The role of multipoles in counterion-mediated interactions between charged surfaces: strong and weak coupling

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Abstract
We present general arguments for the importance, or lack thereof, of structure in the charge distribution of counterions for counterion-mediated interactions between bounding symmetrically charged surfaces. We show that on the mean field or weak coupling level, the charge quadrupole contributes the lowest order modification to the contact value theorem and thus to the intersurface electrostatic interactions. The image effects are non-existent on the mean field level even with multipoles. On the strong coupling level the quadrupoles and higher order multipoles contribute additional terms to the interaction free energy only in the presence of dielectric inhomogeneities. Without them, the monopole is the only multipole that contributes to the strong coupling electrostatics. We explore the consequences of these statements in all their generality.

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

1. Introduction
The assembly of colloidal building blocks with designer engineered size, shape, and chemical anisotropy seems to be the next step in the fundamental and applied colloid science and science of soft materials, vigorously pursued by many researchers [1, 2]. Moving beyond the traditional systems implies the creation of designer colloidal building blocks such as colloidal molecules, Janus spheres, and other patchy particle motifs that have very different propensities for self-assembly [3]. In order to realize this goal one needs detailed control over various aspects of colloid geometry at the nanoscale (e.g., aspect ratio, faceting, branching, roughness) and microscale (e.g., chemical ordering, shape gradients, unary and binary colloidal ‘molecules’). The ability to spatially modify the surface structure of colloids with designed chemical heterogeneity, e.g.,, to form a patchy surface structure, seems to be becoming a realistic goal. Fabrication of stable anisotropic microcapsules was recently accomplished by the layer-by-layer polyelectrolyte adsorption technique combined with the particle lithography technique to produce anisotropic polymer microcapsules with a single nanoscale patch [4].
Granick and co-workers [2] recently reported a highly scalable synthetic pathway for creating bipolar Janus spheres, i.e., particles that consist of oppositely charged hemispheres. Such colloidal ‘animals’ which are probably the simplest example of patchy colloids, exhibit orientation-dependent interactions that go together with localized patches of like/unlike charges or hydrophobic/hydrophilic regions. This heterogeneous interaction landscape promotes the formation of larger colloidal molecules and clusters that are themselves patchy.

These advances in the nanoscopic and microscopic tailoring of (charged) colloids motivated various approaches to generalizations of the existing theories of electrostatic interactions in charged colloids by explicitly including the structure of the counterions as embodied by their multipolar moments. The inclusion of structured counterions of a dipolar [5, 6] and quadrupolar [7, 8] type into the theory of electrostatic colloidal interactions has brought forth some of the salient features of the counterion structure effects, which on face value appear to be quite distinct from the standard Poisson–Boltzmann framework. In view of these advances in the study of interactions between charged colloids it thus seems appropriate to explore the ramifications of the emerging paradigm if applied to these more complicated structured colloidal molecules. In particular, the nonspherically symmetric charge distribution of microscopically tailored colloidal particles might lead to some unexpected properties of electrostatic interactions in systems of this type. With this in mind we thus embark on a thorough examination of the consequences of multipolar charge distribution of mobile counterions that mediate interactions between charged (planar) macromions.

2. Model

In what follows we will consider a system of fixed macromions of surface charge distribution \( \rho_0(\mathbf{r}) \) in an aqueous solution, described as a dielectric continuum with a dielectric constant \( \varepsilon \) and temperature \( T \), containing \( N \) neutralizing counterions. Counterions are assumed to be point-like particles but they do possess a rigid internal structure described by a charge distribution \( \hat{\rho}(\mathbf{r}; \mathbf{R}, \omega_i) \) that we assume can be written as a standard multipolar expansion [9]:

\[
\hat{\rho}(\mathbf{r}; \mathbf{R}, \omega_i) = e_0 q \delta(\mathbf{r} - \mathbf{R}_i) - p_0 (\mathbf{n}_i \cdot \nabla) \delta(\mathbf{r} - \mathbf{R}_i) + t_0 (\mathbf{n}_i \cdot \mathbf{r}) \delta(\mathbf{r} - \mathbf{R}_i) + \cdots,
\]

(1)

for the \( i \)th counterion located at \( \mathbf{R}_i \) with \( \omega_i \) being the orientational variables specifying the angular dependence of the counterion charge density. The monopolar moment of each counterion is \( e_0 q \), where \( q \) is the charge valency and \( e_0 \) the elementary charge, \( \mathbf{p} = p_0 \mathbf{n}_i \) is the dipolar moment and \( \mathbf{Q} = t_0 \mathbf{n}_i \otimes \mathbf{n} \) the quadrupolar moment with director unit vector \( \mathbf{n} \). We must emphasize here that the quadrupolar expansion in (1) is not general but adequately describes only a uniaxial counterion, e.g., a charged particle of rod-like structure; figure 1. One possible implementation of counterions possessing just monopolar and quadrupolar moment is a uniformly charged rod with charge \( e_0 q \) and length \( l \). The counterion’s quadrupolar moment is then \( t_0 = e_0 q l^2 / 24 \).

Another possibility is a negative charge \((-e_2)\) in the center and two positive charges \((+e_1)\) located at both ends of the rod leading to monopolar and quadrupolar moments of \( e_0 q = 2e_1 - e_2 \) and \( t_0 = e_1 l^2 \), respectively.

Counterions interact via a Coulomb interaction potential \( u(\mathbf{r}, \mathbf{r}') \) so the interaction energy of two given counterions \( i \) and \( j \) at set positions and orientational configurations is obtained by integrating the Coulomb interaction over their internal orientational degrees of freedom as

\[
\int \int \mathrm{d}^3 \mathbf{r} \int \mathrm{d}^3 \mathbf{r}' \hat{\rho}(\mathbf{r}; \mathbf{R}, \omega_i) u(\mathbf{r}, \mathbf{r}') \hat{\rho}(\mathbf{r}'; \mathbf{R}_j, \omega_j).
\]

(2)

Although our formalism in this section is in general applicable to macromions of arbitrary shape and charge distribution, we shall primarily focus on the case of two charged planar surfaces located at \( z = \pm a \) with the charge distribution

\[
\rho_0(\mathbf{r}) = \sigma \delta(a - z) + \sigma \delta(a + z).
\]

(3)

We also consider a dielectric inhomogeneity between the bounding surfaces and the ionic solution, such that the ionic solution is described with dielectric constant \( \varepsilon \) and the bounding surfaces with dielectric constant \( \varepsilon' \). In this geometry the Coulomb interaction potential is composed of the direct interaction \( u_{0}(\mathbf{r}, \mathbf{r}') = \left( \frac{4 \pi \varepsilon \varepsilon_0}{|\mathbf{r} - \mathbf{r}'|} \right)^{-1} \) and the (electrostatic) image interaction \( u_{im}(\mathbf{r}, \mathbf{r}') \), so

\[
u(\mathbf{r}, \mathbf{r}') = u_{0}(\mathbf{r}, \mathbf{r}') + u_{im}(\mathbf{r}, \mathbf{r}').
\]

