Review

Elastic deformation and area per lipid of membranes: Atomistic view from solid-state deuterium NMR spectroscopy

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This article reviews the application of solid-state 2H nuclear magnetic resonance (NMR) spectroscopy for investigating the deformation of lipid bilayers at the atomistic level. For liquid-crystalline membranes, the average structure is manifested by the segmental order parameters (S2CD) of the lipids. Solid-state 2H NMR yields observables directly related to the stress field of the lipid bilayer. The extent to which lipid bilayers are deformed by osmotic pressure is integral to how lipid–protein interactions affect membrane functions. Calculations of the average area per lipid and related structural properties are pertinent to bilayer remodeling and molecular dynamics (MD) simulations of membranes. To establish structural quantities, such as area per lipid and volumetric bilayer thickness, a mean-torque analysis of 2H NMR order parameters is applied. Osmotic stress is introduced by adding polymer solutions or by gravimetric dehydration, which are thermodynamically equivalent. Solid-state NMR studies of lipids under osmotic stress probe membrane interactions involving collective bilayer undulations, order-director fluctuations, and lipid molecular protrusions. Removal of water yields a reduction of the mean area per lipid, with a corresponding increase in volumetric bilayer thickness, by up to 20% in the liquid-crystalline state. Hydrophobic mismatch can shift protein states involving mechanosensation, transport, and molecular recognition by G-protein-coupled receptors. Measurements of the order parameters versus osmotic pressure yield the elastic area compressibility modulus and the corresponding bilayer thickness at an atomistic level. Solid-state 2H NMR thus reveals how membrane deformation can affect protein conformational changes within the stress field of the lipid bilayer. This article is part of a Special Issue entitled: NMR Spectroscopy for Atomistic Views of Biomembranes and Cell Surfaces. Guest Editors: Lynette Cegelski and David P. Weliky.

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1. Biomembranes as liquid-crystalline materials

Cellular membranes fulfill a multitude of biological roles that involve the synergy of diverse lipids, proteins, peptides, and carbohydrates. To describe the functions of biomembranes at the molecular level can be daunting even to experienced investigators. Given the myriad of lipids and proteins that exist, how can we begin to understand the interactions among the various membrane constituents? Such questions are addressed by solid-state nuclear magnetic resonance (NMR) spectroscopy, which is among the paramount methods used for studies of biomolecular structure and dynamics. Many new aspects pertinent to interactions among the lipid [1] and protein molecules can be uncovered by solid-state NMR [2], including how membranes respond to external perturbations associated with their functional mechanisms. Here, our focus is on the application of solid-state NMR spectroscopy for studying the dynamical structure of membrane lipid bilayers, with an emphasis on the role of water and osmotic stress [3]. By combining modern NMR methods with well-established concepts from surface chemistry and physics, new insights into the functional mechanisms of biomembranes can be achieved.

Notably, the length scale over which lipid–protein interactions and related properties begin to emerge falls between the atomistic and macrosopic dimensions [4]. To describe how a biomembrane system behaves at the mesoscopic level is not immediately obvious; yet this behavior underlies their roles in a host of biological phenomena. Evidently, two avenues can be taken: the first involves molecular dynamics simulations, either all atom [5–7] or coarse grained [8–10], whereby a finite number of molecules is described in atomic detail in terms of a molecular force field [11,12]. The second is a continuum description [1], in which lipid membranes are treated as liquid–crystalline materials, so that molecular information is relinquished in favor of material properties [13,14]. The two avenues do not conflict or compete with each other—rather they end up at the same place. Actually there is a third way, one that combines the atomistic observables from NMR spectroscopy with a continuum material science viewpoint [1]. The new view takes cognizance of material properties of biomembranes as they emanate from the atomistic or molecular-scale interactions, due to their lipid and protein composition. Accordingly, biophysical studies of membrane lipids [15–18] are fundamental to understanding membrane protein structure and function [19,20]. In applications involving solid–state 2H NMR spectroscopy, the average structure of the membrane bilayer is manifested by the segmental order parameters (ScD) of the lipids. The NMR order parameters are relevant to calculating the area per lipid, corresponding to the mean-square fluctuations of the molecules. Knowledge of structural quantities such as the cross-sectional area per lipid is important for molecular dynamics simulations of lipid bilayers [5,21] and biomembranes [11,12,22]. One can then address the question of how membrane lipids are affected by their interactions with proteins or peptides [23–25], and/or changes in thermodynamic state variables such as osmotic [3] or hydrostatic pressure [26].

Our paper reviews how solid-state NMR spectroscopy can help us achieve a more complete view of membrane lipids [16,17], proteins and/or peptides [23–25,27–29], and carbohydrates [30] in biological function. For lipid membranes, we show how atomistic NMR observables describe the structural remodeling of the lipid ensemble due to interactions with water [3]. First, we summarize how solid-state 2H NMR of deuterated lipids can help fill the gap between molecular structures and the dynamic stress fields in biomembranes [4]. Using the residual quadrupolar couplings (RQCs) as model-free experimental observables, order parameters are derived for the flexible lipid molecules. These quantities are related to the area per lipid, volumetric bilayer thickness, and balance of attractive and repulsive forces within the membrane. Next, we show how solid-state 2H NMR spectroscopy studies the interactions of biomembranes with water. Removal of water by osmolytes such as polyethylene glycol (PEG) yields a striking increase of the absolute order parameters, due to a reduction of the interfacial area occupied per lipid. The order parameters approach the values seen for the liquid-ordered (lo) phase of bilayers containing cholesterol in raft-like lipid mixtures [31]. Third, from the dependence of the RQCs on the osmolyte concentration (osmotic pressure), we obtain the elastic area compressibility modulus as a quantity that can affect the energetics of proteins within the stress field of the lipid bilayer. Solid-state 2H NMR quantifies the emergence of bilayer elasticity and deformation at an atomistic level by a mean-field description of the forces. Employing a mean-torque analysis of the NMR observables, we calculate that the mean area per lipid and the volumetric bilayer thickness change by up to 20% upon introduction of osmotic stress. Molecular-level forces associated with the lipids can thus play a significant role in biological processes involving lipid–protein interactions, as in the case of mechanosensation or signaling by G-protein-coupled receptors (GPCRs).

