Screening of heterogeneous surfaces: charge renormalization of Janus particles

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Received 23 September 2009, in final form 13 October 2009
Published 23 February 2010
Online at stacks.iop.org/JPhysCM/22/104104

Abstract
Nonlinear ionic screening theory for heterogeneously charged spheres is developed in terms of a mode decomposition of the surface charge. A far-field analysis of the resulting electrostatic potential leads to a natural generalization of charge renormalization from purely monopolar to dipolar, quadrupolar, etc, including ‘mode couplings’. Our novel scheme is generally applicable to large classes of surface heterogeneities, and is explicitly applied here to Janus spheres with differently charged upper and lower hemispheres, revealing strong renormalization effects for all multipoles.

(Some figures in this article are in colour only in the electronic version)

1. Introduction
The past few years have seen an explosion of newly synthesized colloidal (nano)particles that are not spherically symmetric, either by shape (e.g. dumbbells, snowmen, cubes) or by surface pattern (patches, stripes) [1]. A broken rotational symmetry also occurs when particles are adsorbed to air–water or oil–water interfaces [2], as in for example, colloidal monolayers [3, 4], Pickering emulsions [5], or bijels [6]. Moreover, recent atomic force microscopy studies have shown that even supposedly homogeneous colloidal surfaces can actually be heterogeneous on length scales as large as 100 nm [7], while atomic corrugations and facets render any nm-sized particle strictly heterogeneous. An important consequence of surface heterogeneity is anisotropy of the mutual effective forces, which directly affects the self-assembly process of the (nano)colloids into large-scale structures, for instance into ill-understood linear chains [4, 7, 8].

A fundamental problem is thus to establish relationships between shape and surface heterogeneity on the one hand and effective interactions and large-scale self-assembly structures on the other [1]. Apart from specific forces (e.g., hydrophobic, van der Waals) the effective interactions between dispersed particles often involve a strong generic electrostatic component, which is well described, for homogeneously charged objects, by linear screening theory provided renormalized charges instead of bare charges are used [9]. Renormalization of heterogeneously distributed surface charge is an open problem, for which we develop a systematic theory in this paper. We go beyond recent linear screening treatments [11] and formulate a new and efficient framework for computing nonlinear ionic screening effects of heterogeneously charged spheres dispersed in a 1:1 electrolyte. Our theory generalizes Alexander’s notion of ion-condensation induced charge renormalization [9] to include not only the monopole but also the dipole, quadrupole, etc, as well as their nonlinear couplings. These multipole modes can be important if one wishes to calculate the electrostatic force between particles, which was already shown for clay platelets [10]. Our scheme is versatile and can be applied to essentially any type of charge heterogeneity. We focus on applications to Janus spheres composed of two differently charged hemispheres [12].

We consider an index-matched suspension of \(N\) colloidal spheres of radius \(a\) in a bulk solvent of dielectric constant \(\varepsilon\) and volume \(V\) at temperature \(T\). The solvent also contains point-like monovalent cations (charge +\(e\)) and anions (charge −\(e\)) at fugacity \(\rho_s\). Here \(e\) is the elementary charge. A relatively simple treatment of this many-body problem is the cell model [9], in which a single colloid is considered in the centre of a spherical cell of radius \(R\) and volume \((4\pi/3)R^3 = V/N\). We denote the surface charge density of this central...
colloidal particle by $e o (\theta, \varphi)$, where $\theta$ and $\varphi$ are the standard polar and azimuthal angle, respectively, with respect to a laboratory frame. Within a mean-field approximation, the concentration profiles of the cations and anions can be written as Boltzmann distributions $\rho_+ (r) = \rho_+ e^{\exp [±\Phi (r)]}$, where $k_B T \Phi (r)/e$ is the electrostatic potential at $r = (r, \theta, \varphi)$, with $k_B$ the Boltzmann constant and $r = |r|$. Note that $\Phi (r) = 0$ in the salt reservoir, and that $\rho_+ (r) = 0$ for $r < a$ due to hard-core exclusion. The potential must satisfy the Poisson equation $\nabla^2 \Phi (r) = -4\pi \lambda_B (\rho_+ (r) - \rho_- (r))$, where we defined the Bjerrum length $\lambda_B = e^2/k_B T$. Combining the Poisson and Boltzmann equations gives

$$\Phi'' (r) + \frac{2\Phi' (r)}{r} - \frac{\mathcal{L}^2 \Phi (r)}{r^2} = \begin{cases} 0 & r < a; \\ \kappa^2 \sinh \Phi (r) & r > a, \end{cases}$$