(4)

The Green’s function in planar geometry can be expressed as a sum of image charge contributions, namely

\[
u(\mathbf{r}, \mathbf{r}') = \frac{1}{4 \pi \varepsilon \varepsilon_0} \left[ \sum_{n \text{ even}} \frac{\Delta[n]}{|\mathbf{r} - \mathbf{r}' - 2na| k_l} + \sum_{n \text{ odd}} \frac{\Delta[n]}{|\mathbf{r} - \mathbf{r}' + (2na - 2z)| k_l} \right].
\]

(5)

In both sums the index \( n \) runs through negative and positive integer values. The term corresponding to \( n = 0 \) represents the direct Green’s function of the particle in a space of a uniform dielectric constant, namely \( u_0(\mathbf{r}, \mathbf{r}') \). The unit vector \( \mathbf{k} = (0, 0, 1) \) points in the +z direction. The relative dielectric
jump at the two bounding surfaces is quantified as \( \Delta = (\varepsilon - \varepsilon')/(\varepsilon + \varepsilon') \). For most relevant situations the outer permittivity is smaller than the inner one, \( \varepsilon' < \varepsilon \), so the dielectric jump is positive, \( \Delta > 0 \), and vanishes in the homogeneous case with \( \varepsilon' \rightarrow \varepsilon \).

The canonical partition function \( Z_N \) of this Coulomb fluid composed of \( N \) charged particles is given by

\[
Z_N = \frac{1}{N!} \int d\mathbf{R}_1 \cdots d\mathbf{R}_N d\omega_1 \cdots d\omega_N \times \exp(-\beta U[\mathbf{R}_i, \omega_i; \mathbf{R}_j, \omega_j]),
\]

with the configurationally dependent ion interaction energy given by

\[
U[\mathbf{R}_i, \omega_i; \mathbf{R}_j, \omega_j] = \frac{1}{2} \sum_{i \neq j} \int \! d\mathbf{r}' \hat{\rho}(\mathbf{r}; \mathbf{R}_i, \omega_i) u(\mathbf{r}, \mathbf{r}') \hat{\rho}(\mathbf{r}'; \mathbf{R}_j, \omega_j) + \sum_i \int \! d\mathbf{r}' \hat{\rho}(\mathbf{r}; \mathbf{R}_i, \omega_i) u(\mathbf{r}, \mathbf{r}') \rho_0(\mathbf{r}') + \frac{1}{2} \int \! d\mathbf{r}' \rho_0(\mathbf{r}) u(\mathbf{r}, \mathbf{r}') \rho_0(\mathbf{r}'),
\]

where \( \beta = 1/k_B T \). The three terms in \( U[\mathbf{R}_i, \omega_i; \mathbf{R}_j, \omega_j] \) correspond to direct electrostatic interaction between counterions, electrostatic interactions between counterions and fixed charges and ones between fixed charges on the walls themselves, respectively. We will furthermore make the standard assumption that the system is overall electroneutral, implying that the mobile countercharge exactly compensates the charge on the surfaces.

We now proceed in the Netz [10] fashion and perform the Hubbard–Stratonovich transformation of the partition function [11], where the configurational integral over counterion positions is transformed into a functional integral over a fluctuating auxiliary electrostatic potential \( \phi(\mathbf{r}) \). In this way the grand canonical partition function is obtained straightforwardly in the form [10]

\[
Z = \sum_{N=0}^{\infty} \lambda^N Z_N = C \int \mathcal{D}[\phi(\mathbf{r})] e^{-\beta H[\phi(\mathbf{r})]},
\]

where \( \lambda \) is the bare fugacity which is the exponential of the chemical potential. The prefactor \( C \) above is the functional determinant of the inverse Coulomb kernel \( u^{-1}(\mathbf{r}, \mathbf{r}') \) while the ‘action’ of the functional integral is given by

\[
H[\phi] = \int \! d\mathbf{r} \left[ \frac{1}{2} \varepsilon(\mathbf{r}) \varepsilon_0 (\nabla \phi)^2 - i \rho_0(\mathbf{r}) \phi(\mathbf{r}) \right] - \lambda' \int \! d\omega \Omega(\mathbf{r}) \exp \left[ i \beta \int \! d\mathbf{r}' \hat{\rho}(\mathbf{r}'; \mathbf{r}, \omega) \phi(\mathbf{r}') \right].
\]

Here we have introduced the renormalized fugacity as \( \lambda' = \lambda \exp(\mu_0(\mathbf{r}, \mathbf{r})), \) where \( \mu_0(\mathbf{r}, \mathbf{r}) \) is the electrostatic direct self-energy of a single counterion with multipolar charge distribution. \( \Omega(\mathbf{r}) \) is the geometric characteristic function of the counterions, being equal to unity in the slab between the bounding surfaces and zero otherwise. The partition function in the above form cannot be evaluated explicitly except in the one-dimensional case [12] where the partition function evaluation is reduced to a solution of a Schrödinger-like equation. Nevertheless the field-theoretical representation of the grand canonical partition function allows one to use quite powerful analytical approaches that eventually lead to an explicit evaluation of the partition function in two well defined and complementary limits [10, 13, 14] that we shall address later in this paper. These limits retain relevance also in the case of structured counterions.

3. Dimensionless representation

One may obtain a dimensionless representation for the present system by rescaling all length scales with a given characteristic length scale. Recall that the characteristic distance at which two unit charges interact with thermal energy \( k_B T \) is known as the Bjerrum length \( \ell_B = e_0^2/(4\pi \varepsilon \varepsilon_0 k_B T) \) (in water at room temperature, the value is \( \ell_B \approx 0.7 \) nm). If the charge valency of counterions is \( q \) then the aforementioned distance scales as \( q^2 \ell_B \). Similarly, the distance at which a counterion interacts with a macromolecular surface of surface charge density \( \sigma \) with an energy \( k_B T \) is called the Gouy–Chapman length, defined as \( \mu_0 = e_0/(2\pi q^2 \ell_B \sigma) \). A competition between ion–ion and ion–surface interactions can thus be quantified by the ratio (\( \Xi \)) of these characteristic lengths, that is

\[
\Xi = q^2 \ell_B/\mu = 2\pi q^3 \ell_B^2 \sigma/\varepsilon_0,
\]

which is referred to as the electrostatic coupling parameter.