2. Implications of membrane deformation due to osmotic stress

The ability of lipid bilayers to transduce physical deformations into useful biological work has been the subject of considerable attention, starting from earlier research [20] and continuing well into the present [3,4,16–18,32–35]. How the shape-inducing properties of lipids [4] affect the functions of various membrane peptides [23,36], G-protein-coupled receptors [19,37–39], aquaporins and ion channels [40–43], and other membrane proteins [44] is at the leading edge of biophysical research [45]. Because the activity of membrane proteins underlies so many biological functions, the effect of external forces such as osmotic pressure on the lipid bilayer matrix is often overlooked or neglected. Using atomistically resolved methods such as NMR spectroscopy, the effect of osmotic pressure can be gauged in the context of bilayer deformation and lipid–protein interactions. Among the relevant structural parameters, the area per lipid at the bilayer interface with water figures prominently [34]. Adopting the area per lipid as a structural measure [3], the question then becomes: do membrane lipid bilayers deform appreciably [46–48] or not at all [49] in response to osmotic pressures in the biological range? Another related aspect is that the area per lipid [34,50] is central to molecular simulations of biomembranes [51,52] and pure lipid bilayers [32–34,53–56]. Establishing the proper initial values and boundary conditions is essential for the validation of simulation outcomes. The area per lipid also gives us a quantitative measure of structure in connection to protein-mediated functions of biomembranes—e.g., through the area elastic modulus κA, the Helfrich spontaneous curvature κ0, and bending rigidity κG, and additional elasticity parameters [57].

Osmotic stress is an effective way to control the hydration of biological specimens [58–60], enabling the measurement of membrane forces involving bilayer undulations, collective order–director fluctuations, and molecular protrusions. Bilayer undulations involve relatively large intermembrane distances, whereas protrusions act over shorter distances,
and between these limits area deformation occurs. Investigations of lipid structural properties are valuable because cellular functions can be modulated through nonspecific lipid–protein interactions [46,51,62]. It is important to understand how membrane structures can deform, and how their hydration state is modified under osmotic stress, which can give insight into the hierarchy of membrane forces [63]. Crowded biological environments can exert a significant osmotic pressure on biomolecular structures [64]. Osmotic pressures can occur due to the competition of various molecular species for available water, and by selective partitioning of solutes across lipid membranes. At the molecular scale, osmotic stress corresponding to pressures on the order of 50–100 atm can significantly affect mechanosensitive ion channels [65,66], as well as G-protein-coupled receptors like rhodopsin [67].

The question of whether lipid bilayers deform [46–48] or not [49] in response to osmotic pressures in the biological range might appear as a clear-cut question—until one realizes the relevant lipid structural parameters are actually quite difficult to measure experimentally. As a rule, X-ray and neutron scattering [49,68–75] are often considered the methods of choice, whereby positional correlations can be accessed directly. But for lipid bilayers in the liquid-crystalline state, scattering peaks are broadened or suppressed due to pronounced membrane shape fluctuations [68,76]. This effect leads to a loss of resolution in the reconstructed electron density profiles, as discussed by Nagle, Tristram-Nagle, and coworkers [69]. Under certain conditions [70] reconstructed electron densities might appear insensitive to applied osmotic stress. Yet a detailed analysis of structural data involving fluctuation corrections indicates that remodeling occurs over a whole range of osmotic pressures [77]. In particular, an X-ray scattering method due to Luzzati [71] does not use electron densities, but relies instead on gravi-metric measurement of water content [78], and has shown a limited range of deformation at high osmotic pressures. The issue of sample inhomogeneity has bedeviled this method, however, e.g. see the discussion by Gawrisch and coworkers [48]. To what extent lipid bilayer remodeling occurs in response to external forces continues to remain in a somewhat uncertain state.

In this context, solid-state $^2$H NMR spectroscopy has long been regarded as one of the premier biophysical techniques applicable to lipid bilayers and biomembranes [79]. One of our aims is to highlight the potential of solid-state $^2$H NMR for the study of membrane structural deformations and molecular fluctuations [3]. Unlike X-ray scattering, it does not measure positional correlations. As an example, Fig. 1 shows a comparison of experimental data from $^2$H NMR spectroscopy with small-angle X-ray scattering (SAXS) results. In particular, an X-ray scattering method due to Luzzati [71] does not use electron densities, but relies instead on gravimetric measurement of water content [78], and has shown a limited range of deformation at high osmotic pressures. The issue of sample inhomogeneity has bedeviled this method, however, e.g. see the discussion by Gawrisch and coworkers [48].

Solid-state $^2$H NMR spectroscopy monitors the orientational dynamics of the lipid molecules, giving information about the lipid chain packing, from which the dynamical structure can be investigated [34,76,79]. The area per lipid can be calculated from the orientational order parameters of the C–$^2$H bonds of deuterium-labeled acyl chains. Knowledge of the statistical chain travel along an axis perpendicular to the bilayer interface with water is needed [34,80]. From the response to osmotic pressure, the material constants are then evaluated for the membrane deformation. Such structural measures can also be used to experimentally validate molecular dynamics (MD) simulations [81,82] of lipid systems and biomembranes [53,54,83]. The molecular force fields encapsulate the data obtained with different experimental techniques [84]. Indeed one of the most fundamental properties of a lipid bilayer—and one of the most common ways to assess whether the system has achieved equilibrium in molecular simulations—is the area per lipid [34,85]. When the area per lipid reaches a stable value, most other structural properties of the lipid bilayer do not change, and the system is viewed as having converged [86]. Because of this feature, the solid-state $^2$H NMR approach plays to an even larger audience than addressed here.

3. Relation of membrane structure to observables from solid-state NMR spectroscopy

3.1. Membrane geometry

Upon hydration, the lipid molecules form a multilamellar dispersion due to their amphiphilic nature, involving the hydrophobic effect.
Nevertheless, in this article we do not focus on aspects involving the shape of the membrane, including the curvature in the membrane. The half-water thickness on either side of the bilayer is half the total thickness of the bilayer hydrocarbon region [34]. The lipid polar head groups contribute to the conformation of a lipid bilayer is depicted, in which water surrounds the lipid membrane) [89] is \( A \) and \( B \), which together with the number of lipids \( N_L \) give the overall surface area of the membrane. Changes in equilibrium structural quantities due to bilayer stress give a membrane-based view of the forces that underlie lipid interactions within the bilayer.

