same reduced adsorption, defined as:\textsuperscript{34}

$$
\hat{\Gamma}(\rho) = -\frac{1}{\rho} \frac{\partial (\Omega^{CM})}{\partial \mu} = \int_0^\infty h(z) dz,
$$

where $\Omega^{CM}/A$ is the surface excess grand potential per unit area $A$. $h(z) = \rho(z)/\rho - 1$, with $\rho(z)$ the cm density profile of the polymer coil at a distance $z$ from the surface. In the monomer representation one should replace $\rho$ by $c = LN/M$ and $h(z)$ by the monomer profile; the two reduced adsorptions are equal and measure the reduction in the number of chains near the surface.

In the low-density limit an RG calculation based on a first order $\epsilon$-expansion gives $\hat{\Gamma}(0) \approx -1.074R_g$,\textsuperscript{27,37} which is slightly less than

$$
\hat{\Gamma}^{id} = 2R_g/\sqrt{\pi} \approx -1.128R_g,
$$

density independent result for an ideal polymer with the same size $R_g$.\textsuperscript{26} (but larger $L$ due to the different scaling of the radius of gyration). This reflects the fact that for a given $R_g$, the polymer–polymer interactions reduce the size of the depletion layer, an effect which becomes more pronounced with increasing density; see e.g., Fig. 1.

For the semidilute regime, de Gennes has proposed an approximate expression for the monomer profile near a wall, $h_m(z) = \tanh^2(z/\xi(\rho))-1$, where $\xi(\rho)$ is the correlation length or blob size.\textsuperscript{22} If we identify $\xi(\rho)$ with $-\hat{\Gamma}(\rho)$ then, as shown in Fig. 1, this form provides a fairly accurate fit to our simulation results. Since $\xi(\rho) \sim \rho^{-1/(3\nu-1)} \sim 0.770$ in the
semidilute regime, this implies that the density profiles should become more narrow with increasing density, a trend clearly seen in Fig. 1.

From the density profiles of Fig. 1, we can derive the adsorption at different densities using Eq. (3). These are shown in Fig. 2, together with a simple fit constrained to give the expected $r=0$ value, and the correct scaling behavior in the semidilute regime where $-\hat{\Gamma}(r) \approx \xi(r) - \rho^{0.770}$, namely

$$
\hat{\Gamma}(r) = -1.074 R_g \left[ 1 + 7.63 \frac{\rho}{\rho_s} + 14.56 \left( \frac{\rho}{\rho_s} \right)^3 \right]^{-(0.2565)}.
$$

(5)

Throughout this paper the value of the radius of gyration is conventionally chosen as that appropriate for an isolated polymer, i.e., $R_g=R_g(\rho=0)$. However, as the polymer concentration increases, the measured $R_g(\rho)$ will decrease with density as shown in Fig. 2. In the semidilute regime this scales as $R_g(\rho)/R_g(\rho=0) \sim \rho^{(2v-1)/(6v-2)} \approx \rho^{-0.115}$, which decreases much more slowly with density than the correlation length $\xi(\rho)$ or the relative adsorption $\hat{\Gamma}(\rho)$. In fact at $\rho/\rho_s=1$, the crossover from the dilute to the semidilute regimes, $\hat{\Gamma}(\rho)$ has dropped to 59% of its $\rho \to 0$ value, while $R_g(\rho)$ has only changed by a few percent. The largest rate of relative change in the adsorption is, therefore, found in the dilute regime, suggesting that theories based on the $\rho \to 0$ limit may start to break down well before the semidilute regime is reached. The border between the dilute and semidilute regimes is not sharp. For the semidilute regime, the asymptotic forms derived by scaling theories appear to be reached at a lower density for $\hat{\Gamma}(\rho)$ than for $R_g(\rho)$.

