Inter–domain dipolar repulsion in lipid monolayers with phase coexistence

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Abstract

A great variety of biologically relevant monolayers present phase coexistence characterized by domains formed by lipids in a long-range ordered phase state dispersed in a continuous, disordered phase. Because of the difference in surface densities the domains possess an excess dipolar density with respect to the surrounding liquid phase. In this work we propose an alternative method to measure the dipolar repulsion for neutral lipid monolayers. The procedure is based on the comparison of the radial distribution function, $g(r)$, from experiments and Brownian dynamic (BD) simulations. The domains were modeled as disks with surface dipolar density, whose strength was varied to best describe the experimentally determined monolayer structure. For comparison, the point dipole approximation was also studied. As an example, we applied the method for mixed monolayers with different proportions of distearoylphosphatidylcholine (DSPC) and dimyristoylphosphatidylcholine (DMPC) and obtained the excess dipolar density, which were in agreement with those obtained from surface potential measurements. A systematic analysis for experimentally relevant parameter range is given, which may be used as a working curve for obtaining the dipolar repulsion in different systems provided that the experimental $g(r)$ can be calculated from a statistically relevant amount of images.

Introduction

In different biomembranes, phase coexistence is frequently observed, depending on the membrane and aqueous composition, temperature and the particular model of membrane (Langmuir monolayers, supported films, free-standing bilayers such as giant unilamellar vesicles or black lipid membranes, among others). When the denser phase is not the continuous phase, domains (named rafts for certain lipid composition) are observed, moving in the more fluid phase. The domains interact between each other, and these interactions affect their own movement as well as the motion of other species present in the membrane. Inter-domain interaction may be related with electrostatic forces (dipolar or Coulombic repulsions), those related with the spontaneous curvature of the coexisting phases and hydrodynamic forces, which appear when domains are in motion. These forces hinder the coalescence of the domains and modulate the availability of the species in the membrane and their dynamics at long time scales. The dipolar repulsion are always present, since the molecules forming the membrane are ordered and dipolar. Coulombic forces appear for charged domains, while curvature effects are important when the spontaneous curvature of the coexisting phases is markedly different, and for large domains with high line tension.

The dipolar repulsion may be estimated by
knowing the difference in dipole density of the surfactants organized at the interface, including the contribution of the hydration water at the polar head group region. A frequently used method for determining the value of the average dipole moment (molecule + hydration water) of a homogeneous film is the determination of the surface potential in Langmuir monolayers at the aqueous/air interface. Alternatively, probes sensitive to the local potential have been used in bilayers, as well as conductance measurements. Therefore, if the composition of the coexisting phases is known, the dipole potential of monolayers or bilayers with the composition of each phase allows estimating the dipole density of each phase, and thus, the difference between them. However, the composition of the coexisting phases is not always easily obtained, particularly for systems with more than two components, thus preventing the estimation of the difference in dipole density from dipole potential measurements. Furthermore, when domains are formed as a consequence of the phase transition of a single-component membrane, the estimation of the dipole potential of each phase at the same temperature and molecular density is not straightforward.

The presence of domains is a common feature in different model membranes, and also in plasmatic membrane of mammals (with nanometer putative sizes), yeast, fungi and plants (of radii in the micrometer range). In these natural systems, the composition of the membrane is highly complex and therefore, it is not possible to know the precise composition of the domains, thus preventing also the estimation of difference in dipole density from dipole potential measurements.

In this work, we propose a novel manner of estimating the inter-domain dipolar repulsion using a passive method that involves the analysis of images of the membrane with phase coexistence. We make use of the fact that the dipolar repulsion between domains promotes a 2-dimensional spatial arrangement of the domains, in which the average domain-domain distance is maximal. Therefore, the repulsion will induce a domain distribution which leads to radial distribution function characteristic of liquid systems. The method is based on the comparison of the pair correlation function obtained from experiments, with Brownian dynamics simulations of a model system. As an example, we determined the difference in dipolar density of a binary monolayer of DSPC/DMPC at the air-water interface from the analysis of the radial distribution of domains and the results are compared with those obtained by surface potential determinations.