**Fig. 2.** Lipid bilayer showing schematic depiction of unit cell and structural measures obtained from solid-state \(^1\)H NMR spectroscopy and small-angle X-ray scattering. Lamellar structure of the phospholipid membrane is shown with the pertinent structural quantities. The lamellar repeat spacing \( D \) is the sum of the interlamellar water distance \( D_W \) and the bilayer thickness \( D_B \). Here \( D_B \) is the hydrocarbon thickness per bilayer leaflet and \( D_B \) is the head group layer thickness. Bilayer dimensions involve the average cross-sectional area per lipid \( (A) \), which together with the number of lipids \( (N_L) \) give the overall surface area of the membrane. Changes in equilibrium structural quantities due to bilayer stress give a membrane-based view of the forces that underlie lipid interactions within the bilayer.

Together with the van der Waals forces, a geometrical representation of a schematic phospholipid membrane is provided in Fig. 2. A small portion of a lipid bilayer is depicted, in which water surrounds the lipid polar head groups [87], and partially penetrates the bilayer up to about the level of the glycerol backbone [88]. This director is called interlamellar water, and its total thickness \( D_W \) corresponds to the level of the phospholipid region [34]. The lipid bilayer membrane is seen to consist of the lipid head groups confined to a layer of thickness \( D_B \) (phosphate groups are depicted as filled spheres) facing toward the water. The half-water thickness on either side of the bilayer is \( D_W \) and \( D_C \) designates the volumetric half-thickness due to the acyl chains (rendered by flexible sticks that project away from water toward the bilayer center). A soft multimellar lattice is formed, as illustrated by the electron density profile in Fig. 1(b).

In Fig. 2 the bilayer thickness is denoted by \( D_B \) and can be written as a sum of the hydrocarbon chain thickness \( (D_C) \) and the head group thickness \( (D_H) \). Note that the volumetric chain thickness \( D_C \) is one-half of the total thickness of the bilayer hydrocarbon region [34]. The total interbilayer distance \( (D) \) is the distance from the lower half of the water to the upper half of the water on either sides of the membrane) [89] is \( D = D_B + D_W = 2(D_C + D_H) + D_W \). For liquid-crystalline bilayers, we are interested in extracting quantitative structural information, such as the area per lipid. The cross-sectional area per lipid \( (A) \) is related to the total volume \( V \) of the lipid unit cell through the relation \( V = AD \) where \( V = 2V_C + V_W \) and \( V_C \) is the lipid volume [90]. However, experimentally it is challenging to determine the area per lipid from scattering methods. One has to rely on the length of the 1D unit cell (designated as the interbilayer distance \( D \)) as the single structural measure. Other important properties involve the shape of the membrane, including the curvature in 2D [4,91,92]. Nevertheless, in this article we do not focus on aspects such as curvature deformation [4,20].

Let us now decompose the total volume of the lipid as \( V_L = V_H + 2V_C \) where \( V_C \) and \( V_H \) are the volumes of one of the hydrocarbon chains (assumed identical) and the lipid head group, respectively (see Fig. 1). Keep in mind that \( D_B \) is constant for a given lipid head group type (9 Å for phosphocholine group) [47,93] and \( D_B \) is likewise constant for a given hydration level. The water thickness is \( D_W = 2NW_\text{vwater}/(A) \), where \( NW_\text{vwater} \) is the total number of waters of hydration per lipid molecule. For neutral phospholipids there are approximately \( NW_\text{vwater} = 18 \) water molecules at full hydration [3]. In addition, \( NW_\text{vwater} \) is the molecular volume of water (30.3 Å³) [34] and \( (A) \) is the area per lipid. We are left with the hydrocarbon thickness \( D_C \), which is related to the hydrocarbon chain volume \( V_C \) through the relation \( D_C = 2V_C/(A) \). Often it is assumed that the volume of the hydrocarbon chains of a membrane bilayer is approximately incompressible [34], and hence it is essentially constant.

Because lipid membrane systems have many internal degrees of freedom, we can measure only ensemble or time-averaged quantities. In particular, the hydrocarbon thickness \( D_C \) is not the same as the average hydrocarbon chain length (see below) [34]. Rather, the two quantities are related by the orientational distribution function for the various acyl segments. The acyl segment orientation is distributed with respect to the bilayer normal (director), in the liquid-crystalline (or liquid-disordered, \( l_d \) state, and so we consider the various acyl segment projections. Fig. 3 enables us to see the methylene chain travel from the head group-water interface toward the bilayer center. The head groups are shown by the large open spheres, with the irregular lines depicting the acyl chains, and the terminal methyl groups designated by the small filled spheres. Upon replacing hydrogen (H) by deuterium, the orientations of the carbon–deuterium bond segments to the bilayer normal allow us to quantify the chain travel away from the water-lipid interface.

Part (a) of Fig. 3 shows how the mean projection \( (L_C) \) of the acyl lengths onto the lamellar normal corresponds to the average end-to-end distance of the tethered acyl chains. Due to the chain terminations, together with the restraint that hydrocarbon density is conserved, the mean end-to-end distance of the chains is not the same as the volumetric bilayer half-thickness [34]. Rather, the terminal methyl groups are broadly distributed along the bilayer normal (director axis), because the individual acyl chains terminate at different lengths from the water interface. Clearly \( D_C \) and \( (L_C) \) are not equivalent—the volumetric thickness must be calculated from the area per lipid at the aqueous interface, and not the projections along the hydrocarbon chain [34].