One route to calculate the surface tensions from the density profiles is to use an extension of the Gibbs adsorption equation to express the surface tension near a single wall in terms of the relative adsorption and the equation of state:

$$
\gamma_s(\rho) = -\frac{\partial \Omega^{ex}}{\partial A} = -\int_0^\rho \left( \frac{\partial \Pi(\rho')}{\partial \rho'} \hat{\Gamma}(\rho') \right) d\rho'.
$$

(6)

The derivation of this equation can be found, for example, in Refs. 24 and 38. By performing one integration by parts w.r.t. density, Eq. (6) can also be expressed as

$$
\gamma_s(\rho) = -\Pi(\rho) \hat{\Gamma}(\rho) + \int_0^\rho \Pi(\rho') \left( \frac{\partial \hat{\Gamma}(\rho')}{\partial \rho'} \right) d\rho'.
$$

(7)

The first term in this equation takes the form of a pressure times a length. For ideal polymers, where $\hat{\Gamma}(\rho)$ is independent of density, this term completely describes the surface tension of the depletion layer. It is just the (entropic) free energy cost per unit area of creating a cavity of volume $\Gamma(\rho)A$. The second (positive) term is therefore only relevant if there are polymer–polymer interactions.

We have previously calculated the equation of state for $L=500$ and $L=2000$ SAW polymers, both of which are well described by analytic RG expressions. Using this for $\Pi(\rho)$ together with the fit to the relative adsorption from Eq. (5), we can now use Eq. (7) to calculate the surface tension of a solution of polymers in good solvent near a single wall. Our results are shown in Fig. 3. In the low density limit the surface tension reduces to the same functional form as for
ideal polymers, i.e., \( \gamma_w(\rho) = -\hat{\Pi}(\rho) \hat{\Gamma}(\rho) \), so that \( \lim_{\rho \to 0} \gamma_w(\rho) \approx 1.074 R g / \rho \). Note that for all but the lowest densities, the surface tension is considerably larger than the ideal polymer result \( \gamma_w^{id}(\rho) = 2 R g / \sqrt{\pi} \). The surface tension for interacting polymers increases more rapidly with increasing density both because \( \hat{\Pi}(\rho) \) increases faster than \( \hat{\Gamma}(\rho) \) decreases in the first term of Eq. (7), and because the second term, which is absent for noninteracting polymers, increases with density as well.

Further simplifications occur in the semidilute regime. For example, when the scaling forms for the osmotic pressure, \( \Pi \sim \rho^3 r (3 r - 1) \), and for the reduced adsorption, \( \hat{\Gamma} \sim \rho^{2 r} r (3 r - 1) \), are used in Eq. (7), then the integral in the second term can be easily performed and turns out to be exactly half the first term, a result that is independent of the value of the exponent \( \nu \). The surface tension, therefore, takes on a very simple form

\[
\gamma_w^{id}(\rho) = -\frac{1}{2} \hat{\Pi}(\rho) \hat{\Gamma}(\rho) \sim \rho^{2 r} r (3 r - 1) = \rho^{1.539}.
\]

As shown in Fig. 3, this expression works remarkably well for larger densities into the semidilute regime. Deviations do occur for the dilute regime where Eq. (8) overestimates the surface tension by a factor 1.5 for \( \rho \to 0 \), as demonstrated in the inset of Fig. 3.

In a recent publication Maassen, Eisenriegler, and Bringer\(^{31} \) have used the renormalized tree approximation to derive a surface tension which compares well with our results, as shown in Fig. 3. A similar asymptotic RG \( \epsilon \)-expansion compares slightly less well. The difference between the two approximations gives an estimate of the error in the RG approach. It should be kept in mind that our simulation approach also incurs small errors through the use of the fitted form of \( \hat{\Gamma}(\rho) \), and because we use polymers of a finite length.

Fuchs and Schweitzer\(^{31} \) recently applied the polymer reference interaction site model (PRISM) approach to polymer–colloid mixtures. In the limit of low colloid density, a number of analytic results can be derived for the insertion free energy \( F_1 \), from which the surface tension can be extracted by using Eqs. (1) and (2)

\[
\gamma_{\text{PRISM}}(\rho) = 1.279 \rho R_g \left( 1 + 1.06 \frac{\rho}{\rho^*} \right).
\]

