Effective non-additive pair potential for lock-and-key interacting particles: the role of the limited valence.

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Theoretical studies of self-assembly processes and condensed phases in colloidal systems are often based on effective inter-particle potentials. Here we show that developing an effective potential for particles interacting with a limited number of “lock-and-key” selective bonds (due to the specificity of bio-molecular interactions) requires — besides the non-sphericity of the potential — a (many body) constraint that prevent multiple bonding on the same site. We show the importance of retaining both valence and bond-selectivity by developing, as a case study, a simple effective potential describing the interaction between colloidal particles coated by four single-strand DNA chains.

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Description of multi-scale phenomena in self-assembling systems and understanding of their collective behavior requires a coarse graining process in which the irrelevant degrees of freedom are integrated out in favor of an effective re-normalized interaction potential. Colloidal systems are often described by this type of approach, in which the solvent and its chemical-physical properties are encoded into parameters of the resulting effective inter-colloid particle potential. Thermodynamic (and often also dynamic) bulk behavior of several complex particles, including polymers, star-polymers, dendrimers, micelles, telecheric polymers has been interpreted on the basis of effective pair-wise potentials, function only of the distance between particle centers.

Very recently, design of new materials has started to capitalize on the possibilities offered by complementary “lock-and-key” selective interactions, aiming to achieve a much better control on the resulting bonding pattern and on the mechanical properties of the self-assembled bulk material. In this Letter we show that effective potentials for these new particles must account for both the valence (maximum number of bonds) and for the selectivity of the bonding process. While at the microscopic level specificity in the bonding interactions is pair-wise additive — it builds on steric incompatibilities, e.g. on excluded volume effects which prevent the possibility of multiple interactions at the same bonding site — we find that the conservation of this important feature at the level of the coarse-grained effective potential may require the breakdown of the pair-wise additive approximation.

As a working example, we discuss the development of an effective potential for a macromolecule composed by a core particle decorated by a small number of short single strands (ss) DNA to show (i) the importance of properly accounting for valence and bond selectivity; (ii) the need for a non-pair-wise additive effective potential. The full model is graphically explained in Fig. 1 (and the chosen length and energy units), and more in detail Ref. [12, 13]. A semi-rigid chain of eight particles, modeling a 8-base ss-DNA, is attached to each of the four particles composing a central tetrahedral core. Each of the 32 base particles carries an additional interaction site (labeled A,T,G or C) which can bind only with complementary site types (only A-T or G-C binding are allowed), resulting in a total of 68 interaction sites per macromolecule. The chosen base sequence is palindromic, so that all eight bases can be simultaneously paired when ss-arms of different macromolecules approach in the correct orientation. All site-site interactions are pair-wise additive. The phase diagram of this model shows a region of gas-liquid phase separation, but only at small number densities \( \rho \). The liquid region is characterized by the presence of a four-coordinated network, whose dynamics progressively slows down on cooling, parallel to the formation of an open gel-like structure.