Referring to part (b) of Fig. 3, we show how the segmental ordering parameters correspond to the orientational fluctuations of the individual carbon–deuterium bonds relative to the bilayer normal. The statistical amplitudes of the fluctuations correspond to the time-averaged second-order Legendre polynomials, \( (P_2(\cos \beta)) \), where \( \beta \) is the angle of the C–H bond axis to the bilayer normal. At any instant the segment orientation can be separated into a time-dependent part \( \beta(t) \) and a time-independent part \( \beta(t) \), a time-independent part. Here \( \beta(t) \) is the time-dependent angle between the ith carbon–deuterium bond (principal axis, \( P \)) and the bilayer normal (director axis, \( D \)). It corresponds to motions that are rapid on the NMR time scale, and lead to the averaging indicated by the angular brackets. On the other hand, \( \beta(t) \) is the time-independent angle between the bilayer normal \( \mathbf{b}_0 \) and the direction of the external magnetic field \( \mathbf{B}_0 \) (the laboratory axis, \( L \)), which characterizes the sample geometry. Motions of the entire membrane are typically too slow to contribute to motional averaging on the NMR time scale, e.g., as in the case of multimellar dispersions or large unilamellar vesicles. To relate the configurational properties of the acyl chains to the area per lipid, we must assume a distribution function, as further discussed below.

In solid-state NMR spectroscopy of membrane lipids, the structural properties are manifested by the RQCs, as given by [79]:

\[
\Delta R^{(i)}_Q = \frac{3}{2}R^Q_s S^{(i)}_o P_2(\cos \beta_Q).
\]
In the above formula, \( \Delta v^i_0 \) is the (absolute) quadrupolar splitting of the \( i \)th lipid segment, and \( \chi_0 = 167 \text{ kHz} \) is the static quadrupolar coupling constant. The dependence on the bilayer orientation is described by the second-order Legendre polynomial \( P_2(\cos \beta_{\text{PL}}) = (3\cos^2 \beta_{\text{PL}} - 1)/2 \) where \( \beta_{\text{PL}} \) is the angle of the bilayer normal (director axis) to the laboratory magnetic field (laboratory, \( L \)). For each lipid acyl segment, the order parameter \( S^i_{\text{CD}} \) is defined with respect to the bilayer director \( (D \) frame) as the time or ensemble average. Referring to Fig. 3, the segmental order parameter \( S^i_{\text{CD}} \) can be represented by the second-order Legendre polynomial \( P_2(\cos \beta_{\text{MD}}) \) or alternatively the second-rank Wigner rotation matrix element, leading to:

\[
S^i_{\text{CD}} = \frac{1}{2}3\cos^2 \beta_{\text{MD}}(t) - 1. \quad (2)
\]

The angular brackets indicate a time or ensemble average over those fluctuations of the segments that are faster than the (quadrupolar) interaction strength in frequency units, see Fig. 3(b). Based on geometrical considerations, the segmental order parameters are assumed to be negative for a polyethylene chain.

Fig. 4 illustrates the various transformations considered in the case of the mean-torque model. Denoting the segment index by \( i \) we have that the distribution of the angles \( \beta \equiv \beta^i_{\text{MD}} \) for the individual acyl segments is related to the statistical travel of the chain along the bilayer normal (director). We can then expand the matrix elements corresponding to \( \beta_{\text{MD}} \) into various coordinate frame transformations by using the closure property from group theory [94]. For \(^2\text{H}\) NMR the orientation of the C–H bond is considered as the principal axis system \( (P) \) for evaluating the segmental order parameter. The main external magnetic field \( B_0 \) corresponds to the laboratory frame \( (L) \). Designations for the Euler angles \( \Omega = (\alpha, \beta, \gamma) \) are: \( P \), principal axis system for \(^2\text{H}\) nucleus (2-axis parallel to C–H bond); \( L \), laboratory frame for methylene group motion (2-axis perpendicular to H–C–H plane); \( M \), molecular coordinate system; \( D \), director frame (2-axis is bilayer normal); and \( I \), laboratory system (2-axis along main external magnetic field \( B_0 \)). The closure property from group theory allows the overall rotation of the C–H bond to the laboratory frame to be expanded or collapsed in terms of various coordinate frames depending on the motional model.

3.2. Equilibrium statistics of membranes

In general, any angular-dependent property denoted by \( A(\beta) \) can be expressed in terms of an orientational distribution function \( f(\beta) \), which gives us the ensemble average,

\[
\langle A(\beta) \rangle = \int_{-1}^{1} A(\beta)f(\beta)d\cos \beta.
\]

As mentioned above, \( \beta \) is a generalized Euler angle (colatitude) (see Fig. 3) whose definition for a particular model will be introduced subsequently. The orientational distribution function \( f(\beta) \) can be expanded in a complete set of orthogonal polynomials, e.g., the Legendre polynomials \( P_j(\cos \beta) \), as

\[
f(\beta) = \sum_{j=-\infty}^{\infty} c_j P_j(\cos \beta), \quad (4)
\]
where \( c_j \) are coefficients. We recall that the Legendre polynomials \( P_j(\cos \beta) \) obey the orthogonality relation

\[
\int_{-1}^{1} P_j(x) P_k(x) \, dx = \frac{2 \delta_{jk}}{2j+1},
\]

where \( x \equiv \cos \beta \) and \( \delta_{jk} \) is the familiar Kronecker delta function.

Next, we left multiply the distribution function by another Legendre polynomial, and integrate over the full angular range. Using Eq. (5) allows us to solve for the expansion coefficients:

\[
c_j = \left( \frac{2j+1}{2} \right) \langle P_j(\cos \beta) \rangle.
\]

Here the values of \( \langle P_j(\cos \beta) \rangle \) correspond to the moments of the \( f(\beta) \) distribution, i.e., order parameters:

\[
\langle P_j(\cos \beta) \rangle = \int_{-1}^{1} P_j(\cos \beta) f(\beta) \, d\cos \beta
\]

By inserting Eq. (6) back into Eq. (4), we obtain our distribution function in terms of the Legendre polynomials, and their corresponding moments:

\[
f(\beta) = \sum_{j=0}^{\infty} \left( \frac{2j+1}{2} \right) \langle P_j(\cos \beta) \rangle \ P_j(\cos \beta).
\]

Evidently, knowledge of all the moments is required to completely specify the distribution function. Yet the order parameter measured by \(^1H\) NMR spectroscopy is only related to the second moment \( \langle P_2(\cos \beta) \rangle \) of the orientational distribution function \( f(\beta) \). As a general rule, \( f(\beta) \) is a function of both even- and odd-rank order parameters, including of particular interest the odd-rank term \( P_j(\cos \beta) \), which is related to the acyl chain segmental projection on the bilayer normal.