### Pair potential and distribution of domains in a monolayer with phase coexistence

We consider a monolayer in its two-phase liquid condensed (LC) and liquid expanded (LE) coexistence region, where the LC phase forms domains in the LE phase that occupies the larger area of the monolayer. Because of the difference in surface densities the LC domains possess an excess dipole density, $\sigma$, with respect to the surrounding LE phase. This originates a dipolar repulsive interactions between the domains. In general in DSPC-DMPC monolayers, the domains exhibit nearly circular shapes. For this reason, we model the mixed monolayer as a 2D–dispersion of circular domains, which interact through a dipolar pair potential, $U_d$. Considering only dipole components perpendicularly oriented to the interface, the repulsive potential can be described by

$$U_d(r) = \frac{\sigma^2}{4\pi \varepsilon_0 \varepsilon} \int A_2 \int A_1 \frac{1}{|\vec{r}_1 - \vec{r}_2|^3} \, da_1$$

where $A_i$ denotes the area of domain $i$, $da_i$ its area element and $\vec{r}_i$ its position vector respect to the domain center, with $i = 1, 2$. $\vec{r}$ is the vector from center of domain 1 to center of domain 2, as shown in Scheme. $\varepsilon_0$ is the vacuum permittivity and $\varepsilon$ is the relative permittivity of the membrane.

[Diagram: Scheme 1: Two domains of equal radii $R$ and excess dipolar density $\sigma$ with center–to–center distance $r$.]
force on domain 1, derived from this potential is
\[ F_d(\vec{r}) = \frac{3\sigma^2}{4\pi\varepsilon_0\varepsilon} \int \frac{d\vec{r}_1}{A_2} \int \frac{d\vec{r}_2}{A_1} \frac{\vec{r}_1 - \vec{r}_2 - \vec{r}}{(|\vec{r}_1 - \vec{r}_2 - \vec{r}|^3 \, da_1}. \]  
\[ (2) \]
This force lies along center–to–center line and its magnitude depends only on the distance \( r \) between the domains. There is no analytic expression for \( F_d(r) \), hence, it must be calculated numerically. However, for the particular case of monodisperse systems (all domain radii equal to \( R \)), Wurlitzer et al.\[^{10}\] found the asymptotic behaviour of this force:
\[ F_d(r) \approx \begin{cases} 
\frac{\sigma^2}{2\pi\varepsilon_0\varepsilon} \frac{1}{\sqrt{r/R-2}} & 2R < r \ll 3R \\
\frac{3\pi\sigma^2 R^4}{4\pi\varepsilon_0\varepsilon} \frac{1}{r^3} & r \gg 3R 
\end{cases} \]  
\[ (3) \]
As expected, for large distances it reduces to the force of two point dipoles \( r^{-4} \). These expressions do not describe the interaction between domains in the experimentally relevant interval \((2 + 0.1)R \leq r \leq 10R\), and hence a numerical integration of Eq. (2) must be performed. This 4D–integral represent much more computational effort in a simulation than an analytical expression. This is the very reason why, in general, the interaction is approximated by point dipoles in the center of the domains with a dipole moment \( \mu \), representing the dipole density over the area of the domain, \( \mu_i = \sigma A_i \). Then, the pair potential for point dipoles perpendicularly oriented to the interface is
\[ U_p(r) = \frac{\mu_1 \mu_2}{4\pi\varepsilon_0\varepsilon} \frac{1}{r^3}, \]  
\[ (4) \]
and the corresponding force over the domain 1 is
\[ \vec{F}_p(\vec{r}) = -\frac{3\mu_1 \mu_2}{4\pi\varepsilon_0\varepsilon} \frac{1}{r^4} \hat{r}, \]  
\[ (5) \]
where \( \hat{r} \) is \( \hat{r} = \vec{r}/r \) the unitary vector in the direction of \( \vec{r} \). For convenience, we define
\[ f_0 = \frac{\sigma^2}{4\pi\varepsilon_0\varepsilon}, \]  
\[ (6) \]
a quantity that characterizes the strength of the forces \( F_d \) and \( F_p \).

Figure 1 shows \( F_d(r) \) (solid line) and \( F_p(r) \) (dashed line) for two circular domains of equal radius \( R \). Superimposed on the numerical solution of Eq. (2), we have added the asymptotic expression of Eq. (3) (dotted line). Note that when the two domains approach to contact \((r = 2R)\) \( F_d \) diverges, while \( F_p \) remains finite. Furthermore, for the range of experimental interest, \( 2R < r < 10R \), the difference between the two forces is appreciable. In particular, even for \( r = 5R \) the difference is of the order of 15%.