To evaluate the effective potential we simulate, using Monte Carlo methods, a system composed of two macromolecules in a box for several temperatures \( T \), covering the \( T \)-region where ss-DNA pairs. From a large ensemble of equilibrium configurations we extract the properties of the degrees of freedom to be retained in the coarse-grained model: (i) the normalized probability \( h_r \) of finding the two particles at relative center-to-center distance \( r \) (i.e. in such a way that \( \frac{1}{4} \int \pi J \approx 1 \)) (ii) the normalized probabilities \( h_{\theta_1} \) and \( h_{\theta_2} \) of the angle \( \theta_1 \) and \( \theta_2 \) between \( r \) and the closest ss-DNA arm on each particle (so that \( \frac{1}{2} \int_0^\pi h_{\theta \sin\theta d\theta} = 1 \)). The variables \( r \), \( \theta_1 \) and \( \theta_2 \) are defined in Fig. 1. The \( T \)-dependence of \( h_r(r) \) and \( h_{\theta}(\theta) \) is shown in Fig. 2. On cooling, angular and spatial correlations build up, reflecting the formation of a stable double-strand configuration, with a characteristic center-to-center distance of the order of the DNA arm length (8.5 \( \lesssim r \lesssim 10.5 \)) and a preferential orientation for linear bonding (\( \theta_1 \approx 0 \)). At the highest and lowest investigated \( T \), \( h_r \) and \( h_{\theta} \) describe, to a good approximation, the fully non-bonded (\( nb \)) and fully bonded (\( b \)) state, respectively. The non-bonded high \( T \) state is characterized by essentially unstructured \( h_{\theta}^{nb} \) and \( h_{\theta}^{nb} \), except for the \( r \)-region close to the origin where excluded volume interactions play a role. The function \( h_{\theta}^{b} \) well described by a gaussian distribution \( h_{\theta}^{b} = \frac{A}{\sqrt{2\pi\sigma_2}} e^{-\frac{\theta^2}{2\sigma_2^2}} \) (where the normalization constant is \( A = 22.32 \) and the best-fit \( \sigma^2 = 0.052 \text{ rad}^2 \)).
A similar angular distribution has been recently chosen to model self-assembly of patchy particles\textsuperscript{14}. The distribution $h_b^\theta$ is essentially localized only between $8.5 \lesssim r \lesssim 10.5$, corresponding to the bonded distance. We have not been able to provide a simple analytic best-fit expression for $h_b^\theta$ and $h_{nb}^\theta$ and, in the following, we have used their numerical values. The sharp separation in configuration space between the bonded and the non-bonded states (due to the localized nature of the interaction and to its specificity) suggests to describe the distributions at intermediate $T$ as a linear combination of the $b$ and $nb$ distributions, weighted respectively by the probability of being bonded ($p_b(T)$) or non-bonded ($1-p_b(T)$). Lines in Fig.\textsuperscript{2} show that both $h_r$ and $h_{\theta}$ can indeed be described as linear combinations (with the same $p_b(T)$ coefficient) of the $b$ and $nb$ distributions. Moreover, as shown in the inset of Fig.\textsuperscript{2} the resulting $T$-dependence of $p_b(T)$ is very well fitted by the two-state expected theoretical expression $p_b(T) = 1/(1 + e^{-\Delta U/k_B T})$, where $\Delta U$ and $\Delta S$ measure the change in energy and entropy associated to the formation of a double strand. The important result is that the $T$-dependence decouples from the $r$ and $\theta$ dependence and it becomes possible to write the probability of finding the two macromolecules at relative distance $r$ and angles $\theta_1$ and $\theta_2$ as

$$P(r, \theta_1, \theta_2, T) = p_b(T) P^b(r, \theta_1, \theta_2) + (1-p_b(T)) P^{nb}(r, \theta_1, \theta_2).$$

We approximate $P^b(r, \theta_1, \theta_2) = h_b^r(r) h_b^\theta(\theta_1) h_b^\theta(\theta_2)$ and analogously $P^{nb}$ for the non-bonded case. In this approximation, correlation between distances and angles within the $b$ (or the $nb$) state are neglected, since both $P^b$ and $P^{nb}$ are written as a product of their arguments. We have tested that this is indeed a good approximation. Correlation between angles and distances arising from the differences between $b$ and $nb$ states (small $\theta_1$ and $\theta_2$ are found when $r$ is within the bonding distance) are retained in $P$, which indeed can not be factorized.

Thus the four functions $h_b^r, h_b^\theta, h_{nb}^r, h_{nb}^\theta$ and $p_b(T)$ fully specify $P(r, \theta_1, \theta_2, T)$ and make it possible to model the macromolecule as a particle decorated by four tetrahedrally oriented arms and evaluate an effective potential between these particles. The simplest approach is to neglect the angular correlation and develop a spherical ($s$)
The disagreement is so large that no state points of liquid-gas instability, which includes the region where phase separation is expected for the model studied in Ref. [13] (estimates based on the extrapolated behavior of the low wavevector structure factor). Filled squares, outside the dashed area, denote the stable state points studied in Ref. [13]. The graph also shows the boundary between homogeneous and phase-separated states for $V_s$ and $V_{ns}$. Note that most of the filled squares are located in a region of the phase-diagram where phase separation is observed in $V_s$ or $V_{ns}$. The filled circles indicate the gas-liquid coexistence for the valency constrained effective model.