Therefore we must introduce a model for the segmental conformations to reconstruct \( \langle P_j(\cos \beta) \rangle \) from the given \( \langle P_2(\cos \beta) \rangle \) value. In other words, we need to assume a functional form for the orientational distribution function \( f(\beta) \).

3.3. Connecting dynamics to structure

Given the preceding framework, we are now equipped to address the configurational statistics of the various acyl segments of a flexible membrane lipid. For a methylene group, the relevant Euler angle \( \beta \equiv \beta_{CH2} \) is between the normal to the H–C–H plane of the intermediate frame (l) and the average molecular long axis, designated as the molecular frame (M). This approach lends itself to a liquid-crystalline picture for the individual segments of the flexible bilayer [1,95]. Alternatively, for each carbon segment (index i) we can consider the three carbon atoms from \( C_{i-1} \) to \( C_{i+1} \) in terms of a virtual bond of length 2.54 Å in the case of a methylene group [96]. The virtual bonds then correspond to a freely jointed chain, or other models used in polymer physics for chain molecules. Each definition has its own merits and limitations [34]. Here we utilize the treatment of three-carbon segments of the polymethylene chain [34].

The average segment projection onto the bilayer normal can then be written in terms of the first moment \( \langle D_i \rangle / D_M = \langle \cos \beta_i \rangle \) where \( D_i \) is the distance between carbon atoms \( C_{i-1} \) and \( C_{i+1} \) projected onto the bilayer normal, and \( D_M \) is the maximum projection of 2.54 Å. For a given acyl configuration, the sum of all of the three-carbon segment projections gives the total projected length \( \langle L_C \rangle \) or travel of the hydrocarbon chain. We can now address the problem of calculating the area \( \langle A \rangle \) per lipid. If we imagine a chain segment to be fluctuating in space, the degrees of freedom are limited by the volume within it moves. Calculating the average travel of a methylene chain segment near the lipid head group leads us to the average area \( \langle A_C \rangle \) per chain. For a symmetric (like chain) lipid, the area \( \langle A \rangle \) per lipid molecule is twice this value [34]. With the assumption that the average shape is a geometrical prism, the cross-sectional area for a statistical segment comprising two methylene groups is

\[
\langle A^{(i)} \rangle = \frac{2V_{CH2}}{D_i} = \frac{2V_{CH2}}{D_M} \left( \frac{1}{\cos \beta_i} \right).
\]

As discussed by Nagle and coworkers, \( V_{CH2} \) is the volume of a methylene group as obtained from density measurements [97–99]. The factor of two appears because the volume of the statistical segment represents two equivalent CH2 groups.

For calculating the average cross-sectional area (per chain) \( \langle A^{(i)} \rangle \), the value of the area factor \( q_i = 1/\cos \beta_i \) is clearly needed. Expanding to second order about \( x = 1 \) and truncating the Taylor series gives [34]:

\[
q_i = \frac{1}{\cos \beta_i} \approx 3 - 3(\cos \beta_i) + (\cos^2 \beta_i).
\]

Upturns (or back-folding) of a methylene segment are assumed to be negligible for the top part of the chain. Such an approximation is necessary, as \( 1/\cos \beta_i \) has a singularity at \( \beta_i = 0 \). The area calculation is less accurate for highly mobile methylene segments, and applies to methylene segments near the lipid head group (so-called plateau region of the order parameter profile). Supposing the index \( i \) the average cross-sectional area of a chain in terms of \( q \) is denoted by

\[
\langle A_C \rangle = \frac{2V_{CH2}}{D_M} q_i.
\]

Note that in the limit of a rotating all-trans chain with axial symmetry, \( \langle \cos^2 \beta_i \rangle = \langle \cos \beta_i \rangle = 1 \), giving \( q = 1 \) as expected. The limiting area per chain is \( 2V_{CH2}/D_M \) according to Eqs. (10) and (11). For a mixture of chains, the area factor \( q \) is the weighted sum, and the calculated value of \( \langle A \rangle \) is the number-weighted average over the components, according to the theory of moments [100].

Lastly, given the area per chain in Eq. (11) and the volume \( V_c \) of a hydrocarbon chain, we can calculate the volumetric thickness of an acyl chain (for an individual monolayer):

\[
D_c = \frac{n_c V_{CH2}}{2q},
\]
of a continuous orientational potential is equivalent to an average torque (potential of mean force) for the individual methylene segments; hence the appellation mean-torque model. The approach is akin to a liquid crystal view of the membrane, whereby the various segments of the flexible lipid molecule are subject to a orienting potential [95]. Our strategy is to reconstruct the first moment of the segmental or molecular orientational distribution \(\langle P_1(\cos \beta_j)\rangle\) from the second moment \(\langle P_2(\cos \beta_j)\rangle\), which allows us to calculate the average membrane structure [50] in terms of the orientational distribution function using Eq. (4) [34]. An advantage is that specific orientations of the methylene segments are not assumed for calculating structural parameters.

We begin with the distribution function corresponding to a given orientational potential [34]. The orientational distribution for each methylene segment is written in terms of the Boltzmann factor

\[
f(\beta) = \frac{1}{Z} \exp\left( - \frac{U(\cos \beta)}{k_B T} \right),
\]

in which the partition function is

\[
Z = \int_{-1}^{1} \exp\left( - \frac{U(\cos \beta)}{k_B T} \right) \, d \cos \beta.
\]

In the above formula, \(U(\cos \beta)\) is the orientational potential for an individual carbon segment, and \(x \equiv \cos \beta\). To simplify the notation, for the mean-torque model, \(\beta = \beta_{\text{mol}}\) where the suffix and superscripts (i) representing the segment index are suppressed.