The point dipole approximation deviates from the dipolar density force for short and intermediate distances. As a consequence, the structural properties of the monolayer obtained with both schemes have remarkable differences.

A key quantity to characterize the structure of the monolayer is the radial distribution function \( g(r) \). Considering an homogeneous distribution of domains in the monolayer plane, \( g(r) \) represents the probability to find a domain at the distance \( r \) of another domain chosen as a reference point:
\[ g(r) = \rho \left\langle \frac{1}{N} \sum_{i,j=1 \atop i \neq j}^N \delta(\vec{r}_i - \vec{r}_j) \right\rangle. \]  
\[ (7) \]
Here, \( \rho = A/N \) is the number density, \( N \) the number of domains, \( A \) the total monolayer area and the angular brackets indicate an equilibrium ensemble average.

In this work, we use radial distribution function in order to calculate the dipolar repulsion, \( \sigma/\sqrt{\varepsilon} \), by fitting the \( g(r) \) obtained from the experiments with that of the simulations, being \( f_0 \) (i.e. \( \sigma/\sqrt{\varepsilon} \)
Experimental Section

Mixed distearoylphosphatidylcholine (DSPC) and dimyristoylphosphatidylcholine (DMPC) monolayers present a wide range of compositions and lateral pressures where they exhibit phase coexistence. At room temperature, the mixed monolayers show two-phase coexistence, with LC micrometer sized domains dispersed in a LE continuous phase, at a lateral pressure of 10 mN/m and for composition of DSPC higher than 24 mol%.

We take micrographs of mixed monolayers at different DSPC concentration using fluorescence microscopy. We perform the experiments with DSPC, DMPC, and the lipophilic fluorescent probe L-R-phosphatidylethanolamine-N-(lissamine rhodamine B sulfonyl) ammonium salt (chicken egg, trans-phosphatidylated) (Rho-PE) purchased from Avanti Polar Lipids (Alabaster, AL). The water used for the subphase was from a Milli-Q system (Millipore) with a resistivity of 18 MΩ cm and a surface tension of 72 mN/m.

The fluorescent probe (Rho-PE) was incorporated into the lipid solution before being spread at a concentration of 1 mol% or less. Monolayers were formed in a Langmuir trough (Microtrough-XS, Kibron Finland) on subphases of pure water. The lipid mixture was dissolved in chloroform:methanol (2:1) to obtain a solution of 1 nmol/µl, which was spread onto the aqueous surface. We performed the experiments at room temperature $T = (20 \pm 1)^\circ C$, and films were compressed up to a lateral pressure $\pi = (10 \pm 1)$ mN/m determined with a Pt plate using the Wilhelmy method.

After spreading the lipid layer on an area 1.5 times the lift-off, the subphase level was reduced to a thickness of about 3 mm, in order to minimize convection. The Langmuir balance was placed on the stage of an inverted microscope (Axiovert 200, Zeiss) equipped with a CCD IxonEM+ model DU-897 (Andor Technology) camera, a 100x objective, a continuous solid state laser (TEM00, 532 nm up to 200 mW, Roithner Lasertech), and rhodamine emission filters. The fluorescent probe partitions preferentially on the LE phase and therefore the domains formed by lipid in a condensed phase appear darker in the images.

The surface potential was determined in Langmuir monolayers using the KSV NIMA Surface Potential Sensor (Helsinki, Finland) with the vibrating plate method. The films were prepared with pure DSPC (composition of the condensed state$^3$) or with a mixture of DSPC and DMPC with 24 mole% of DSPC (composition of the expanded phase$^3$).

Image analysis and Radial distribution function

From the micrographs we calculated the condensed area fraction, $\phi$, defined as the ratio of the area occupied by the domains over the monolayer area. To do this, we determined the amount of each phase, converting the original gray scale images into black/white images using the image processing software ImageJ. Then, the total area occupied by the black regions, which corresponds to the area occupied by domains, was determined.