In what follows, we may rescale the length scales with the Gouy–Chapman length, i.e., \( \mathbf{r} \rightarrow r/\mu \); hence, the surface separation will be rescaled as \( D = D/\mu \) or the rescaled half-distance as \( \tilde{a} = a/\mu \). Other dimensionless quantities can be defined as follows. The dimensionless multipolar moments are defined as

\[
p = p_0/e_0 q \mu \quad \text{and} \quad t = t_0/e_0 q \mu^2,
\]

and the dimensionless pressure as

\[
\tilde{p} = \frac{\beta p}{2\pi \ell_B (\sigma/\varepsilon_0)^2}.
\]

For simple structureless counterions the functional integral (9) can be rescaled [10] yielding

\[
Z \rightarrow C \int \mathcal{D}[\phi(\mathbf{r})] e^{-H[\phi(\mathbf{r})]/\Xi},
\]

with dimensionless \( H'[\phi(\mathbf{r})] \). As will be shown later an identical representation can be obtained also for structured counterions, described on the level of a multipolar expansion of their charge density.

4. Strong and weak coupling dichotomy

In the absence of a general approach that would cover thoroughly all the regions of the parameter space one has to take recourse to various partial formulations that take into account only this or that facet of the problem [15]. The traditional approach to these one-component Coulomb fluids
has been the mean field Poisson–Boltzmann (PB) formalism applicable at weak surface charges, low counterion valency and high temperature [13, 16, 17]. The limitations of this approach become practically important in highly charged systems where counterion-mediated interactions between charged bodies start to deviate substantially from the mean field accepted wisdom [15]. One of the most important recent advances in this field has been the systematicatization of these non-PB effects based on the notions of weak and strong coupling approximations. They are based on the field-theoretical representation of grand canonical partition function (10), whose behavior depends on a single dimensionless coupling parameter $\Xi$ [10] (see below). The weak coupling (WC) limit of $\Xi \to 0$ coincides exactly with the mean field Poisson–Boltzmann theory [20] and is based on a collective description of the counterion density. The strong coupling (SC) limit of $\Xi \to \infty$ is diametrically opposite and corresponds to a single-particle description. It has been pioneered by Rouzina and Bloomfield [21], elaborated later by Shklovskii et al [22–24] and Levin et al [25], and eventually brought into a final form by Netz et al [10, 13, 15, 26–28].

These two limits are distinguished by the pertaining values of the coupling parameter $\Xi$ which can be introduced in the following way.

The regime of $\Xi \ll 1$ is the case of low counterionic valency and/or weakly charged surfaces, and is referred to as the weak coupling limit. It is characterized by the fact that the width of the counterion layer $\mu$ is much larger than the separation between two neighboring counterions in solution and thus the counterion layer behaves basically as a three-dimensional gas. Each counterion in this case interacts with many others and the collective mean field approach of the Poisson–Boltzmann type is completely justified.

On the other hand in the strong coupling regime $\Xi \gg 1$, which is true for high valency of counterions and/or highly charged surfaces. In this case the mean distance between counterions, $a_\perp \sim \sqrt{e_0 q/\sigma}$, is much larger than the layer width (i.e., $a_\perp/\mu \sim \sqrt{\Xi} \gg 1$), indicating that the counterions are highly localized laterally and form a strongly correlated quasi-two-dimensional layer next to a charged surface. In this case, the weak coupling approach breaks down due to strong counterion–surface and counterion–counterion correlations. Since counterions can move almost independently from the others along the direction perpendicular to the surface, the collective many-body effects that enable a mean field description are absent, necessitating a complementary SC description [10, 18, 19].

These two approximations allow for an explicit and exact treatment of charged systems at two disjoint limiting conditions whereas the parameter space in between can be analyzed only approximately and is mostly accessible solely via computer simulations. As will become clear in what follows the WC and SC limits remain valid also for structured counterions described with a multipolar charge distribution.

### 5. Mean field limit

The mean field limit, $\Xi \to 0$, is defined via the saddle-point configuration of the Hamiltonian (10) [20] and is valid for a weakly charged system. This leads to the following generalization of the Poisson–Boltzmann equation

$$\varepsilon \varepsilon_0 \nabla^2 \psi_0(\mathbf{r}) = -\lambda' \int \text{d}r \, \delta(\mathbf{r} - \mathbf{r}', \omega) \Omega(\mathbf{r}') \times \exp \left[ -\beta \int \text{d}r' \, \hat{\rho}(\mathbf{r}'; r', \omega) \psi_0(\mathbf{r}') \right]$$

for the real-valued potential field $\psi_0 = -i\phi_0$. Note that the dielectric discontinuity at the boundaries is irrelevant within the mean field theory [29] since in the planar geometry this is effectively one-dimensional theory. Taking now the counterion density function as a sum of the monopolar, dipolar and quadrupolar terms, where the mean field depends only on the transverse coordinate $z$, the above equation can be written in dimensionless form as

$$\psi'' = -\frac{1}{\Xi} \int_{z_1}^{z_2} \text{d}x \Omega(x, z)(u(z) - pxu'(z) + tz^2u''(z)), \quad \propto \rho_1(z) + \rho_2(z) + \rho_3(z),$$

where we have defined the dimensionless potential $\psi = \beta e_0 q \psi_0$ with corresponding derivatives $\psi' = \beta e_0 q \mu \psi_0'$ and $\psi'' = \beta e_0 q \mu^2 \psi_0''$ and dimensionless multipolar moments $p$ and $t$. We defined the orientational variable $x = \cos \theta$, where $\theta$ is the angle from the $z$ axis, assumed to coincide with the normal to the bounding surfaces, the director of the uniaxial counterion is $\mathbf{n}$ and the integral over this variable gives the orientational average. We have also introduced

$$u(z) = C \exp \left( -\psi - px \psi' - tz^2 \psi'' \right),$$

which is obviously the locally orientationally dependent number density of the counterions and $\rho(z)$ is the corresponding orientationally averaged charge density of the counterions. In the case of higher multipoles the number density and the charge density of the counterions are not proportional. Expressions $\rho_i(z)$ are simply the orientationally averaged multipolar charge densities, i.e., $i = 1$ for monopolar charge, $i = 2$ for dipolar charge etc, that are simply proportional to the three terms in the integrand of (14). The corresponding prefactor is simply obtained by appropriate normalization of $\rho_1$ to satisfy the electroneutrality condition of the system.

The above PB equation has to be supplemented by an appropriate boundary condition at $z = \pm a$ corresponding to the electroneutrality of the system by taking into account surface charges, $\sigma$. The constant $C$ is set by these boundary conditions, as is the case in the standard PB theory. In dimensionless units these boundary conditions read $\psi'(-a) = \frac{1}{2}$. Obviously in the case of counterions with only monopolar charge distribution the above set of equations reduces to the standard Poisson–Boltzmann theory.