For statistical treatment of the possible orientations of the methylene segments, the mean-torque model assumes the orientational order is described by a potential of mean force. In a first-order approximation, the potential is given by

\[
U(\cos \beta) = \sum_i U_i P_i(\cos \beta) \approx U_1 \cos \beta,
\]

where \(U_i\) is the first-order mean-torque parameter. Knowing these parameters for each chain segment gives us information about the stress profile of the bilayer. The first and second moments are obtained from integrating Eq. (7) with use of Eqs. (14) and (15) for the distribution function:

\[
\langle P_1(\cos \beta)\rangle = \frac{1}{Z} \int_{-1}^{1} P_1(\cos \beta) \exp\left( - \frac{U(\cos \beta)}{k_B T} \right) \, d \cos \beta.
\]

Evaluation of the integral in Eq. (16) in closed form then yields the desired analytical results,

\[
\langle P_1(\cos \beta)\rangle = \frac{1}{Z} \int_{-1}^{1} \frac{U_1}{U_1 + k_B T} \exp\left( - \frac{U(\cos \beta)}{k_B T} \right) \, d \cos \beta = \frac{1}{Z} \int_{-1}^{1} \frac{U_1}{U_1 + k_B T} \, d \cos \beta = \frac{1}{Z} \int_{-1}^{1} \frac{U_1}{U_1 + k_B T} \, d \cos \beta = \frac{1}{Z} \int_{-1}^{1} \frac{U_1}{U_1 + k_B T} \, d \cos \beta.
\]

and

\[
\langle P_2(\cos \beta)\rangle = 1 + \frac{3}{2} \left( \frac{k_B T}{U_1} \right)^2 + \frac{3}{2} \frac{k_B T}{U_1} \coth\left( - \frac{U_1}{k_B T} \right).
\]

In Eq. (18), the second moment of the distribution is measured directly from solid-state \(^2\)H NMR experiments by the segmental order parameter \(P_2(\cos \beta_{\text{mol}}) = S_{\text{CD}}^2\), where the segment index (i) has now been reintroduced. Taking account of the factor of \(P_2(\cos \beta_{\text{mol}}) = P_2(\cos 90^\circ) = -1/2\) for the methylene segments gives \(1 - 4S_{\text{CD}}^2/3 \equiv \cos^2 \beta_i\). Eq. (18) can be then solved numerically to deduce the first-order mean-torque parameter \(U_1\) for any segment in the chain. If \(U_1\) is known, the average projection, Eq. (17), can be found, and the average acyl length projection \(\langle L_\text{C} \rangle\) calculated. Alternatively, an analytical solution for \(\langle \cos \beta_i \rangle\) is obtained by using the approximation \(\coth( - U_1/k_B T) \approx 1\), which for individual segments leads to:

\[
\langle P_1(\cos \beta_i)\rangle = \langle \cos \beta_i \rangle = \frac{\langle D_i \rangle}{D_M} \approx \frac{1}{2} \left( 1 + \sqrt{-8S_{\text{CD}}^2 - 1} \right). \quad (19)
\]

It is assumed there are no upturns of the segment for a very strong orienting potential. This relation is valid only for order parameters in the range of \(-1/8 < S_{\text{CD}} < -1/2\), because their values are assumed negative.

Using Eq. (19) and knowing the order parameters along the acyl chain, we calculate the average projected acyl length as the sum of the average segment projections:

\[
\langle L_\text{C} \rangle = \frac{n_s}{2} \sum_{i=1}^{n_s} \left( 1 + \sqrt{-8S_{\text{CD}}^2 - 1} \right) + \frac{n_m}{3} \sum_{m=1}^{n_m} \left( - \frac{24S_{\text{CD}}^2}{3} \right) - 1.
\]

For highly mobile lipids, the absolute order parameter for the terminal methyl groups is very low; so Eq. (18) should then be solved numerically. The methyl segment requires special treatment, as the carbon–deuterium bond is oriented differently than for the methylene segments. The three-fold rotational symmetry projects the residual quadrupolar coupling along the carbon–carbon bond, leading to \(S_{\text{CD}}^2 P_2(\cos 109.5^\circ) = S_{\text{CD}}^2 + S_{\text{CD}}^6 = -3S_{\text{CD}}^2\). The result is

\[
\langle L_\text{C} \rangle = \frac{n_s}{2} \sum_{i=1}^{n_s} \left( 1 + \sqrt{-8S_{\text{CD}}^2 - 1} \right) + \frac{n_m}{3} \sum_{m=1}^{n_m} \left( - \frac{24S_{\text{CD}}^2}{3} \right) - 1.
\]

By combining Eqs. (2) and (19) with Eqs. (10) and (11), we then obtain the mean-torque expression for the average area per chain

\[
\langle A_c \rangle = \frac{2V_{\text{CH}}}{D_M} \left( \frac{11}{6} \left( \frac{1}{2} \sqrt{3( -8S_{\text{CD}}^2 - 1) - 4} \right) - \frac{4}{3} \right).
\]

where the area factor \(q\) is contained in the parentheses, and the order parameters are negative. This method of calculating the chain cross-sectional area by using the mean-torque model has been shown to be in agreement with other experimental methods [102]. Last, using Eq. (12) together with Eq. (22), the volumetric thickness is found to be:

\[
D_{L} = \frac{n_s}{2} \sum_{i=1}^{n_s} \left( 1 + \sqrt{-8S_{\text{CD}}^2 - 1} \right) + \frac{n_m}{3} \sum_{m=1}^{n_m} \left( - \frac{24S_{\text{CD}}^2}{3} \right) - 1.
\]

One should recall that the maximum order parameter (plateau region) is used for this calculation.

4. Thermodynamics of membrane deformation and dehydration

We next turn our attention to how the structural parameters obtained from solid-state \(^2\)H NMR spectroscopy can help us to understand the forces governing membrane organization, remodeling, and deformation. In exploring the molecular interactions in phospholipid bilayer membranes, the osmotic stress method [103], surface forces apparatus [104], and micropipette aspiration method [105,106] have figured prominently. Each of these methods essentially involves consideration of the membrane as a macroscopic material. Because solid-state \(^2\)H NMR spectroscopy yields atomistic knowledge for liquid-crystalline phospholipids, Fig. 2(a), it has the potential to transform our comprehension of how the material properties begin to emerge from intermolecular forces [107–109].
multilamellar lipid phase with the osmotic stress due to application of an external force. The free energy of the system is reduced by transferring water from the lipid membrane phase to a stressing polymer solution, or by gravimetric removal of water. In either case water is removed, thereby maximizing entropy. Deformation of the membrane lipid phase occurs by reduction of the water volume at the aqueous interface, reducing the area per lipid with a concomitant increase in volumetric bilayer thickness. The lipid phase is separated from the polymer solution by either a semipermeable membrane, or a virtual (imaginary) dividing surface that bisects the system into thermodynamically distinct lipid and osmolyte phases [103,112,113]. Due to an unfavorable loss of entropy, the stressing polymer is not admitted to the multilamellar lipid phase. Hence the stressing polymer solution does irreversible work on the lipid phase by removing water. For the osmolyte phase, the additional pressure increases the chemical potential of the water, which then becomes equal to the solvent chemical potential in the lipid phase. Deformation of the lipid phase occurs due to changing the water volume, with temperature and pressure held constant.