(1)

where a prime denotes a radial derivative, $\mathcal{L}^2 = -[\sin \theta]^{-1} \partial_\theta \sin \theta \partial_\theta + [\sin \theta]^{-2} \partial_\varphi^2$ the angular momentum operator, and $\kappa^{-1} = (8\pi \lambda_B \rho_+)^{-1/2}$ the screening length. On the colloidal surface, $r = a$, Gauss’ law imposes the boundary condition (BC)

$$\lim_{r \to a^+} \Phi' (r, \theta, \varphi) = \lim_{r \to a^-} \Phi' (r, \theta, \varphi) - 4\pi \lambda_B \sigma (\theta, \varphi).$$

(2)

Electro-neutrality of the cell imposes $\int d\varphi \sin \theta \partial_\theta \tan \theta \Phi' (r, \theta, \varphi) = 0$ at $r = R$, which is a sufficiently stringent BC to close the system of equations in the spherically symmetric case. Now, however, an additional BC is to be specified for the angular dependence at $r = R$, depending on the environment of the cell. For now, we assume an environment that is characterized by ‘isotropic’ boundary conditions,

$$\partial_\theta \Phi (R, \theta, \varphi) = \partial_\varphi \Phi (R, \theta, \varphi) = 0.$$  (3)

We will discuss this choice, and its consequences, in section 4.

2. Theory

For a given $\sigma (\theta, \varphi)$ one can solve (1) with BCs for $\Phi (r)$, for example, numerically on a discrete $(r, \theta, \varphi)$ grid. The approach we take, however, avoids a cumbersome two-dimensional grid in favour of a systematic expansion of the angular dependence in spherical harmonics. For notational convenience and illustration purposes we restrict attention here to $\varphi$-independent cases where the expansion involves only Legendre polynomials $P_l (x)$ with $x = \cos \theta$.

The first step in this analysis is the decomposition of the colloidal surface charge into surface multipoles $\sigma_\ell = \frac{2\pi}{4\ell + 1} \int_0^1 \sin \theta d\theta \sin \theta \sigma (x) P_\ell (x)$, such that $\sigma (x) = \sum_{\ell = 0}^\infty \sigma_\ell P_\ell (x)$. Similarly we decompose $\Phi (r, x) = \sum_{\ell = 0}^\infty \Phi_\ell (r) P_\ell (x)$. With $\Phi_0 (r) = (r/a)^4 \Phi (a)$ the regular solution to (1) for $r \in [0, a]$, the BCs for $r \in [a, R]$, $\Phi_0' (a) = \frac{b}{a} \Phi_0 (a) - 4\pi \lambda_B \sigma \ell \quad (\ell \geq 0);$  

(4)

$$\Phi_0 (R) = 0 \quad \text{and} \quad \Phi_\ell (R) = 0 \quad (\ell \geq 1),$$  (5)

conveniently decouple for the different $\ell$’s. By contrast, the nonlinear sinh term in (1) induces ‘mode coupling’ between all Legendre components $\Phi_\ell (r)$—not to be confused with mode couplings in dynamical slowing down. This coupling is obviously impractical for a numerical treatment.

The second step of our analysis resolves this mode coupling problem by ‘ordering’ the modes systematically. We introduce a dimensionless ‘switching’ parameter $A$, and consider the auxiliary distribution $\sigma^{(A)} (x) = \sum_{\ell = 0}^\infty A^2 \sigma_\ell P_\ell (x)$, such that $A = 0$ describes a homogeneous distribution and $A = 1$ the heterogeneous one of interest. We also define the corresponding auxiliary potential $\Phi^{(A)} (r, x) = \sum_{\ell = 0}^\infty A^2 \Phi_\ell (r, x)$, where $L$ sets the order of the truncation and where the expansion coefficients can themselves be expanded as $\Phi_\ell (r, x) = \sum_{x = 0}^\infty f_{\ell x} (r) P_\ell (x)$. The functions $f_{\ell x} (r)$ are independent of $A$ and will be calculated numerically below for the cases of interest $n \geq \ell$ (which is assumed implicitly from now on). Since the problem is invariant under the simultaneous transformation $x \to -A$ and $x \to -x$ one checks that $f_{\ell x} (r) = 0$ for $n + \ell + 1$, odd, that is, we only consider $\ell$ and $n$ both even or both odd.