The characteristic function $\Omega(x, z)$ for the parallel plane geometry simply excludes the counterion configurations that would penetrate the bounding walls and depends on the geometric form of the counterions. Since in what follows we will not be interested in steric effects, we will assume that all counterions are point-like, and so $\Omega(x, z) = 1$. Steric effects have been studied elsewhere [6–8].

Because of the similarity with the standard PB equation for monopolar charges one is led to believe that the pressure
in an inhomogeneous system of multipolar countercharge can
derived in the same way as in the standard PB theory, via the s.c. contact value theorem [30]. Indeed this can be
proven exactly, namely the mean field pressure can be obtained
from the first integral of the PB equation (14). After some
manipulations its first integral is obtained in the form

$$\tilde{P} = -\frac{1}{2} \psi^{\prime 2} + \frac{1}{2} \int_{-1}^{1} dx \left[ u + pxu \psi^{\prime} + tx^2 (u \psi^{\prime\prime} - u^{\prime} \psi^\prime) \right].$$

(15)

Here $\tilde{P}$ is the dimensionless equilibrium pressure in the
system, defined as (12). The rhs of the above identity can be
calculated at any arbitrary point $|z_0| < a$ as its value is
independent of $z_0$. The choice of the actual point is governed
by the symmetry of the system, as is usual also in the standard
PB theory. Since we delve only into the symmetrical solution,
with inversion symmetry centered on $z = 0$, the derivative of
the mean potential at the mid-point must vanish, i.e., $\psi^{\prime}(z_0 = 0) = 0$. Thus in this case

$$\tilde{P} = \frac{1}{2} \int_{-1}^{1} dx \ u(0)(1 - tx^2 |\psi^{\prime\prime}(0)|).$$

(16)

While the first—van’t Hoff—term corresponds to repulsive
interactions of the standard PB type, the second term in this
particular geometry and symmetry entails attractive
interactions between the bounding surfaces. Note that $\psi^{\prime\prime} < 0$.

Let us take a closer look at the above expression. For a
polyelectrolyte chain, i.e., an extended and flexible counterion,
one can derive a similar type of pressure formula [31], with
a negative term that contributes an attractive part to the force
equilibrium. This attractive part in the case of polyelectrolytes
is due to polyelectrolyte bridging interactions [32] that stem
from the connectivity and flexibility of the polyelectrolyte
chain.

Does equation (16) have a similar physical content? This
interpretation certainly does not seem likely for point
counterions, for which the above pressure formula was derived.
The structure of the first integral of the PB equation (16)
seems to be saying that apart from the ideal contribution to
the equilibrium pressure, a term proportional to the mid-point
seems to be saying that apart from the ideal contribution to
the interaction pressure is to induce a small attractive contribution.
In order to show this we expand (14) to the first order in $t$ and
put $p = 0$, which then reads

$$\psi^{\prime\prime} = -Ce^{-\psi}(1 + \frac{1}{4}t \psi^{\prime 2} - \frac{1}{2}t \psi^{\prime\prime}).$$

(17)

This equation can be solved much more easily with
typical numerical methods. Although we must be aware
that it is valid only for small $t$. The corresponding formula for
interaction pressure in the expansion to first order in $t$ reads

$$\tilde{P} = \frac{1}{2} Ce^{-\psi(0)}.$$  

(18)

Due to a convenient cancelation the formula contains no
explicit dependence on $t$, but the pressure still depends on
$\psi$ implicitly, due to the $t$-dependent potential $\psi$. As can be
shown numerically for small $t$, the correction to the standard
PB pressure depends linearly on $t$.

5.2. Numerical results in the mean field limit

Here we are not interested in the details of the mean
field results—they were analyzed in detail before [6–8]—
but list them for completeness anyway. In what follows we
limit ourselves to counterions that possess monopolar and
quadrupolar charge, so we do not take into account any dipoles
($p = 0$).

The density profile in figure 2(a) corresponds to
monopolar density $\rho_{1}(z)$, (14). Counterions are localized
preferentially in the vicinity of the two surfaces due to mutual
repulsion that prevents them from being localized at the center.
What we then observe is that with increase of the quadrupolar
moment, $t$, counterions concentrate at surfaces even more.
This can be easily explained. The potential energy of every
quadrupolar particle in the electrostatic potential is

$$\beta W = t \psi^{\prime\prime} \cos^2 \theta.$$  

(19)

Since the second derivative of the mean field potential $\psi^{\prime\prime}$
in the symmetric case considered here is a concave function of
the coordinate $z$, this means that the quadrupolar force, $F =
-\partial W / \partial z$, acts away from the center toward both surfaces.

Insight into the orientation of quadrupolar moments can be
obtained through the average of the square of variable $x$, namely
($x^2$),

$$\langle x^2 \rangle = \frac{\int x^2 u(x) \, dx}{\int u(x) \, dx}.$$  

(20)

It is instructive to define an orientational order parameter

$$S = \frac{1}{3}(3 \langle x^2 \rangle - 1).$$  

(21)

This order parameter can have values in the range from $-1/2$
to 1. In a totally disordered system where quadrupoles
point randomly in all possible directions, $\langle x^2 \rangle = 1/3,$ and
the orientational order parameter vanishes, $S = 0$. In
the completely ordered case, $S = 1$ and $\langle x^2 \rangle = 1$, so
all quadrupolar moments are parallel with the $z$ axis and
perpendicular to the walls, $\theta = 0$. The other extremum,
$S = -1/2$, corresponds to $\langle x^2 \rangle = 0$ so all quadrupolar
moments are perpendicular to the $z$ axis and parallel with the walls, $\theta = \pi/2$.

Figure 2(b) shows the numerically obtained profile of the orientational order parameter. It increases with increasing quadrupolar strength $t$ and is larger at both surfaces than at the center. The order parameter $S > 0$ indicates that quadrupolar moments are preferentially aligned parallel to the $z$ axis. According to (19) the electrostatic energy can be minimized by increasing $\cos^2 \theta$, since $\psi'' < 0$. So $(\cos^2 \theta)$ really is preferentially increased above the value of $1/3$.

The quadrupolar contribution to the pressure for $t > 0$ turns out to be attractive (figure 2), as was shown before [6–8]. Again this attractive contribution is obtained here on the mean field level where the inter-counterionic correlations, essential in the strong coupling limit, play no role. It is indeed the intra-counterion correlations (entering here via the rigid structure assumed for counterions) that lead to such attractive contributions. At this juncture it does not make much sense to us to continue the multipolar expansion to yet higher orders, that would become relevant for even more aspherical and elongated charge distributions that would obviously entail also some molecular flexibility. In that case one could use the well developed theory of polyelectrolyte-mediated interactions with much better confidence [35].