4.1. Free energy of the lipid phase

For the lipid phase, we are interested in how the work content (Helmholtz free energy) changes with the water volume under the constant osmotic pressure. The total differential of the Helmholtz free energy ($F$) is given by [114]

$$dF = -PdV - SdT + \sum_k \mu_k dn_k,$$

(24)

where $F$ is an extensive thermodynamic state variable. In the above formula $S$ is the entropy, $T$ is the temperature, and the chemical potentials are defined by $\mu_k = (\partial F/\partial n_k)_{V,T,n_{\text{other}}}$, where $n_k$ is the moles of the $k$th component, holding the natural variables ($T$ and $V$) constant. The first two terms on the right correspond to a closed system, where $\partial F/\partial V|_{T,n_k} = -P$ and $\partial F/\partial T|_{V,n_k} = -S$. The summation gives the change due to mass transfer of $dn_k$ moles of the $k$th component with chemical potential $\mu_k$ for an open system.

Essentially, the volume of the lipid phase can change in two ways—that is to say, either by compression due to a change in pressure at constant number of waters ($N_w$), or by changing $N_w$ at constant pressure. In the osmotic stress method, we assume the lipid phase is incompressible, i.e. the density is approximately constant, and hence the partial lipid and water volumes remain $\approx$ unchanged. Only a mass transfer of water is involved with the osmotic pressure held constant, and hence the changes in either the Gibbs or Helmholtz free energies holding their natural variables ($T$ and $P$, or $T$ and $V$, respectively) constant are the same. They both depend on the chemical potential $\mu_{nw}$ of the aqueous solvent, together with the moles of water transferred across the thermodynamic dividing surface.

For a given composition, if we hold the volume of the lipid phase and the temperature constant, then the total differential of the free energy, Eq. (24), is simplified accordingly. Identifying $F$ as the Helmholtz free energy per lipid molecule, and $n_{nw}$ as the moles of associated waters per lipid, the total differential becomes: $dF = \mu_{nw}dn_{nw}$. Conservation of energy (first law of thermodynamics) thus implies that the reversible work $\mu_{nw}dn_{nw}$ done on the lipid phase is equal but opposite to the work done by the osmolyte phase. Substituting $\mu_{nw} = \Pi_{nw}$ for the osmolyte phase leads to the result that:

$$dF = -\Pi_{nw}dV_{nw}. $$

(25)

Here, we have formulated the water volume per lipid as: $V_{nw} = V_{nw}n_{nw} = v_{nw}N_w$, where $v_{nw} = V_{nw}/N_w$ is the (partial) molecular volume of water, $N_w$ is the Avogadro constant, and $N_w$ is the number of waters per lipid molecule. Typically, it is assumed that the partial molar volume $V_{nw}$ is approximately equal to the water molar volume $V_{nw}$ and that it remains $\approx$ constant. The effect of osmotic pressure on the (total) volume of the lipid phase is analogous to the reduction in volume of a gas that occurs by application of a constant external pressure. Because the volumetric reduction of the lipid phase occurs in the same direction as the external osmotic pressure, the reversible work is positive.

Eq. (25) states that the reversible work of deforming the lipid phase—due to changing the bilayer separation plus any structural deformation of the bilayer—corresponds to the directly measured removal of water from the lipid phase. The work is positive because $dV_{nw}$ is negative for movement of water from the lipid phase to the osmolyte phase. The removal of water can be accomplished either osmotically or gravimetrically. By introducing the area per lipid ($A$) and the water thickness $D_{nw}/2$ as the lattice variables [34] (see Fig. 2), the total differential can be written as

$$dF = \left(\frac{\partial F}{\partial (A)}\right)_{D_{nw}/2} d(A) + \left(\frac{\partial F}{\partial D_{nw}/2}\right)_{A} dD_{nw}/2.$$

(26)

The above formula states that for the lipid phase, the free energy depends only on the area per lipid ($A$) and $D_{nw}/2$, which represents the interlamellar water spacing. We can then write the water volume in terms of the area per lipid molecule and the water spacing for a geometrical prism (see Fig. 2), giving $V_{nw} = \langle A \rangle D_{nw}/2$ as the result. Upon differentiation and combination with Eq. (25), we obtain

$$dF = -\Pi D_{nw}/2 d\langle A \rangle - \Pi(A) dD_{nw}/2.$$

(27)

Here we recall that the osmotic pressure $\Pi \approx$ constant due to a large excess of the stressing polymer solution, or due to gravimetric removal of water.

From the above total differential, we then obtain the following thermodynamic relations [46,115]:

$$\left(\frac{\partial F}{\partial (A)}\right)_{D_{nw}/2} = -\Pi D_{nw}/2 = \tau,$$

(28)

and

$$\left(\frac{\partial F}{\partial D_{nw}/2}\right)_{A} = -\Pi(A) = -F_K,$$

(29)

where $F_K$ is the repulsive force acting between the various bilayers. The first equation, Eq. (28), tells us that the change in Helmholtz free energy $F$ with respect to the interfacial area ($A$) per lipid corresponds to the lateral tension $\tau$ acting on a lipid molecule in a bilayer. The second equation, Eq. (29), states that the free energy per lipid due to a change in the bilayer separation gives the force ($F_K$) acting perpendicularly to the bilayer surface. Reduction of the area per lipid ($d\langle A \rangle$ negative) as the bilayer separation decreases ($dD_{nw}/2$ negative) is unfavorable ($dF$ positive), meaning that work is done by the stressing polymer solution on the lipid phase. Our next question is: how much of this work goes into bilayer separation, and how much goes into bilayer deformation?