Here we have used the PRISM results arising from local packing information (see Ref. 31 for details). As can be seen in Fig. 3 this PRISM approach does not show the right scaling behavior. This is not surprising since these results are based on a simplified PRISM model with an ideal (Gaussian) description of the internal polymer correlations. This has the advantage of being analytically solvable, but the disadvantage of exhibiting the wrong scaling behavior. In the semidilute regime, this can be remedied by choosing an effective step-length,\(^{40} \) which sets the correct scaling behavior of the radius of gyration \( R_g \sim \rho^{1-(1/2)(1+2/3 \nu)} \). When this correction is applied, Eq. (9) shows the right scaling behavior in the semidilute regime, as shown in Fig. 3. In principle, a fully self-consistent PRISM calculation, which includes the correct internal polymer statistics, should result in the correct scaling without the need for an effective step-length. This would have the advantage that the surface tension would also be correctly described in the dilute regime, where the use of an effective step-length results in a spurious divergence as \( \rho \to 0 \).

### III. Density Profiles and Surface Tension Around a Hard Sphere

Having described the surface tension for a polymer solution near a single hard wall, we next investigate the related problem of a polymer solution near a single hard sphere (HS) of radius \( R_c \). As discussed in the Introduction, adding a single HS to a polymer solution reduces the number of configurations available to the polymers, and results in a finite insertion free energy or chemical potential described by Eq. (1). Besides the configurations directly excluded by the sphere of volume \( \frac{4}{3} \pi R_c^3 \), there are also configurations excluded near the surface of the sphere, an effect which manifests itself in an entropically driven depletion layer, just as was found for the case of a hard wall. However, the curvature of the sphere leads to corrections to the planar surface tension, as described by Eq. (2), i.e., the surface tension \( \gamma_s(\rho) \) depends not only on the polymer density \( \rho \), but also on \( R_c \) through the ratio \( q = R_g / R_c \).

#### A. Ideal polymers

The free energy cost of inserting a single HS into a bath of ideal polymers is known\(^{26,41} \)

\[
F_{1}^{id} = \rho \rho^* q \left( \frac{6 q}{\sqrt{\pi}} + 3 q^2 \right).
\]

By combining this result with Eq. (1), it follows that the ideal polymer surface tension takes the form

\[
\gamma_{s}^{id}(\rho) = \gamma_{s}^{id}(\rho) + \rho R_c q.
\]

The curvature corrections defined in Eq. (2) take on a particularly simple form here, since \( k_{s}^{id}(\rho) = \rho R_c \), and \( k_s = 0 \) for \( i \geq 2 \). Note that this expression is not simply an expansion in \( q = R_g / R_c \); it is valid for all size ratios.

In 1958 Asakura and Oosawa\(^{14} \) (AO) introduced a model where the ideal polymers are approximated as interpenetrable spheres of radius \( R_{AO} \). This corresponds to approximating the true depletion layer by a step-function. The free-energy of insertion of a single hard sphere into a bath of AO particles can be easily calculated to be

\[
F_1^{AO} = \eta_{AO} \frac{1}{q_{AO}} (1 + q_{AO})^3,
\]

where \( \eta_{AO} = \frac{1}{4} \pi R_{AO}^3 \) is analogous to \( \rho / \rho^* \), and we have defined the size ratio \( q_{AO} = R_{AO} / R_c \). The surface tension is, therefore, given by

\[
\gamma_{s}^{AO}(\rho) = \gamma_{s}^{AO}(\rho) + \rho R_{AO} q_{AO} + \frac{\rho R_{AO}^3}{3} (q_{AO})^2.
\]
monotonically with decreasing sphere size $R_c$. For infinite sphere size (i.e., a wall) $R_{AO}=(2/\sqrt{\pi})R_c$. As the relative sphere size $R_c/R_g$ decreases this effective parameter decreases due to the deformation of the polymers around a sphere. The symbols denote direct simulations $L=200$ ideal polymers on a lattice, taken from Ref. 12. The small differences are due to the discrete nature of the lattice used in the simulations.

Since the pressures in the two systems are the same, i.e., $\Pi = \rho$, this is equivalent to equating the two insertion free-energies $F_1$ of Eqs. (10) and (12), as done in Refs. 12 and 26. For $q=R_c/R_g \to 0$ this expression reduces to $R_{AO}^{\text{eff}}/R_g = 2/\sqrt{\pi}$, the known result for a single wall. For large $q$ on the other hand, the effective AO radius scales as $R_{AO}^{\text{eff}}/R_g \sim q^{-1/3}$. For fixed $R_g$, the effective radius $R_{AO}^{\text{eff}}$ decreases monotonically with decreasing sphere size $R_c$, as shown in Fig. 4.