6. Strong coupling limit

In the strong coupling limit, $\Sigma \gg 1$, the system is highly charged and counterions are highly correlated. The partition function (9) can be approximated by the two lowest order terms in the virial expansion with respect to the (renormalized) fugacity [10]

$$
Z = Z_{SC}^{(0)} + \lambda Z_{SC}^{(1)} + O(\lambda^2).
$$

(22)

The second term corresponds to a one-particle partition function as the SC limit is effectively a single-particle theory. $Z_{SC}^{(0)}$ and $Z_{SC}^{(1)}$ are then given by

$$
Z_{SC}^{(0)} = \exp \left[ -\frac{1}{\beta} \int \int \rho_0(r) u(r, r') \rho_0(r') \right],
$$

(23)

which is the exponential of the interaction between bare surface charges, and

$$
Z_{SC}^{(1)} = \int \int \mathrm{dR} \mathrm{d} \omega \exp \left[ -\beta \int \int \mathrm{d}r \mathrm{d}r' \rho(r; R, \omega) u(r, r') \right. \\
\times \rho_0(r') - \frac{1}{\beta} \int \mathrm{d}r \mathrm{d}r' \\
\left. \times \rho(r; R, \omega) u(r, r') \rho(r'; R, \omega) \right].
$$

(24)

which is the partition function of all possible (single-) counterion configurations. The interaction potential in this part of the partition function is composed of the direct and image electrostatic interactions (4).

In the case of two charged surfaces with uniformly smeared surface charge density (3), the surface charge electrostatic potential does not depend on the $z$ coordinate; it is spatially homogeneous and is given by

$$
\int \mathrm{d}r' u(r, r') \rho_0(r') = -\frac{\sigma \beta a}{\epsilon \epsilon_0}.
$$

(25)

The corresponding potential energy of a counterion in this surface electrostatic potential is given by

$$
\int \int \mathrm{d}r' \rho(r; R, \omega) u(r, r') \rho_0(r') = -\frac{\sigma \beta a}{\epsilon \epsilon_0} \epsilon_0 q.
$$

(26)

Since all the terms in density operator (1), except the first one, depend on the gradients, i.e., spatial derivatives, the counterion energy in a homogeneous external electrostatic potential depends only on the first, monopolar term (26). That means that higher multipoles do not interact directly with planar surface charge, but we must emphasize that this is not the case in an inhomogeneous potential at curved surfaces, e.g., at cylindrical or spherical surfaces or inhomogeneously charged surfaces.

As for the self-energy contribution, the second term in the exponent of (24), it only picks up terms from the $z$-dependent parts of the image self-interaction, $u_{im}$. The direct self-interaction, $u_{0i}$, does not depend on coordinates so it can
be discarded. The self-image energy is then

$$\int \int d\mathbf{r}d\mathbf{r}'\hat{\rho}(\mathbf{r}, \mathbf{R}, \omega)u_{im}(\mathbf{r}, \mathbf{r}')\hat{\rho}(\mathbf{r}', \mathbf{R}, \omega)$$

$$= \sum_{i=1}^{9} w_i(z, \cos \theta). \quad (27)$$

Self-image contributions are among monopolar, dipolar and quadrupolar moments of the counterions interacting with their own electrostatic images of monopolar, dipolar and quadrupolar moments. Summing up all these contributions we are left with nine terms of which only six are different. Writing them up in {	extit{extenso}} we obtain

$$w_1(z, \cos \theta) = q^2e_0^2 u_{im}(\mathbf{R}, \mathbf{R}),$$

$$w_2(z, \cos \theta) = q_0^2 \phi_0^2 (\mathbf{n} \cdot \nabla)u_{im}(\mathbf{r}, \mathbf{r})|_{\mathbf{r}=\mathbf{R}},$$

$$w_3(z, \cos \theta) = q_0^2 \phi_0^2 (\mathbf{n} \cdot \nabla)^2 u_{im}(\mathbf{r}, \mathbf{r})|_{\mathbf{r}=\mathbf{R}},$$

$$w_4(z, \cos \theta) = w_2(z, \cos \theta),$$

$$w_5(z, \cos \theta) = p_0^2 (\mathbf{n} \cdot \nabla')u_{im}(\mathbf{r}, \mathbf{r})|_{\mathbf{r}=\mathbf{r}'},$$

$$w_6(z, \cos \theta) = p_0^2 (\mathbf{n} \cdot \nabla')^2 u_{im}(\mathbf{r}, \mathbf{r}')|_{\mathbf{r}=\mathbf{r}'},$$

$$w_7(z, \cos \theta) = w_5(z, \cos \theta),$$

$$w_8(z, \cos \theta) = w_6(z, \cos \theta),$$

$$w_9(z, \cos \theta) = t_0^2 (\mathbf{n} \cdot \nabla)^2 (\mathbf{n} \cdot \nabla')^2 u_{im}(\mathbf{r}, \mathbf{r}')|_{\mathbf{r}=\mathbf{r}'}. \quad (28)$$

Here, the first expression, $w_1$, corresponds to the interaction between a monopole and its own monopole image; therefore it is proportional to $q^2$. The second term, $w_2$, corresponds to the monopole–dipole image interaction which is obviously the same as the dipole–monopole image interaction, $w_4$, and so on for all the higher order terms.

These image self-interaction terms can be expressed in a dimensionless form as

$$\tilde{w}_i(\tilde{z}, \cos \theta) = \frac{1}{\tilde{\beta}} w_i(z, \cos \theta). \quad (29)$$

The above expressions were derived for the planar geometry. Similar expressions but with different image potentials can be derived also in other geometries with dielectric inhomogeneities. Note that if there is no dielectric discontinuity, and so $u_{im}(\mathbf{r}, \mathbf{r}') = 0$, they are all identically zero, $w_i = 0$!

Thus one can conclude at this point that in the dielectrically homogeneous case the higher order multipoles are completely irrelevant on the SC level. Only the monopolar term survives in the partition function. This is probably one of the most important conclusions of this work, so let us reiterate it: in a dielectrically homogeneous case in plane-parallel geometry the structure of the counterions as codified by their multipolar moments plays absolutely no role in the strong coupling limit!