FIG. 4: Phase diagram of the model for colloidal particles coated by four single-strand DNA chains. The dashed (red) area approximately indicates the region where phase separation is expected for the model studied in Ref. [13].

FIG. 3: (a) Shape of the non-spherical effective potential $\beta V_{ns}(r)$ for given angles $\theta_1 = 0$ and $\theta_2 = 0$, and (b) the spherically averaged effective potential $\beta V_s(r)$, for different temperatures.

The radial shape of the two resulting potentials is shown in Fig. 3. $V_{ns}$ has a deeper minimum than $V_s$, to compensate for the reduction of the solid angle associated to bonding. To assess the ability of the two potentials in reproducing the bulk behavior of the original model and to estimate the importance of the directional interactions, we compare the phase diagram of the two potentials, with the (previously studied) behavior of the original potential [12 13]. Both $V_s$ and $V_{ns}$ perform very badly, predicting a very wide region of liquid-gas instability, which includes the region where the tetrahedral network of bonded particles is known to be the stable phase (see Fig. 4). The unstable gas-liquid region is even wider in the case of $V_{ns}$, due to its deeper minimum. The disagreement is so large that no state points can be chosen to compare the original and the effective models in the region where a network develops, since $V_s$ and $V_{ns}$ always generate a phase-separated configuration. The failure in reproducing the bulk behavior can be ascribed to the absence, in both $V_s$ and $V_{ns}$, of the lock-and-key character of the interaction. In the real model, once a ss-DNA is linked to a complementary strand of a different particles, it is not available for further bonding. The specificity of the interaction implies that each ss-DNA can not be simultaneously bonded to more than one other arm. This important rule is not enforced neither by $V_s$ (which allows for a large number of bonded neighbors) nor by $V_{ns}$ since the width of the angular part of $V_{ns}$ does not prevent multiple bonding at the same arm direction. We are forced to abandon the pair-wise additivity approximation and develop a new potential which enforces the bond selectivity and the valence (so that each particle can not be engaged in more than four bonds). To this aim, we complement $V_{ns}$ with an ad-hoc rule imposing that each arm can be engaged only in one bond, very much in the same spirit as the maximum valence models [13]. From a computational point of view, during the simulation, we retain a list of all pairs of bonded arms and make sure that each arm is involved at most in one bonded interaction. The resulting effective valency-constrained non-spherical model is able to generate homogenous equilibrium states in the expected region of the phase diagram. This opens the possibility to compare the predictions of the effective model with those of the original model [13], at the same $T$ and $\rho$. A comparison in real and in reciprocal space is shown in Fig. 5. The limited-valence effective model is able to reproduce with satisfactory accuracy the structure of the
tetrahedral network: both the peculiar ratio between the position of the first and second peak in the center-center radial distribution function $g(r)$ as well as the development of a pre-peak in the corresponding structure factor $S(q)$ on cooling are well captured by the effective potential. To locate precisely the gas-liquid coexistence for the limited-valence effective potential we perform umbrella sampling grand canonical Monte Carlo simulations \cite{16} (and histogram re-weighting techniques \cite{17}). The resulting gas-liquid coexistence curve is shown in Fig. 4. As compared to the unconstrained $V_s$ and $V_{ns}$, the region of gas-liquid instability is now confined to small $\rho$, opening up a window of densities (above the liquid coexistence branch) in which equilibrium liquid states can be accessed.

In summary, we have shown that in the process of developing effective potentials for particles with low valence and with selective interactions (as in functionalized colloids) it is mandatory to account for the specificity of the interaction which intrinsically limit the maximum number of bonds that can be formed. The present study shows that colloidal particles with specific interactions constitute another class of colloidal systems for which pair additivity approximation breaks down in the effective potential, in addition to the recently reported case of charge-stabilized colloidal crystal \cite{18}. The importance of retaining, beside the directionality, the valence of the original model is clearly evidenced by the fact that valence strongly controls the width of the gas-liquid coexistence \cite{19,20}.

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