B. Interacting polymers

1. Low density limit for interacting polymers

For interacting polymers, the $\rho \to 0$ limit of the curvature corrections to the surface tension have been calculated to first order in an $\epsilon$-expansion by Hanke, Eisenriegler, and Dietrich. For large spheres (small $R_c/R_g$) they find

$$\lim_{\rho \to 0} \gamma_s(\rho) = 1 + 0.849q - 0.0375q^2 + \mathcal{O}(q^3).$$

In this low density and small $q$ regime, the curvature corrections for interacting polymers are quite similar to those found for noninteracting polymers. Compare, for example, the first relative curvature correction coefficient, which is 0.849 for interacting polymers, and 0.886 for ideal polymers. In the opposite (small sphere) $q \to \infty$ limit the differences are more pronounced: $\gamma_s^{\text{id}}/\gamma_s^{\text{id}} \sim q$, $\gamma_s^{AO}/\gamma_s^{AO} - (d_{AO})^2$ while for interacting polymers RG and scaling theory approaches predict that $\gamma_s^{\text{id}}/\gamma_s^{\text{id}} \sim (q)^{1/2} \approx q^{0.701}$.

2. Interacting polymers at finite densities

We have calculated the density profiles $h(r)=\rho(r)/\rho - 1$ for polymers around spheres of radius $R_c = 1.49R_g$, $R_g = 0.95R_g$, and $R_g = 0.59R_g$ from simulations of $L=500$ SAW polymers. These are shown in Fig. 5 in the cm representation. Just as was found for the case of a planar wall, the
depletion layers shrink with increasing bulk density $\rho$. Because the polymers can deform around the colloid, the density profiles in the cm representation can penetrate into the HS region, an effect which becomes more pronounced for smaller colloids (larger $q$). (For an interesting proposal that describes the monomer density around a spherical particle we refer to Ref. 43.)

The relative adsorption around a sphere is defined as

$$4 \pi R_c^2 \Gamma_s(\rho) = \frac{4 \pi R_c^2}{\rho} \left[ \frac{\partial (\Omega^s/A_t)}{\partial \mu} \right]_{\mu=0}$$

$$= \int_0^\infty 4 \pi \rho^2 h(r) dr + \frac{4}{3} \pi R_c^3. \tag{16}$$

Here $h(r)$ is defined from the center of the sphere, and $\Gamma_s(\rho)$ has the dimension of a length. The volume of a single HS was subtracted off so that the adsorption only describes the effects of the depletion layer around a sphere. For low density the relative adsorption of a sphere is larger than that of a planar wall by a curvature correction factor term similar to those described in Eq. (15). As the density increases, the relative adsorption decreases and tends asymptotically to the same value as for a planar wall. This can be understood from the simple “blob” picture in the semidilute regime: Since the ratio of the blob-size to the sphere $\xi(\rho)/R_c$ decreases with increasing density, the curvature corrections to the relative adsorption are also expected to become relatively less important with increasing density.