For our case of two charged planar surfaces we evaluate the above expressions by using the Green’s function (5), which was derived for planar geometry with a step-function dielectric profile at the two boundaries $z = \pm a$. We get the following somewhat cumbersome expressions for the dimensionless self-image interactions defined above:

$$\tilde{w}_1(\tilde{z}, x) = \frac{1}{2} 2 \sum_{n \text{ odd}} \frac{n\Delta^n}{(n\Delta)^2 - \tilde{z}^2} \frac{\Delta^n}{\ln(1 - \Delta^2)},$$

$$\tilde{w}_2(\tilde{z}, x) = \frac{1}{2} \sum_{n \text{ even}} \frac{n\Delta^n}{(n\Delta)^2 - \tilde{z}^2} \Delta^n,$$

$$\tilde{w}_3(\tilde{z}, x) = -\frac{1}{4} \Delta t(1 - 3\tilde{x}^2) \Delta^n,$$

$$\tilde{w}_4(\tilde{z}, x) = \frac{1}{8} \sum_{n \text{ even}} \frac{\Delta^n}{2\Delta_t} \Delta^n + (1 + \tilde{x}^2),$$

$$\tilde{w}_5(\tilde{z}, x) = \frac{1}{8} \Delta p t(1 + x^2) x \tilde{z} \tilde{t},$$

$$\tilde{w}_6(\tilde{z}, x) = \frac{1}{2} \sum_{n \text{ odd}} \frac{n\Delta^n}{(n\Delta)^2 - \tilde{z}^2} \Delta^n,$$

$$\tilde{w}_7(\tilde{z}, x) = \frac{1}{2} \sum_{n \text{ odd}} \frac{n\Delta^n}{(n\Delta)^2 - \tilde{z}^2} \Delta^n,$$

$$\tilde{w}_8(\tilde{z}, x) = \frac{1}{32} \Delta t^2 \left( (3 - 30x^2 + 35x^4) \sum_{n \text{ even}} \frac{\Delta^n}{(n\Delta)^2} + \frac{3\tilde{x}^2 + 3x^4}{(n\Delta)^2} \Delta^n \right),$$

All indices $n$ here run only through positive values. Note again that self-image contributions, $\tilde{w}_i$, vanish when $\Delta = 0$. Taking all this into account, the strong coupling interaction free energy, $\beta F = -k_B T \ln Z_{\text{SC}}^{(1)}$, i.e., the part of the free energy that depends on the intersurface separation, can be written in dimensionless form as

$$\tilde{F}/\tilde{A} = 2\tilde{a} - 2 \ln \int_{-\tilde{a}}^{\tilde{a}} d\tilde{z} \int_{-1}^{1} dx \exp \left( -\frac{1}{9} \sum_{i=1}^{9} \tilde{w}_i(\tilde{z}, x) \right). \quad (31)$$

Just as in the WC case we have again assumed that the counterions are point-like particles so that the characteristic function $\Omega$ does not depend on their coordinate $\tilde{z}$ and therefore the integration goes from $-\tilde{a}$ to $\tilde{a}$. This means that we also disregard the possible entropic effects due to the finite size and anisotropy in the shape of counterions. These entropic contributions to the partition function are relevant only for intersurface separation on the order of the size of the counterion or smaller. In that regime of separations other, much stronger effects would come into play and compete with ionic finite size effects, thus such effects are not the focus of this paper.

The corresponding dimensionless pressure is simply obtained by taking the derivative of the free energy (31) with respect to wall separation, $\tilde{D} = 2\tilde{a}$,

$$\tilde{P} = -\frac{\partial \tilde{F}/\tilde{A}}{\partial \tilde{a}}. \quad (32)$$

It is again obvious from here that in the dielectrically homogeneous case, i.e., the case with no electrostatic images, where $w_i(\tilde{z}, x) = 0$, the strong coupling limit is given exactly
by the monopolar term (the first term in (24)). Thus without the images we are left with the same form of the strong coupling interaction free energy as in the case of monopolar point charges. It is given by

$$\tilde{F}/\tilde{A} = 2\tilde{a} - 2\ln\tilde{a},$$

with the corresponding pressure

$$\tilde{P} = \frac{1}{\tilde{a}} - 1.$$

The higher order multipoles thus make no direct contribution to the strong coupling interaction free energy or forces between the bounding charged surfaces without dielectric inhomogeneities.

This is a very powerful result whose ramifications in fact impose rather stringent limits on the significance of the multipolar expansion of counterionic charge. It appears that for highly charged counterions, that are in fact the only ones where the multipolar expansion really makes sense, most of the electrostatics is properly captured by the monopolar term. Higher order multipoles simply do not contribute to the pressure in the system in the SC limit. Of course all of this is valid in the limit of homogeneous dielectric properties for planar surfaces without any image effects.

The counterion density profile can be extracted from (31) as an integrand in the second term,

$$\tilde{\rho}(\tilde{z}) \propto \int_{-1}^{1} dx \exp \left( - \sum_{i=1}^{9} \tilde{w}_i(\tilde{z}, x) \right).$$

In the case without the dielectric mismatch the density profile is simply homogeneous, $\rho(z) = \text{const}$. Dielectric images induce an additional repulsion between the counterions and the surface charges pushing them towards the mid-plane region between the two charged bounding surfaces.

Orientational order of quadrupoles can be inspected via averaged $\langle x^2 \rangle$ which is here defined as

$$\langle x^2 \rangle = \frac{\int x^2 \exp \left( - \sum_{i=1}^{9} \tilde{w}_i(\tilde{z}, x) \right) dx}{\int \exp \left( - \sum_{i=1}^{9} \tilde{w}_i(\tilde{z}, x) \right) dx}.$$

We can now use the same definition of order parameter $S$ as in WC, (21).

6.1. Numerical results in the SC limit

We now present some numerical results for the SC limit. Without any dielectric mismatches the density profile of the monopolar counterions is homogeneous, i.e., it does not depend on $z$, which is a well known result [10]. In the case of multipoles with dielectric images this is no longer true, but the counterion density does remain an even function of $z$.

In figure 3 we show the numerical results for coupling parameter strength $\Xi = 50$, dielectric jump $\Delta = 0.95$ and $\tilde{a} = 1$. Contrary to the WC situation, here counterions are localized in the vicinity of the mid-point between the two bounding surfaces due to image repulsion. Even without quadrupolar contributions ($t = 0$) the monopolar interaction, $w_1$, is repulsive and long ranged, $w_1 \sim z^{-1}$ [29], compressing the counterions to the mid-point.

When the quadrupolar parameter, $t$, is increased a bit the mid-point peak first starts to widen. This is due to the attractive monopole–quadrupole image contribution, represented by the $w_3$ and $w_7$ terms which are proportional to $t$. As $t$ increases further (approximately for $t > 8\tilde{a}^2/45$ for large $\Delta \Xi$) the repulsive quadrupole–quadrupole image contribution, $w_9$, which is proportional to $t^2$, takes over and confines counterions to the center even more, so the density peak sharpens up.

For the value $\Delta \Xi$ large enough, so that density profile becomes bell-shaped, the SC density profile can be approximated by a Gaussian curve with a variance, i.e., peak width, of the form

$$\delta \tilde{z}^2 \approx \frac{1}{\Delta \Xi \tilde{a}^4} - \frac{\tilde{a}^7}{180} + \frac{\Delta \Xi \tilde{a}^2}{10}.$$

As can be seen, small $t$ widens the density peak, but on further increasing $t$ over the value $t > 8\tilde{a}^2/45$ the peak narrows. Also on increasing parameters $\Xi$, $\Delta$ the mid-point density peak narrows. This also means that the repulsive pressure between the walls increases. As the density peak narrows and the density of the counterions at the mid-point plane increases, the counterions become more ordered in the...
quasi-two-dimensional mid-point layer, implying also that the correlations are stronger and therefore the SC limit should \textit{a fortiori} be even more appropriate.