To process the images we removed the slightly nonuniform illumination in the images (due to the intensity distribution across the laser beam profile) using a band-pass filter. Then, we selected a particular gray scale level, and all pixels with intensities above this threshold were converted to “white”, while pixels with intensities below this threshold level were converted to “black”. The value of the threshold level was determined on the basis of an optimal resolution of the structures by performing a constant eye comparison with the original photo. The threshold value must be carefully selected, since it determines the principal source of error for $\phi$. Different threshold values change the size of the domains; thus, the determination of $g(r)$ is not significantly affected, since it only depends on the position of the domain centers. However, selection of a low threshold value may led to an underestimation of the number of domains, since the smallest domains (with a domain area less than 4 pixels) appear lighter than the larger ones. Therefore, the threshold value was selected in order not to modify the total number of domains in more than 10%.

We calculate the radial distribution function for each monolayer as an histogram of domain center–to–center distance $r$. For each condition the or-
der of 1000 micrographs were used of a size of 122 × 122 μm², and a binning of 0.5 px (0.12 μm) was selected. This size is bigger than the error in r, Δr = 0.3 px (0.07 μm), and is small enough to obtain a well characterized curve. The error of g(r) is calculated independently for each value of r using the standard deviation.

Simulations

We model the mixed monolayer as a two dimensional Brownian suspension of interacting hard disks of equal radii (monodisperse) immersed in an effective fluid, each disk representing an idealized lipid domain. The inter–domain interactions are described by the full dipolar density pair potential given by Eq. (1) plus a hard core repulsive part. The same system is also studied using the point dipole potential Eq. (4), in order to analyze the validity of this approximation which is easier to implement and widely used.

To study the static properties of the two mixed monolayer models we performed Brownian dynamics (BD) simulations. In this scheme, the finite difference equation describing the in-plane displacement of N identical Brownian disks immersed in a fluid during the time step Δt is given by [17]

\[
\vec{r}_i(t + \Delta t) - \vec{r}_i(t) = \sum_{j=1}^{N} \frac{D_0}{k_B T} F^P_j \Delta t + \vec{X}_i, \tag{8}
\]

where \(F^P_j\) is the direct total force on disk j due to all other N-1 disks, \(D_0\) the disk diffusion coefficient, \(k_B\) the Boltzmann constant, \(T\) the temperature, and \(\vec{X}_i\) a random displacement vector of particle \(i\) originated from solvent particle collisions. \(\vec{X}_i\) is sampled from a Gaussian distribution with zero mean and covariance matrix:

\[
\left\langle \vec{X}_i \right\rangle = 0; \left\langle \vec{X}_i \vec{X}_j \right\rangle = 2D_0 \mathbb{I} \delta_{i,j} \Delta t \tag{9}
\]

where \(\mathbb{I}\) is the identity matrix, and \(\delta_{i,j}\) the Kronecker delta.

The inter–domain dipolar density force, Eq. (2), was calculated in advance, and the values were tabulated for later use in the simulations. The 4d–integral in Eq. (2) was calculated using the Monte Carlo algorithm for \(2.003 < r/R < 22\), and the analytical expressions for the asymptotes, Eq. (3), were used outside this range.

The simulated systems consisted of \(N\) disks of radius \(R\) under periodic boundary conditions, using the minimum image convention. The size of the simulation box, \(L\), was determined using the expression of the condensed area fraction \(\phi = N\pi R^2 / L^2\).

In our simulations, we use \(\phi\) determined from the monolayer micrographs, \(N = 144\) disks and \(\Delta t = 2 \times 10^{-4} R^2 / D_0\). The remaining parameter to determine the system, \(f_0\), is systematically varied in order to find the best agreement with the experimental g(r).

We verified that, for the studied systems, there is no system size dependency in the structural quantities. The accuracy of our BD simulation method was tested for specific examples by comparison with published simulation data on 2D systems.[18]

Results and Discussions

We analyzed monolayers of three DSPC:DMPC compositions with different area fraction. We used a pure water subphase, 10 mN m⁻¹ lateral pressure, and DSPC mol %: 40, 50 and 60. A representative micrograph of each monolayer is presented in Figure 2. The monolayers were photographed at 12.1 frame/s for 100 s, during while domains suffer Brownian motion and the monolayer is subjected to drift, allowing to image different regions of it. The condensed area fraction was calculated for each frame using the process described in Experimental Section. Then, the value of \(\phi\) was obtained from the average. From the frame area, \(a\), and the number of domains in the micrographs, \(n\), the number density was calculated as \(\rho = n/a\).
The values of $\phi$ and $\rho$ for the considered systems are shown in Table I. The domain radius distributions were determined from the domain sizes (areas) assuming circular domains. The corresponding histograms are shown in Figure [3], and the average radii, $\bar{R}$, are presented in Table [I].