The surface tension $\gamma_s(\rho)$ can now be calculated from Eq. (6) using the adsorption defined in Eq. (16). In Fig. 6, we compare the surface tension for three different sphere sizes to $\gamma_w(\rho)$, the value for a planar wall. As expected from the results for low densities, for a given density $\rho$, the surface tension increases with decreasing $R_c$. The ratio $\gamma_s(\rho)/\gamma_w(\rho)$, shown in Fig. 7, decreases with increasing $\rho/\rho^\ast$. Again, the rate of change is largest in the dilute regime; for increasing $\rho/\rho^\ast$ the two terms appear to approach each other asymptotically. Just as we argued for the adsorptions, the blob picture in the semidilute regime implies that the curvature corrections should decrease with increasing density, which is what we observe. This also implies that $\gamma_s(\rho) = \gamma_w(\rho) = \rho^{1.539}$ in the semidilute regime. Of course the smaller the HS, the higher the density one needs for the curvature corrections to become negligible. This picture is confirmed by recent scaling and RG arguments, which show that the first curvature correction coefficient $\kappa_1 \sim \rho^{6(3\nu-1)-0.770}$, implying that with increasing density, the contribution of the curvature corrections defined in Eq. (2) becomes relatively smaller, so that $\gamma_s$ approaches $\gamma_w$. In the inset of Fig. 7 we compare our results to the RG calculations, valid to lowest order in $q$, i.e., $\gamma_s(\rho) = \gamma_w(\rho) + \kappa_1(\rho) q$. Although only the results for the ratio $q = 1.05$ are shown, they are similar to those at the other two size-ratios, which also show an overestimate by the RG. The difference may be due in part to higher order $\kappa_1(\rho)$ terms which have not yet been calculated by RG. To confirm this picture further simulations are needed since: (a) our simulations of the adsorption are only for $\rho/\rho^\ast \approx 2.32$, and we extrapolated to higher densities using a fit form which scales as $-\Gamma_s(\rho) \sim \xi(\rho) \rho^{-0.770}$ at high densities; (b) we only examined three different sphere sizes so that it is difficult to directly extract $\kappa_1(\rho)$, and for that matter the higher order $\kappa_i(\rho)$.

Finally, we reemphasize how much the density dependence of the surface tension of the interacting polymers differs from that of ideal polymers or the related Asakura–Oosawa model, where the ratio of the wall to the sphere surface tensions is independent of density, and close to that of interacting polymers in the low-density limit [compare Eqs. (11), (13), and (15)].
3. Comparison with a hard-sphere model

One might inquire what would happen if the polymers were modeled as HS instead. By using the very accurate Rosenfeld fundamental measure density functional technique, an explicit form for the surface tension of a HS fluid with radius \( R_p \) around a single HS (radius \( R_c \)) has been calculated\(^{45} \)

\[
\frac{\gamma_{\text{HS}}(\eta_p)}{\gamma_w(\eta_p)} = 1 + \frac{2(1 - \eta_p)R_p}{(2 + \eta_p)R_c} - \frac{2(1 - \eta_p)^2 \ln(1 - \eta_p)}{3 \eta_p(2 + \eta_p)} \left( \frac{R_p}{R_c} \right)^2,
\]

where we have defined the packing fraction \( \eta_p = \frac{4}{3}\pi \rho_p R_p^3 \), for a number density \( \rho_p \). The planar wall-surface tension is given by

\[
\gamma_w(\eta_p) = \frac{3 \eta_p(2 + \eta_p)}{8 \pi R_c^2 (1 - \eta_p)^2}.
\]

We note that this result for the surface tension of a HS fluid around a sphere was also derived independently by scaled particle theory.\(^{46} \) Equation (17) can be generalized to the nonadditive HS model, for which the cross diameter \( R_{c,p} \neq \frac{1}{2}(R_c + R_p) \), so that one can smoothly interpolate between the fully repulsive HS model and the fully nonadditive AO model.\(^{45,47} \)

To lowest order in density, the surface tension of a HS system near a planar wall is \( \gamma_w \approx R_p \rho_R \), i.e., the same as that of the AO model, which is close to that of interacting polymers in the same limit where \( \gamma_w \approx 1.074 R_p \rho_R \). However, the terms of order \( \rho^2 \) are already significantly larger in the HS case. Therefore, as illustrated in the inset of Fig. 6, the HS model gives a large relative overestimate of the surface tension well before reaching the packing fraction at which the system freezes. (Here we took units where \( R_p = R_g \) so that \( \eta_p \) is equivalent to \( \rho / \rho^* \).)