Replacing $\sigma (x)$ by $\sigma^{(A)} (x)$ and $\Phi (r, x)$ by $\Phi^{(A)} (r, x)$, inserting the corresponding expansions into the BCs, and equating all orders of $A$ yields at $r \downarrow a$ and $r = R$

$$f_{\ell n} (a) = \frac{\ell}{a} f_{\ell 0} (a) - 4\pi \lambda_B \sigma_\ell \delta_{\ell n};$$

(6)

$$f_{\ell 0} (R) = 0 \quad (\ell = 0) \quad \text{and} \quad f_{\ell n} (R) = 0 \quad (\ell \geq 1),$$

(7)

where $\delta_{\ell n}$ is the Kronecker delta. When the same replacement and expansion procedure is applied to the PB equation (1), one finds upon expanding the argument of the sinh term with respect to $A$ a hierarchy of second-order differential equations for $f_{\ell n} (r)$, with a structure that allows for an order-by-order sequential solution. For $n = \ell = 0$ we obtain for $r \in [a, R]$ the spherically symmetric nonlinear PB equation in the cell,

$$f_{\ell 0} (r) = \frac{9\pi}{8} f_{\ell 0} (a) \quad (\ell = 0),$$

(8)

where $S_{\ell 0} (r)$ acts as a source term of the form $S_{11} = 0$, $S_{22} = \frac{1}{2} f_{11} (r) \sinh f_{00} (r) + \frac{1}{2} f_{22} (r) \sinh f_{00} (r)$, and explicit expressions for higher-order terms can be generated straightforwardly. The key observation is that $S_{\ell 0} (r)$ only depends on $f_{10} (r)$’s with $n < \ell$, that is, a hierarchy of terms follows spontaneously. Thus (8) with the BCs (6) and (7) can be solved for $n = \ell = 1$, which in turn determines $S_{20} (r)$ and $S_{22} (r)$ such that $f_{20} (r)$ and $f_{22} (r)$ can be solved, etc. The nonlinear mode coupling, represented explicitly by $\cos f_{00} (r)$ and $S_{\ell 0} (r)$, renders the linear equation (8) highly nontrivial, yet numerical solution on a radial grid $r \in [a, R]$ is straightforward. With $f_{\ell n} (r)$ determined for $L \geq n \geq \ell \geq 0$ for some cut-off $L$, we can set $A = 1$ to explicitly construct the potential of interest $\Phi (r, x) = \sum_{\ell = 0}^L \Phi_\ell (r) P_\ell (x)$ with $\Phi_\ell (r) = \sum_{x = 0}^\infty f_{\ell x} (r)$.
3. Results

The theory developed so far is directly applicable to any uniaxial charge distribution, while generalizations to azimuthal dependencies and nonspherical shapes are feasible. In this paper we illustrate our scheme for the prototype heterogeneous charge distribution of Janus spheres characterized by surface charge densities $\sigma_N$ and $\sigma_S$ on the northern ($x > 0$) and southern ($x < 0$) hemisphere, respectively [12, 13]. The non-vanishing modes are thus $\sigma_0 = (\sigma_N + \sigma_S)/2$ and $\sigma_\ell = g\sigma_0(2\ell + 1)^{-1}2^{(\ell-2)/2}\Gamma((\ell+1)/2)$ for $\ell$ odd. Here we defined the dimensionless heterogeneity parameter

$$\gamma = \frac{\sigma_N - \sigma_S}{\sigma_N + \sigma_S}, \quad (9)$$

which together with the total charge $Z_0 = 4\pi a^2 \sigma_0$ fully characterizes the distribution. Below we set $\kappa a = 1$, $R/a = 3$, and $L = 13$ throughout unless stated otherwise, and we identify $Z_0\lambda_B/a$ as the only other relevant dimensionless combination. For $Z_0\lambda_B/a = 22.5$ and $g \in \{0, 0.5, 0.8\}$ figure 1(a) shows the $\theta$-dependence of $\sigma$ (inset) and $\Phi$ for several $r$, revealing isotropy (as expected) for a homogeneous surface charge ($g = 0$, arrows) and strong anisotropy for the heterogeneous cases $g = 0.5$ ($\sigma_N = 3\sigma_S$, dot–dashed) and $g = 0.8$ ($\sigma_N = 9\sigma_S$, solid lines); the (small) oscillations with $\theta$ at $r = a$ are numerical artefacts due to the truncation at $L = 13$. The $\theta$-dependence of the potential weakens, as expected, for increasing distances $r$. Figure 1(b) shows $\Phi_\ell(r)$ for $\ell = 0, 1, 2$ and $g = 0.5$. Interestingly, the modes with $\ell = 1, 2$ have a non-vanishing electric field in the interior of the particle. The overall magnitude and spatial variation of $\Phi$ in figure 1 show the need for nonlinear screening theory. Nevertheless, in analogy to the spherically symmetric case [9] one can describe the far-field potential ($r \approx R$) and hence the colloidal interactions in terms of linear screening theory (dashed curves in figure 1(b)) with a renormalized surface charge distribution $\sigma^*(x) \equiv \sum_i \sigma_i^R P_i(x)$ that we will calculate below.