Table 1: Condensed area fraction, number density, average radius and effective radius for the monolayers considered. Errors are determined by standard deviation, except for $\phi$ which is determined varying the threshold values, as explained in Experimental Section.

| DSPC±2 mol% | $\phi \pm 0.02$ | $\rho \pm 0.003 [\mu m^{-2}]$ | $\bar{R} [\mu m]$ | $R_{eff} [\mu m]$ |
|-------------|-----------------|---------------------|-----------------|-----------------|
| 40          | 0.17            | 0.028               | 1.2 ± 0.6       | 1.4 ± 0.2       |
| 50          | 0.20            | 0.050               | 1.0 ± 0.5       | 1.1 ± 0.1       |
| 60          | 0.23            | 0.122               | 0.7 ± 0.3       | 0.77 ± 0.07     |

Figure 3: Domain radius distribution of monolayers with different condensed area fraction.

In a monodisperse model the area fraction, the number density and the domain radii are not independent quantities. Since $\phi$ and $\rho$ are more straightforwardly calculated than the average radii, we determine the effective radius from these quantities as $R_{eff} = (\phi / (\pi \rho))^{1/2}$. Note that this radius is slightly different from the average domain radius, due to the fact that $R_{eff}$ is equivalent to the square root of the second moment of the radius distribution (see Table [I]).

Figure 4 shows the radial distribution functions for the three experimental systems. In all cases, the $g(r)$ resembles that of a liquid, showing a well defined first peak (at $r_{max}$), which characterizes the first coordination shell, i.e., the nearest neighbors. Besides, a marked first minimum and a second maximum, which corresponds to the second coordination shell, is observed for the systems with $\phi = 0.20$ and $\phi = 0.23$. Due to the “strong” repulsive interaction, the nearest neighbor distance is much larger than the average domain diameter. For this reason the hard core repulsion is not determinant for the structure, and the characteristic length of the system is the mean geometric distance $r_m = \rho^{-1/2}$.

In order to calculate the dipolar repulsion, $\sigma / \sqrt{\epsilon}$, we fitted the experimental $g(r)$ to the BD one by varying the interaction strength $f_0$. The best fit is selected by matching the height of the first peak. Due to the polydispersity in domain sizes, the experimental $g(r)$ has, in general, a broader first peak and a shallower first minimum. For this reason, we used the first peak height as fitting criteria instead of fitting the whole curve.

Figure 5 shows the experimental $g(r)$ compared with the BD simulations with the full dipolar density interaction, Eq. (2). Here, for convenience, the distance $r$ is scaled by the mean geometric distance, $r_m$. For the experimental $g(r)$, we selected the value of $r_m$ from the 90% confidence interval that best matches the peak position of the BD $g(r)$. In general we observe a remarkable agreement between experiments and simulation. Simulations results overestimate the first minimum depth and underestimate the first peak width. Both differences might be attributed to polydispersity effects, not taken into account in our BD simulations.

The fitting procedure, that consists in matching
For the studied systems the values of $\sigma/\sqrt{\varepsilon}$ are summarized in Table 2 where the uncertainties were determined from $\Delta f_0$ discussed above.

In order to compare the dipolar density obtained with other experimental techniques, the value of the relative permittivity of the membrane $\varepsilon$ is needed. Several studies were done to determine the relative permittivity in the interfacial region, obtaining values that lie between the characteristic one for the membrane interior and that of the bulk solution, and depends also on the precise site and technique of measurement. In the present work we use the approximated value of 2, which leads to $\sigma = (0.8 \pm 0.3) \times 10^{-12}$ C/m, $(1.6 \pm 0.3) \times 10^{-12}$ C/m and $(2.1 \pm 0.3) \times 10^{-12}$ C/m for the monolayers with $\phi = 0.17$, 0.21 and 0.23, respectively. The order of magnitude of the dipolar density obtained are in agreement with previous result obtained from domain migration under an externally applied electric field.