For a fixed size-ratio \( R_p / R_c \), the curvature corrections for a HS system vary with density as

\[
\frac{\gamma_{\text{HS}}(\eta_p)}{\gamma_w(\eta_p)} = \frac{\gamma_{\text{AO}}}{\gamma_w} - \frac{3 R_p^2}{2 R_c^2 + 3 R_p^2} \eta_p + O(\eta_p^2),
\]

where the ratio for the AO model comes from Eq. (13) with \( R_{\text{AO}} = R_p \). As illustrated for a 1:1 size ratio in the inset of Fig. 7, for small \( \eta_p \) this ratio is indeed almost linear. The change with density is more pronounced than that found for a polymer–colloid system with a similar size-ratio, suggesting (not surprisingly) that a full HS system is not such a good model of interacting polymers, even at relatively low densities. Making the spheres nonadditive does not fundamentally alter this picture—the behavior of polymers falls into the class of “mean-field fluids,” i.e., they do not behave like hard-core particles.

4. Direct calculation of \( F_1 \) by the Widom insertion method

We also performed direct computer simulations of the free energy \( F_1 \) by measuring the insertion probability of a single sphere in a bath of polymers at fixed density \( \rho / \rho^* = 1.16 \). This is closely related to the so-called Widom insertion technique to find the chemical potential.\(^{35} \) Figure 8 shows that \( F_1 \) grows with increasing sphere size as expected. The same is true for the contribution due to the depletion layer, i.e., the contribution proportional to \( 4 \pi \gamma R_g^2 \) in Eq. (1). However, the relative importance of this surface tension term increases with \( \gamma \). Using the adsorption method was used. We also compare this ratio is indeed almost linear. The planar wall-surface tension is given by

\[
\gamma_{\text{w}}(\eta_p) = \frac{3 \eta_p(2 + \eta_p)}{8 \pi R_c^2 (1 - \eta_p)^2}.
\]

We note that this result for the surface tension of a HS fluid around a sphere was also derived independently by scaled particle theory.\(^{46} \) Equation (17) can be generalized to the nonadditive HS model, for which the cross diameter \( R_{c,p} \neq \frac{1}{2}(R_c + R_p) \), so that one can smoothly interpolate between the fully repulsive HS model and the fully nonadditive AO model.\(^{45,47} \)

To lowest order in density, the surface tension of a HS system near a planar wall is \( \gamma_w \approx R_p \rho_R \), i.e., the same as that of the AO model, which is close to that of interacting polymers in the same limit where \( \gamma_w \approx 1.074 R_p \rho_R \). However, the terms of order \( \rho^2 \) are already significantly larger in the HS case. Therefore, as illustrated in the inset of Fig. 6, the HS model gives a large relative overestimate of the surface tension well before reaching the packing fraction at which the system freezes. (Here we took units where \( R_p = R_g \) so that \( \eta_p \) is equivalent to \( \rho / \rho^* \).)

For a fixed size-ratio \( R_p / R_c \), the curvature corrections for a HS system vary with density as

\[
\frac{\gamma_{\text{HS}}(\eta_p)}{\gamma_w(\eta_p)} = \frac{\gamma_{\text{AO}}}{\gamma_w} - \frac{3 R_p^2}{2 R_c^2 + 3 R_p^2} \eta_p + O(\eta_p^2),
\]

where the ratio for the AO model comes from Eq. (13) with \( R_{\text{AO}} = R_p \). As illustrated for a 1:1 size ratio in the inset of Fig. 7, for small \( \eta_p \) this ratio is indeed almost linear. The change with density is more pronounced than that found for a polymer–colloid system with a similar size-ratio, suggesting (not surprisingly) that a full HS system is not such a good model of interacting polymers, even at relatively low densities. Making the spheres nonadditive does not fundamentally alter this picture—the behavior of polymers falls into the class of “mean-field fluids,” i.e., they do not behave like hard-core particles.

\[
F_1 = A_g R_g^{d-1/v} \rho R_g^{1/v}.
\]

Where \( A_g \) is a universal numerical pre-factor that can be calculated from an RG technique.\(^{27,28} \) For \( d = 3 \) Eq. (20) reduces to \( F_1 \approx 18.4 \rho R_g^{1.30} R_g^{1.70} \). This expression is directly compared to our simulations in Fig. 8. By comparing to Eq. (1) we can extract the surface tension from the insertion free energies. This was done for the \( F_1 \) at \( \rho / \rho^* = 1.16 \) shown Fig. 8, and also for \( L = 2000 \) polymers at \( \rho / \rho^* = 0.94 \). Using the longer polymers allows effectively smaller colloidal \( R_c \)’s to
be used in our lattice simulations. The surface tensions are depicted in Fig. 9. At small $q$ our computer simulation results correspond reasonably well with the asymptotic RG results. We expect there to be small errors due to the discreteness of our lattice simulations, similar to those depicted for ideal polymers in Fig. 4. These discretization errors become more important as the spheres become relatively smaller. We have made some small corrections\textsuperscript{50} to take this into account, but a more systematic study, possibly with longer polymers, would be necessary to completely test the RG results.