In the far-field $r \approx R$ we treat the deviation of $\Phi(r, x)$ from its angular average $\Phi_0(R) \equiv \chi_0$ at $r = R$ as a small expansion parameter, such that (1) for $r > a$ can be linearized as $\nabla^2 \Phi(r, x) \approx k^2 \left[\tan \chi_0 + (\Phi(r, x) - \chi_0)\right]$ with $k^2 = \kappa^2 \cos \chi_0$. The uniaxial solutions to this linear PB (LPB) equation read $\Phi(r, x) \approx \chi_0 - \tanh \chi_0 + \sum_{i=0}^\infty a_i i_i(k r) + b_i k_i(k r) P_i(x)$ where $i_i$ and $k_i$ are modified spherical Bessel functions. The coefficients $a_i$ and $b_i$ are integration constants that we fix by matching the LPB solution at $r = R$, for each $\ell$, to $\Phi_\ell(R) \equiv \chi_\ell$ and $\Phi'_\ell(R) \equiv \chi'_\ell$ of the nonlinear problem. This leads for every $\ell$ to the linear problem two by two

$$\chi_\ell = (\chi_0 - \tanh \chi_0) \delta_{\ell 0} + a_i i_i(k R) + b_i k_i(k R); \quad \chi'_\ell = k (a_i i'_i(k R) + b_i k'_i(k R)), \quad (10)$$

which results in explicit expressions for $a_i$ and $b_i$ given by

$$a_i = \frac{v_i (k_i k'_i(k R) - \frac{1}{k_i} i'_i(k R)) + \chi'_i k_i(k R)}{i_i(k R) k_{i+1}(k R) + i_{i+1}(k R) k_i(k R)}; \quad (11)$$

$$b_i = \frac{v_i (i_{i+1}(k R) + \frac{1}{k_i} i'_i(k R)) - \chi'_i i'_i(k R)}{i_i(k R) k_{i+1}(k R) + i_{i+1}(k R) k_i(k R)}. \quad (12)$$

where $v_i = \chi'_i = (\chi_0 - \tanh \chi_0) \delta_{\ell 0}$. The dashed curves in figure 1(b) are the result of such a far-field fit. With $a_i$ and $b_i$ explicitly known, one can extrapolate the LPB solution to $r = a$ to yield, with (4) and standard Bessel function relations, the renormalized multipoles

$$\sigma^*_\ell = -\frac{k}{4\pi \lambda_B} (a_i i_{\ell+1}(k a) + b_i k_{\ell+1}(k a)). \quad (13)$$

This expression is the multipole generalization of the well-known charge renormalization [9].

A first illustration of multipole renormalization is shown by the angles in figure 1(b), which represent slope discontinuities at $r = a$ which are proportional to $\sigma_\ell$ (nonlinear theory) and $\sigma^*_\ell$ (far-field fit). For $\ell = 0, 1$ we see $\sigma^*_\ell < \sigma_\ell$, which means that the effective charge and dipole are