Figure 3(b) shows also the orientational order parameter for the SC limit. Contrary to the WC results, here the orientational order parameter is negative, implying that quadrupoles are preferentially aligned perpendicular to \( z \) axis. The parameter reaches its extremum \(-1/2\) at both walls where quadrupoles are perfectly perpendicular to the \( z \) axis and parallel to the walls. This is due to the strong quadrupole–quadrupole image repulsion, \( w_q \), which has its minimum at \( x = 0 \) and is not a purely steric effect as in simulations of long charged stiff rods [36].

Since the monopole–monopole image interaction, \( w_1 \), has the longest range, we can expect that for large separations between the walls, \( 2a \), this interaction will dominate and higher multipoles can again be neglected. Figure 3(c) shows the pressure difference \( \Delta P \), which is the difference between the pressure with finite \( t \) and the pressure without the quadrupolar contributions, \( t = 0 \). As expected, this difference does indeed go to zero at large separations. At moderate separations the quadrupole contribution becomes negative, which is caused by monopole–quadrupole image attractions, \( w_3 \) and \( w_5 \), that are shorter ranged than \( w_1 \). At even smaller separations quadrupole–quadrupole image repulsion, \( w_q \), becomes dominant, since it has the shortest range and varies approximately as \( \sim a^{-5} \). The larger the \( r \), the smaller the maximal quadrupolar attraction at larger distance thus, since it is overwhelmed by repulsive contributions.

7. Multipoles, correlations, bridging and all that jazz

There appear some similarities between the polyelectrolyte bridging interaction [32, 33] and the attraction seen with strongly charged counterrions. Recent simulations and density functional results [36] in fact do accentuate a close connection between polyelectrolyte bridging and strong coupling electrostatics, a line of reasoning that we will explore here in finer detail.

Turesson \textit{et al} [36] indeed find that for flexible polyelectrolytes, bridging attraction appears to be the dominant source of attractive interaction. On the other hand, for stiff charged rods the strong adsorption to the bounding charged walls prevents bridges from forming. Instead a very strong correlation attraction is apparent at shorter separations. Flexibility thus extends the range of attractive interactions as their nature changes from short range correlation attraction to longer ranged polyelectrolyte bridging. Similar conclusions have also been reached in [37].

Here we will formulate the exact correspondence between the intuitive notion of ‘bridging’ in the case of multivalent counterrions and the definition of ‘bridging interaction’ in the case of charged polymers. Let us analyze first the partition function for two disparate systems: a counterrion Coulomb fluid confined between two strongly charged walls, and a flexible polyelectrolyte chain confined in the same geometry with weakly charged walls. For point counterrions between two charged surfaces the grand canonical partition function in the SC limit [38] can be written to the first order in fugacity as

\[
Z = Z^{(0)}_{\text{SC}} + \lambda' Z^{(1)}_{\text{SC}} + \cdots \tag{37}
\]

where

\[
Z^{(0)}_{\text{SC}} = C \int D[\phi(r)] \exp\left( -\frac{\beta}{2} \delta_0 \int \varepsilon(\mathbf{r})(\nabla \phi(\mathbf{r}))^2 d\mathbf{r} \right) + i\beta \int \rho_0(\mathbf{r})\phi(\mathbf{r}) d\mathbf{r} \tag{38}
\]

and

\[
Z^{(1)}_{\text{SC}} = C \int dR_0(\mathbf{R}) \int D[\phi(\mathbf{r})] \exp\left( -\frac{\beta}{2} \delta_0 \int \varepsilon(\mathbf{r}) \right) \times (\nabla \phi(\mathbf{r}))^2 d\mathbf{r} + i\beta \int \rho_0(\mathbf{r})\phi(\mathbf{r}) d\mathbf{r} + i\beta q e_0 \times \int \rho(\mathbf{r} - \mathbf{R})\phi(\mathbf{r}) d\mathbf{r} \right). \tag{39}
\]

The above partition function is of course exactly the same as the one in (22) after one evaluates the functional integrals indicated above explicitly. Here \( \rho_0(\mathbf{r}) \) is again the external fixed charge on the two bounding surfaces located at \( z = \pm \frac{a}{2} \) and the shorthand \( \rho(\mathbf{r} - \mathbf{R}) \) is introduced for \( \rho(\mathbf{r} - \mathbf{R}) = \delta^3(\mathbf{r} - \mathbf{R}) \).

On the other hand the partition function for a flexible polyelectrolyte chain is given by [39]

\[
Z_P = C \int D[\phi(\mathbf{r})] \exp\left( -\frac{\beta}{2} \delta_0 \int \varepsilon(\mathbf{r})(\nabla \phi(\mathbf{r}))^2 d\mathbf{r} \right) + i\beta \int \rho_0(\mathbf{r})\phi(\mathbf{r}) d\mathbf{r} + \ln \int D[\phi(\mathbf{r}; N) d\mathbf{r}] \tag{40}
\]

where \( G_\phi(\mathbf{r}, \mathbf{r}') \) is the Green function of a polyelectrolyte chain in external electrostatic field \( \phi \) given by [40]

\[
G_\phi(\mathbf{r}, \mathbf{r}'; N) = \int_{\mathbf{R}(0)}^{\mathbf{R}(N)} \int \mathcal{D}[\mathbf{R}(n)] \times \exp\left( -\frac{3}{2\ell} \int_0^N \left( \frac{d\mathbf{R}(n)}{dn} \right)^2 dn + i\beta q e_0 \right) \times \int_0^N \phi(\mathbf{R}(n)) dn, \tag{41}
\]

where \( \mathbf{R}(n) \) is the coordinate of a polyelectrolyte segment \( n \) and the boundary condition is set as \( \lim_{N \to 0} G_\phi(\mathbf{r}, \mathbf{r}'; N) = \delta^3(\mathbf{r} - \mathbf{r}') \). Representing the Green function via a corresponding Edwards equation one can derive that the polyelectrolyte monomer density \( \rho_\phi(\mathbf{r}; N) \) is given by a functional derivative of \( \ln \int D[\phi(\mathbf{r}; N) d\mathbf{r}] \) with respect to the fluctuating electrostatic potential \( \phi(\mathbf{r}) \) [40]. The polyelectrolyte monomer density has to satisfy the condition \( \lim_{N \to 0} \rho_\phi(0; N) = \delta(\mathbf{r}) \).