When the colloids are much smaller than the polymers, one expects that they only probe the local monomer density, and not the overall number density of polymer coils. In fact, Eq. (20) implies just that since $F_1 \approx \rho R_g^{1.30} / R_c - \rho L R_c$. The reason $F_1$ scales linearly with the monomer density $c$ is that by definition this is very small ($c \ll 1$) in the dilute and semidilute regime. The small colloidal particles probe what is effectively an ideal gas of monomers.

For ideal polymers $F_1 \approx c R_c$ in the limit of small $R_c$, which implies that for a given $R_g$ and $\rho$, and for a small enough $R_c$, it is easier to insert a hard-sphere into a bath of interacting polymers than it is to insert it into a bath of noninteracting polymers. At first sight this may seem surprising, but the reason is as follows: Inside an interacting polymer, the monomer concentration scales as $c \sim R_g^{-1.30}$ while for ideal polymers it scales as $c \sim R_g^{-1}$. In other words, the interactions swell a polymer and make it less dense; for a given $R_g$, the monomer density $c$ is larger for ideal polymers than for interaction polymers, and since the small colloids only probe the local monomer density it is easier to insert the sphere into an interacting system than into a noninteracting system at the same $R_c$. This effect is illustrated in Fig. 8 for $\rho/\rho^* = 1.16$, where the crossover is at about $R_c \approx 0.5 R_g$.

This analysis should not be confused with a comparison at fixed $L$. Note that the PRISM results also overestimate $F_1$ at small $R_c$. This is in part because the simplified PRISM model we compare to also includes ideal polymer statistics, resulting in an overestimate of the monomer density compared to a true interacting system, an effect already pointed out in Ref. 31.

For large spheres, on the other hand, where $F_1 \approx \frac{1}{2} \pi R_g^2 (\rho R_g^1)$ which scales as $F_1 \approx \rho^{2.30} R_g^3$ in the semidilute regime, the spheres do directly probe the number density of polymer coils, and the insertion free energy for interacting polymers is always higher than that of ideal polymers at the same $R_g$ and $\rho$. Note how differently the large and small $R_c/R_g$ limits of $F_1$ scale both with $\rho$ and with $R_g$. Significant differences can also be found for the scaling of the surface tensions since for large $R_c/R_g$, $\gamma \sim \gamma_0(\rho)(1 + C (R_g/R_c)) \sim \rho^{1.30} (1 + C (\rho^{-0.70} R_g/R_c))$, while for small $R_c/R_g$, the RG expressions imply that $\gamma \sim c R_c^{-0.7}$.

**C. Surface tension for polymers as soft colloids**

We have recently modeled polymers as single soft colloids interacting with a pair potential between their cm.\textsuperscript{18–21} These pair potentials were derived by a liquid state theory based inversion procedure such that the soft colloids have exactly the same radial distribution function $g(r)$ as those generated by a fully interacting polymer simulation. A similar inversion procedure was used to derive the potential between the soft colloids and a planar wall or a HS. These wall–polymer or sphere–polymer potentials are such that they exactly reproduce the one-body density profiles $\rho(r)$.

Since our effective polymer–polymer potentials provide a very accurate representation of the pressure $\Pi(\rho)$,\textsuperscript{19,21} while the polymer–wall or polymer–sphere interactions are constrained to reproduce the correct density profiles, and therefore the correct adsorption $\Gamma(\rho)$, Eq. (6) implies that our soft-colloid approach has the correct surface tensions automatically built in. Similarly Eq. (1) implies that this approach correctly reproduces $F_1$ for a sphere immersed in a polymer solution.