We performed measurements of the surface potential, $\Delta V$, in two homogeneous monolayers; one in the LE phase and the other in the LC phase, both with the same proportions of DSPC as in the analyzed experiments, as explained in the experimental section. We obtained $\Delta V_{LC} = (555 \pm 12)$ mV and $\Delta V_{LE} = (402 \pm 8)$ mV, for the pure LC phase and pure LE phase, respectively. These values are in agreement with previous results for monolayers with the same and different subphase, and for bilayers.

Using the parallel plate capacitor model, the excess dipolar density can be obtained from the difference of the surface potential, $\Delta(\Delta V) = \Delta V_{LC} - \Delta V_{LE}$, through the equation $\sigma = \Delta(\Delta V)\varepsilon_0\varepsilon$ and assuming the same value for the relative permittivity in both phases. The resulting dipolar density obtained from this model is $\sigma_c = (2.7 \pm 0.4) \times 10^{-12}$ C/m, consistent with the results obtained with the method proposed here.

| $\phi$ | $f_0 [k_BT/R_{eff}]$ | $f_0 \times 10^{-14}[N]$ | $\sigma/\sqrt{\varepsilon} \times 10^{-12}[C/m]$ |
|-------|----------------------|--------------------------|-----------------------------------------------|
| 0.17  | 1.2 \pm 0.9          | 0.3 \pm 0.2              | 0.6 \pm 0.2                                   |
| 0.20  | 3 \pm 1              | 1.1 \pm 0.4              | 1.1 \pm 0.2                                   |
| 0.23  | 3.7 \pm 0.8          | 1.9 \pm 0.4              | 1.5 \pm 0.2                                   |

We observe that the obtained values for the dipole...
lar repulsion, $\sigma/\sqrt{\epsilon}$, are indistinguishable, except for the monolayer with $\phi = 0.17$. This could be attributed to the fact that this monolayer has a larger size dispersion and presents characteristics compatible with a bimodal distribution. In general, unimodal size polydispersity affects the peak height and width of the $g(r)$, while for more complex size distributions these effects are pronounced, and eventually could also lead to a splitting of the first peak. In particular, for the monolayer with $\phi = 0.17$ these effects are responsible for the fact that the fitted $g(r)$, obtained with a monodisperse model, represents a much less interacting system. As a consequence, the fitting procedure results in an underestimated value of $\sigma/\sqrt{\epsilon}$.

**Radial distribution of domains as a tool for the determination of the inter-domain dipolar repulsion: working curve**

With the aim of obtaining the dipolar repulsion for a range of area fraction and interaction strength we performed simulations and analyzed the dependence of $g(r_{\text{max}})$ with $\phi$ and $f_0$. Figure 6 shows $g(r_{\text{max}})$ as a function of $f_0$ in units of $k_B T / R_{\text{eff}}$ for three representative values of the area fraction.

These results may be used to estimate $\sigma/\sqrt{\epsilon}$ from the experiments without performing simulations. For this purpose, first $\phi$, $\rho$ and $g(r)$ are determined, then $g(r_{\text{max}})$ and $R_{\text{eff}}$ are calculated, and finally $f_0$ is obtained using Fig. 6. Note that the value of $\rho$ is used only to calculate $R_{\text{eff}}$, needed to express $f_0$ in physical units.

**Point dipole approximation**

A widely used approximation for the domain interactions is to consider each domain as a point dipole with excluded volume. This scheme has the advantage of having a simple analytic expression for the interaction force, Eq. (5), hence being easier to implement than the full dipolar density interaction. To assess the validity of this approximation in the considered area fraction range, we systematically studied the structural properties of the point dipole system, and compared them to the corresponding dipolar density system. Figure 7 shows $g(r)$ for both schemes using the same set of parameters [$\phi, f_0, \rho$]. It is clearly observed that the point dipole approximation does not represent properly the structure; in all cases it underestimates the pair correlations. We found that a good agreement can be achieved if $f_0^p$ of the point dipole model is considered as an effective interaction strength, and adjusted to match the first peak height of the dipolar density $g(r)$. This is shown in Fig. 8 for the three systems analyzed in the previous section. Slight differences are observed in the first peak position and in the small region where $g(r)$ starts to deviate from zero. The agreement improves when the systems are more structured. In particular, for weak interacting domains the differences start to be noticeable. These deviations arise from the fact that the point dipole force tends to a finite value at contact, while in the dipolar density case it diverges. For this reason, the hard disk interaction becomes relevant for the point dipole model.