![Figure 1](image-url)
renormalized downward. Interestingly, however, for ℓ = 2 the inset in the lowest panel reveals upward renormalization since σ2∗ ≠ 0 while σ2 = 0, that is, the Janus particle has a mode coupling induced far-field quadrupole signature. Multipole renormalization is quantified further in figures 2–4, where (scaled) renormalized multipoles Zℓ∗ = 4πa2σℓ∗ are shown for the monopole ℓ = 0 in figure 2, the dipole ℓ = 1 in figure 3 (scaled with g), and the higher-order multipoles ℓ = 1, . . ., 6 in figure 4. All multipoles are shown as a function of Z0λB/a. Furthermore, in figures 2 and 3 we chose several truncation levels L and heterogeneities g, and also picked two cell radii R. The dot–dashed curves denote the linear limit Zℓ∗ = Zℓ. Figure 2 and 3 show that all curves for L = 13 superimpose on those of L = 9, 11 for all Z0 and g ≤ 0.8, indicative of excellent convergence in this parameter regime; for g = 1 the convergence deteriorates for Z0λB/a ≥ 15. The R dependence in figure 2 and 3 shows the strongest renormalization in the largest cell, not unlike the homogeneous charge case [9]. Interestingly, in the nonlinear regime Z0λB/a ≥ 10 figure 2 shows a mode coupling induced reduction of Z0∗ by 10’s of per cents when g increases from 0 to 1. In other words, in contrast to the more usual ‘linear’ electrostatics we now have a far-field monopole potential that is not only determined by the net charge but in fact also by its heterogeneity. This is a key finding, relevant for understanding patchy-particle interactions. The mode coupling has an even stronger effect on renormalization of Zℓ∗, for which figures 3(a) and (b) show a pronounced maximum in between the low-Z0 linear screening regime and the high-Z0 regime in which Zℓ∗ becomes even vanishingly small for all g’s considered. Figure 4 shows, for g = 0.4, that in fact all Zℓ∗ with ℓ ≠ 0 vanish in the limit of large Z0, while they all show an intermediate regime with finite values even for ℓ = 2, 4, 6 for which σℓ = 0. The underlying physics for non-oppositely charged hemispheres with σH > σS > 0 (that is, with 0 < g < 1) is that both σH and σS renormalize, if both are high enough, to the same saturated value, giving rise to a pure far-field monopole without multipoles.

Figure 2. Renormalized (scaled) monopole Z∗_0 for several g, L (see text) of a Janus sphere as a function of the total charge Z0λB/a. The graph includes data for cell radii R/a = 1.5 and R/a = 3.0. The dot–dashed curve denotes Z∗_0 = Z_0.

Figure 3. Renormalized (scaled) dipole Z∗_1 for several g, L (see text) of a Janus sphere as a function of the total charge Z0λB/a. The upper and the lower graph represent cell sizes R/a = 3.0 and R/a = 1.5 respectively. The thick dotted lines denote the (scaled) difference of the renormalized northern and southern charge presumed distributed homogeneously (see text) for g = 0.8. The dot–dashed curves in both graphs denote Z∗_0 = Z_0.

Figure 4. Renormalized (scaled) multipoles Zℓ∗ of a Janus sphere as a function of the total charge Z0λB/a. The parameters here are g = 0.4 and L = 13.

The idea might emerge that both hemispheres renormalize their charge independent of each other, that is, some of the results could suggest that the renormalized charge density on the colloidal surface is a function which depends only locally on the bare charge density. If that were the case, it would suffice to calculate the renormalized surface charge density for σN and σS, as if both were the charge density of a monopole. The cell radius is to be kept unchanged. The thick dotted lines in figures 3(a) and (b) denote the resulting dipole charge for g = 0.8, which is calculated with the obtained
screening theory could be a firm basis to further study these intricate features of heterogeneously charged particles. The category of particles which are described by \( g > 1 \), carrying positive and negative charges on the two hemispheres, are particularly interesting. Because the present method is not very efficient at high values of \( g \), we have developed an alternative method to treat these mainly dipolar particles within a mode expansion as well. This will be presented in further work.

It is tempting to model Janus spheres by a Yukawa-segment model [13] in which every surface element \( dS \) contributes \( \sigma^*(x)\lambda\delta s^{-1}\exp(-\kappa s)dS \) to the (dimensionless) potential at a distance \( s \). Here the renormalized charge densities on both hemispheres is obtained from the renormalized monopole and dipole charge density via

\[
\sigma^*(x) = \begin{cases} 
\sigma^*_{\theta} + \frac{2}{\lambda} \sigma^*_{\phi} & x < 0; \\
\sigma^*_{\phi} - \frac{2}{\lambda} \sigma^*_{\theta} & x \geq 0.
\end{cases}
\]  

The dotted curves in figure 1(a) show that agreement with our full calculations is reasonable though not quantitative; the Yukawa model ignores the ion hard-core exclusion in the interior of the particle. Therefore, effectively it describes the (dimensionless) potential of a charge configuration in which oppositely charged ions were able to approach the heterogeneously charged surface from two sides, such that this potential is more suppressed compared to the full theory.