**IV. CONCLUSIONS**

In summary then, we have used computer simulations of SAW polymers on a cubic lattice to calculate the density profiles for nonadsorbing polymers near a planar wall, and near HS’s. From this we were able to calculate and fit the relative adsorption $\Gamma(\rho)$. Together with the equation of state, which is well understood for polymer solutions, this provides the needed ingredients to calculate the surface tensions through Eq. (6).

The surface tension of interacting polymers near a planar wall was shown to differ significantly from that of ideal polymers, or other simple models such as the Asakura–Oosawa penetrable-sphere model, or a pure HS fluid. Similarly, a recent PRISM calculation\textsuperscript{31} also shows large qualitative differences with our results, which could have been anticipated in view of its use of simplified ideal polymer statistics. However, if an effective step-length is used, the
correct scaling behavior is obtained in the semidilute regime. On the other hand, some recent RG results\textsuperscript{29} compare very well to our calculations. In the semidilute regime, the surface tension simplifies to the form given in Eq. (6), which implies that $\gamma_s (r) \sim r^{-0.770}$ in the semidilute regime. This is again consistent with some recent RG calculations of the correction coefficient $\kappa_1 (r)$,\textsuperscript{29} although further simulations are needed to confirm the scaling and form of the coefficients $\kappa_2 (r)$.

For smaller colloids, it is advantageous to use a direct Widom insertion technique to calculate the free-energy $F_1$. For very small colloids (large $q$), our simulations were consistent with asymptotic RG results which suggest that $F_1 \propto c q^{1/3}$. This insertion free-energy is dominated by the contribution of the depletion layer; its behavior is qualitatively different from the behavior found at smaller $q$, and suggests that the expansion of Eq. (6) breaks down for large $q$.

Because our soft-colloid approach was derived to reproduce the correct one-particle density profiles near hard walls or hard-spheres, it will automatically reproduce the correct adsorptions, and therefore also the correct surface tensions and insertion free-energies $F_1$.

The walls and spheres in this study are purely repulsive. Adding a wall–polymer attraction should decrease the amount of depletion, and therefore, also lower the surface tensions. More subtle effects could be expected if in addition the solvent quality is decreased. New effects are also expected for binary mixtures of polymers with selective adsorption of one of the species. These systems will be the subject of future investigations.

The next step is to move from the one-sphere or one-wall problem to the case of a two-sphere or a two-wall system, and calculate the effective interactions between the two particles. This is the subject of a forthcoming paper.\textsuperscript{51}

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\textsuperscript{36}In our earlier work (Refs. 18–20) we used $R_g = 16.5$ instead of the correct $R_g = 16.83$ value for the $L = 500$ SAW polymers. This means a slight adjustment in the values of $R_g$ and in $\rho_p^{*}$ quoted in our papers. For example, the values quoted for $\rho_p^{*}$ should increase by a factor 1.06.
\textsuperscript{37}To compare with (Ref. 27), where results are expressed in terms of $R_g$, the end-to-end radius, we have used $R_g / \rho_p^{*}$. $R_g$, the value expected for interacting polymers (Ref. 32). This is close to the ideal-polymer result $R_g = 16.83$ value for the $L = 500$ SAW polymers, a procedure similar to that used when comparing to experiments (Ref. 32).
where the measured $B_2$ is used to set the fit parameter. In principle one can also extract $B_2$ from an RG calculation, so that the fit is not really necessary in the scaling limit (Ref. 25). However, we kept the fit form to make our results self-consistent for $L = 500$ SAW polymers.

41 P. B. Warren, private communication.
47 To correctly obtain the fully nonadditive AO limit, the explicit $R_p$ in Eq. (17) should be equated with $R_{AO}$, while the packing fractions $\eta_p \rightarrow 0$.
50 In the simulation we insert the colloid with its center a lattice site. We then check all the sites that are within the radius for overlap with one of the polymers. The number of these lattice sites, i.e., the volume occupied by the sphere can be transformed into an effective radius. This radius takes the discreteness effects partially into account, and is used in the analysis.
51 A. A. Louis, P. G. Bolhuis, E. J. Meijer, and J. P. Hansen, preprint cond-mat/0203144.