In the planar geometry with two bounding surfaces, one can center the monomer density at the origin of the coordinate system, whose \( z \) axis is directed along the surface normal and has the origin at the mid-plane. Defining \( \mathbf{R}_0 = (0, 0, 0) \) one can thus write \( \rho_\phi(0; N) = \rho_\phi(\mathbf{r} - \mathbf{R}_0; N) \). In the limit of small electrostatic potentials, the lowest order (i.e., the first-order) solution of the polyelectrolyte Green function can be
written as \[40\]
\[
\ln \int G_\phi(r, r'; N) \, \text{d}r \, \text{d}r' \equiv i \beta q e_0 \\
\times \int \rho_{\phi=0}(r - R_0; N) \phi(r) \, \text{d}r.
\] (42)

By construction, this solution clearly corresponds to a weak coupling of the polyelectrolyte to the external field generated by the fixed charges at the boundary of the system. For large values of this charge, higher orders in the expansion (42) would certainly have to be taken into account.

In the case of weak coupling the partition function of a polyelectrolyte chain is then given approximately by
\[
Z_p \equiv Z_{WC}^{(0)}[\rho_{\phi=0}(r - R_0; N)] \\
= C \int D[\phi(r)] \exp\left(-\frac{1}{2} \beta \delta_0 \int \epsilon(r)(\nabla \phi(r))^2 \, \text{d}r \right) \\
+ i \beta \int \rho_0(r) \phi(r) \, \text{d}r + i \beta q e_0 \\
\times \int \rho_{\phi=0}(r - R_0; N) \phi(r) \, \text{d}r. (43)
\]

If we now compare the expressions (39) and (43), we can establish the following identity:
\[
Z_{SC}^{(1)} = \lim_{N \to \infty} \int \text{dR} \Omega(R) \, Z_{WC}^{(0)}[\rho_{\phi=0}(r - R; N)], (44)
\]

with the simple (first-order SC) counterion partition function on the lhs and the polyelectrolyte (zeroth-order WC) partition function on the rhs. This ‘duality’ relation connects the partition function of a strongly coupled point counterion system with the partition function of a weakly coupled flexible polyelectrolyte system. Since \(Z_{WC}^{(0)}\) for finite \(N\) describes polyelectrolyte bridging interactions it is clear that the above formula establishes a formal connection between strong coupling point counterion attractive interactions and weak coupling polyelectrolyte bridging interactions.

8. Conclusions

Using field-theoretic methods we derived a description of the counterion-mediated electrostatic interaction for when the counterions possess internal degrees of freedom such as a rotational axis. We concentrate on the symmetrically charged planar surfaces with a dielectric mismatch on both sides and counterions with quadrupolar moments in between. We analyze this system in two different regimes, namely in the weak coupling (mean field) and the strong coupling limits. The former describes the case of low surface charge and low counterion valency, whereas the latter describes high surface charge and high counterion valency.

In the mean field limit, we derived the Poisson–Boltzmann equation for counterion density in the case where the counterions possess dipolar as well as quadrupolar moments. Since the equation is highly non-linear, we solved it numerically only in the first-order approximation, which should be valid for small quadrupolar moments, \(t\). As is already known, the dielectric mismatches play no role in the mean field theory and therefore have no effect on quadrupoles \[29\]. One could add those corrections in by hand \[41\], but such an approach does not strictly correspond to the mean field analysis. Quadrupolar interaction affects the counterion density distribution as well as interactions between the charged surfaces. Counterions are depleted from the central region and concentrated in the vicinity of both surfaces. The orientation of quadrupoles is preferentially parallel to the \(z\) axis as the moment \(t\) is increased. The highest alignment is reached right next to both bounding surfaces. The quadrupolar contribution to the interaction pressure is attractive.

In the strong coupling limit higher order multipole interactions also play an important role in the intersurface interactions but only for a dielectrically inhomogeneous case. Without any dielectric discontinuities in the system the higher multipoles simply do not matter in the plane-parallel case and the monopolar part of the partition function captures all the strong coupling effects. This is a completely general result for plane-parallel surfaces and should be of some importance in assessing the electrostatically mediated forces between strongly charged planar surfaces. As already mentioned, the multipoles can have effects near curved surfaces even without dielectric discontinuities. In contrast to the case for monopolar counterions, higher multipoles interact only via the orientationally dependent part of the image self-interaction. In our analysis we focused on the counterions with monopolar and quadrupolar moments, so the interaction is composed of three contributions. These are the monopole–monopole image interaction, which is long ranged and repulsive, the monopole–quadrupole image, as well as quadrupole–monopole image interactions that are shorter range and attractive. Finally, the very short ranged quadrupole–quadrupole image contribution is repulsive and plays a dominant role at very small distances.

We are at present involved in carrying out extensive Monte Carlo simulations in order to explore the validity of the formal developments described above. Preliminary results completely vindicate the theoretical analysis.

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References

[1] van Blaaderen A 2006 *Nature* **439** 545
[2] Hong L, Cacciuto A, Luijtjen E and Granick S 2006 *Nano Lett.* **6** 2510
[3] Glotzer S C and Solomon M J 2007 *Nat. Mater.* **6** 557
[4] Jerri H A, Dutter R A and Velegol D 2008 *Soft Matter* **5** 827
[5] Abrashkin A, Andelman D and Orland H 2007 *Phys. Rev. Lett.* **99** 077801
[6] Maset S and Bohinc K 2007 *Phys. Rev. Lett.* **101** 208305
[7] Deraad L L, Milton K A, Tsai W Y and Schwinger J S 1998 *Classical Electrodynamics, the Advanced Book Program* (New York: Perseus Books Group)
[8] Netz R R 2001 *Eur. Phys. J.* **E** 5 557
[9] Podgornik R 1992 *Phys. Rev. E* **5** 557

Moreira A G and Netz R R 2002 *Eur. Phys. J.* **E** 23 265

Jho Y S, Kanduˇc M, Naji A, Podgornik R, Kim M W and Pincus P A 2008 *Phys. Rev. Lett.* **101** 188101

Poon W C K and Andelman D 2006 *Soft Condensed Matter Physics in Molecular and Cell Biology* (New York: Taylor and Francis)

Podgornik R 2004 *J. Polym. Sci.* **B** 42 3539

[30] Turesson M, Forsman J and Åkesson T 2006 *Langmuir* **22** 5734

Turesson M, Forsman J and Åkesson T 2005 *Langmuir* **352** 131

[37] DeRouchey J, Netz R R and Rädler J O 2005 *Eur. Phys. J. E* **16** 17

[38] Netz R R 2001 *Eur. Phys. J. E* **5** 557

[39] Podgornik R 1991 *J. Phys. Chem.* **95** 5249

[40] Doi M and Edwards S F 1986 *The Theory of Polymer Dynamics* (Oxford: Oxford University Press)

[41] Onuki A 2006 *Phys. Rev. E* **73** 021506