As expected $f_0^p$ is always larger than $f_0$. Therefore, even if a good agreement is achieved, fitting the experimental data with the point dipole model leads to an overestimated value of the dipolar repulsion. For the monolayer with $\phi = 0.20$, fitted with $f_0^p = 6.6 R_{\text{eff}} / (k_B T)$, a dipolar repulsion $\sigma/\sqrt{\epsilon} = 1.6 \times 10^{-12} C/m$ would result, that is 45% larger than the one calculated with the full dipolar density model (see Table 2).

Due to the point dipole model simplicity, it is
Figure 7: Radial distribution functions obtained from BD simulations using dipolar density and point dipole interactions. For each area fraction the value of $f_0$ is the same for both models, and is given in Table 2. For clarity the data have been vertically shifted.

It is worthwhile to look for a correspondence between both models. With this purpose, we determined the effective interaction strength, $f_p^0$, for a set of $f_0$ at three different area fractions. For the range of parameters of interest the relation between $f_p^0$ and $f_0$, as can be seen in Fig. 9. This figure can be helpful for selecting the appropriate $f_p^0$ in order to compute the structural properties of a monolayer by only performing simulations based on the simpler point dipole model. Alternatively, if the experimental pair correlation function is modeled by the point dipole approximation, the value of $\sigma/\sqrt{\varepsilon}$ can be calculated using the corresponding $f_0$ estimated from this figure.

**Conclusions**

In the present work, the inter–domain dipolar repulsion in a mixed monolayer was calculated from the experimental radial distribution functions using Brownian dynamics simulations. The domains were modeled as monodisperse disks with constant dipolar density, and their pair interaction obtained from Monte Carlo integration. We found good agreement between experiments and simulations for the structure of the monolayers, where $\sigma/\sqrt{\varepsilon}$ was used as fitting parameter. Mixed monolayers with three different condensed area fractions were analyzed.

With the exception of the experiment with $\phi = 0.17$, the obtained values of $\sigma/\sqrt{\varepsilon}$ were indistinguishable. This finding is in agreement with the fact that the studied monolayers are located in the regime of condensed area fraction where the lever rule is valid, and hence no $\phi$–dependence of $\sigma$ is expected. Furthermore, the dipolar density value obtained is of the same order of magnitude as the obtained with the parallel plate capacitor model. The dipolar repulsion obtained from the experiment with $\phi = 0.17$ has a lower value. This can be attributed to the fact that this monolayer has a broad and non–unimodal size distribution, whose effects are not taken into account in the simulations. Here, for simplicity, we have modeled the domains as monodisperse disks, while experimental monolayers show a distribution of sizes. It is known that this assumption induces a “softening”
of the pair correlation function due to averaging over different particle sizes. These effects are expected to be relevant specially when the size distribution presents a complex structure or has large variance. The inclusion of polydispersity in our scheme is feasible, and is left for future work.

We have systematically studied the dependence of \( g(r_{\text{max}}) \) with the interaction strength and area fraction. The results, summarized in Figure 9, can be used as a working curve to estimate \( \sigma / \sqrt{\varepsilon} \) directly from the experimental data without implementing any simulation, for the range of parameters studied.

We have also studied the point dipole model to probe its validity as an approximation to the dipolar density model. Analyzing the radial distribution function, we found that the structure is strongly underestimated by the point dipole model. However, if the point dipole strength is considered as a fitting parameter, i.e. as an effective interaction strength, a good agreement can be achieved. For different area fractions we determined \( f_0^p \) for a wide range of \( f_0 \). This relation between the models allows the use of the simpler point dipole model both, to calculate the structure for a monolayer with known dipolar repulsion, and to obtain the dipolar repulsion from an experimental \( g(r) \).

We conclude that the proposed procedure to estimate \( \sigma / \sqrt{\varepsilon} \) from the experimental pair correlation function constitutes an alternative to conventional methods, with the advantage that it only requires monolayer micrographs.

The proposed method may be used not only for binary lipid monolayers but also for single-component monolayers and for more complex monolayers (such as those prepared from isolated membranes, i.e. with all the original components of the cell membrane) and also for bilayers, provided that a time-averaged radial distribution can be obtained, and that the predominating interactions corresponds to those derived from dipolar repulsion. Other functional forms may also be explored in a similar manner as that shown here.

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