Being a point of discussion, we return to the choice of the boundary conditions on the cell’s surface (3), which we called ‘isotropic’ BCs. This denomination follows from the fact that, considering two randomly oriented neighbouring cells, the cell-surface potentials of two cells should match on the spot where they touch, giving rise to a constant cell-surface potential. Nevertheless the choice of BCs is not unique. We can also supply the system with ‘nematic’ boundary conditions, corresponding to the situation that all cells are perfectly aligned such that cells only touch on opposite spots. On these spots the electrostatic potential should match, and we can also demand continuity of the electric field. The BCs then become

\[
\Phi(R, \theta, \varphi) = \Phi(R, \pi - \theta, \pi + \varphi),
\]

\[
\Phi'(R, \theta, \varphi) = -\Phi'(R, \pi - \theta, \pi + \varphi). \tag{15}
\]

In fact one can even interpolate between ‘isotropic’ and ‘nematic’ BCs by introducing an orientation distribution function [14]. In this article, we assume a system in which the cell boundary is best described by isotropic BCs, given by (3). However, for the parameters used in this article, it turns out that this particular choice for the BCs did not noticeably affect the values of the renormalized charges. We do not see a significant change by turning to nematic BCs (15). This insensitivity to the choice of BCs is due to the fact that the nonlinear behaviour is an effect which takes place close to the colloidal surface, where these BCs have the least influence on the electrostatic potential. Furthermore, the only multipoles which are directly affected by the particular choice of BCs are the nonzero (\( \ell > 0 \)) even multipoles, which are small for Janus particles. We therefore think that the obtained values for the

\[\text{Figure 5. Renormalized monopole charge } Z^*_0 \text{ for several } g \text{ of a Janus sphere as a function of the cell size } R, \text{ both for } Z_0 \lambda B/a = 22.5, \text{ in (a) for } \kappa a = 1 \text{ and in (b) for } \kappa a = 3. \text{ The thick solid line corresponds to a pure bare monopole, yielding } g = 0. \text{ The diamonds (} g = 0.4 \text{) and circles (} g = 0.8 \text{) show the data from the full theory, using } L = 13. \text{ The thinner solid lines denote the average of the renormalized northern and southern charge presumed distributed homogeneously (see text) for } g = 0.4 \text{ and 0.8.}\
\]
renormalized multipoles can be applied in a model to describe the behaviour of a many-body system within linear theory, no matter what the orientations of the surrounding colloids are. Since the monopole and multipole potentials decay for large $r$ equally fast as $\exp(-\kappa r)/r$ [15], where $\kappa^{-1}$ is the decay length, the renormalized multipole charges are expected to contribute in dense as well as dilute systems.

In summary, we have developed a systematic framework for nonlinear ionic screening of heterogeneously charged spheres. The scheme allows for an explicit far-field analysis that generalizes charge renormalization from the well-studied homogeneous case (pure monopole) [9] to the heterogeneous case (dipoles, quadrupoles, etc and their nonlinear couplings). Application to charged Janus spheres shows (i) a 40% reduction of the effective monopole for $g = 1$ (charged and uncharged hemisphere) compared to $g = 0$ (homogeneously charged sphere), (ii) a mode coupling induced far-field effective quadrupole component without an actual surface quadrupole, (iii) a pure far-field monopole with vanishing higher-order multipoles in the saturated high charge limit, and (iv) no quantitative agreement with a simple Yukawa-segment model based on renormalized multipoles. Our study opens the way to systematic microscopic calculations of effective electrostatic interactions between Janus (and other patchy) particles. In addition, our analyses also reveal non-vanishing electric fields inside heterogeneously charged particles, which could couple to interior dipoles and affect (anisotropic) mutual Van der Waals forces. Given that the presently introduced expansion technique can be generalized to other geometries (for example, patterned planar surfaces or ellipsoidal patchy colloids), our technique and findings are directly relevant for nonlinear ionic screening of heterogeneously charged spheres. The scheme allows for an explicit far-field analysis of patchy or patterned nanoparticles, colloids, or proteins [1].

Acknowledgments

This work is part of the research programme of the Stichting voor Fundamenteel Onderzoek der Materie (FOM), which is financially supported by the Nederlandse Organisatie voor Wetenschappelijk Onderzoek (NWO). Financial support from NWO-CW-ECHO, NWO-VICI, and FOM-SFB-TR6 is acknowledged.

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