4.2. Separation work versus area deformation

The above results allow us to divide the effect of osmotic pressure into the influences of separation forces, and those of lateral tension (which is zero for a flaccid bilayer in equilibrium with excess water). We have used the definition of the lateral tension [116] to obtain $\tau = -\Pi D_{nw}/2$ in Eq. (28). Clearly, the lateral tension for a lipid bilayer is a function of the area per lipid molecule. Because the tension $\tau$ corresponds to a negative pressure, condensing the bilayer costs
work, thereby giving an increase in free energy. If we define the repulsive
pressure as $P_k = f_k/A$, then $\Pi = \Pi_k$ in accord with Eq. (29). The
osmotic pressure $\Pi$ is a positive quantity due to a positive repulsive
force in Eq. (29), which implies there is a tendency for the multilamellar
lipids to expand indefinitely. At some point, however, the swelling from
the repulsion is counterbalanced by the long-range attractive force [34],
due to van der Waals interactions.

We can then calculate the fraction of work that goes into reducing the
bilayer separation versus the area deformation. The ratio of separation
work to area work $x$ is defined as [46]:

$$
x = \frac{\text{separation work}}{\text{area work}} = \left( \frac{\partial F}{\partial D_{W/2} / A} \right) \frac{dD_{W/2}}{d(A)}.
$$

(30)

Following Rand and Parsegian et al. [46], Eqs. (28) and (29) allow us to
simplify Eq. (30) yielding:

$$
x = \frac{-\Pi(A) D_{W/2}}{-\Pi D_{W/2} / d(A) d(A)} = \frac{\ln D_{W/2}}{\ln A}.
$$

(31)

The result above corresponds to the fraction of area work $\theta$ by the
relation: $\theta = x / (1 + x)$. The fraction of area work allows us to calculate
the percentage of energy that goes into deforming the lipid membrane,
as opposed to reducing the interlamellar distance. One should take note that
Eqs. (28) and (29) do not contain the fraction of area work, because the
partial derivatives involve separate contributions from the lattice
variables ($A$) and $D_{W/2}$.

To obtain the area compressibility of the surface film, we first recall
that the lateral tension $\tau$ is defined in terms of the Helmholtz free energy
as [116]:

$$
\left( \frac{\partial F}{\partial A} \right)_{T, V, n_L} = \tau,
$$

(32)

(or alternatively $(\partial G / \partial A)_{T, P, n_L}$ in terms of the Gibbs free energy),
where all symbols have their usual meanings; see also Eq. (28). In
the absence of osmotic pressure, the lipid bilayer is flaccid and not
under tension; and hence the area per lipid is the equilibrium value
[34]. Knowing the water associated with the lipid head group allows us
to recast the expression for the lateral tension $\tau$ in Eq. (28). Substituting
the relation $D_{W/2} = \nu_w \Pi / (A)$ into Eq. (28) gives us the result that:

$$
\tau = -\left( \frac{\nu_w \Pi}{A} \right) \Pi,
$$

(33)

where $\nu_w$ is the partial molar volume of water at the bilayer aqueous
interface, and $\Pi$ is the osmotic pressure. For a lipid surface film, the
area compressibility is defined as [117]

$$
C_A \equiv \frac{1}{K_A} \equiv \left( \frac{\partial \Pi}{\partial \Pi} \right)_{T, V, n_L} = \left( -\frac{1}{\nu_w} \ln \Pi / (A) \right) \left( \frac{\partial (A)}{\partial \Pi} \right)_{T, V, n_L},
$$

(34)

in which $K_A$ is the area compressibility modulus. Because the osmotic
stress is applied equally to both sides of the interface, this relation
holds also for bilayers. Upon integration over the applied pressure
range, we can then rewrite our expression for the cross-sectional area
in terms of osmotic pressure as

$$
\langle A \rangle = \left( -\left( \frac{\nu_w \Pi}{K_A} \right) \Pi \right) + \langle A \rangle_0.
$$

(35)

where $\langle A \rangle_0$ represents the average cross-sectional area per lipid [34]
at zero osmotic pressure (full hydration) and constant temperature $T$. It is
typically assumed that $\nu_w = \nu_w^* i.e.$ the partial molar volume is
approximately equal to the molar volume of pure water (vide infra).

4.3. Osmotic pressure and nonideality of solvent water

Especially the solvent water is expected to behave nonideally in
both the multilamellar lipid phase and the stressing polymer solution.
According to classical thermodynamics, deviations from ideality are accounted for in terms of an activity coefficient. For the
two phases in thermodynamic equilibrium, the common reference
state is pure water, with $\mu_w$ as its chemical potential. In the case of
a binary solution, with water as the solvent, the chemical potential
depends on its activity $\alpha_w$ by $\mu_w = \alpha_w \gamma_w$ as observed by Raoult’s
law. However, direct measurement of the solvent vapor pressure
$P_w$ for multilamellar lipids is fraught with difficulty [33,69,113].
Multilamellar lipid dispersions under osmotic stress require very ac-
curate vapor pressure measurements [46], giving a paradox [48, 118–120] that has bedeviled previous investigators. Using vapor
pressure osmometry, it is challenging to measure the water activity
in both the osmolyte phase and the lamellar lipid phase over the
full range of interest.

The osmotic pressure $\Pi$ can be treated for a nonideal solution by
introducing a virial expansion for the solvent chemical potential in
terms of the solute concentration. Alternatively, a semiempirical
equation of state can be employed, as introduced by Parsegian and
coworkers [110]. Here we use experimentally measured osmotic
pressures rather than theoretical values. The water activity is measured
experimentally, which is related to the polymer solute activity by the
Gibbs–Duhem equation. We are thus able to effectively bypass the
nonideality of the stressing polymer solution [113]. Introduction of
an osmotic coefficient $\varphi$ allows us to simplify the treatment of the
nonideality of water in both the multilamellar lipid dispersion and
the osmolyte solution [3]. By equating the solvent chemical potential
$\mu$ of the two phases in equilibrium, we can connect the nonideality
of the aqueous solvent of the multilamellar lipid phase to the bilayer
forces. We are thus able to obtain knowledge of the repulsive inter-
lamellar forces and the forces acting between the lipids molecules
in the bilayer.

The following equation of state has been introduced [3] to describe
how the osmotic pressure acts upon multilamellar lipid membranes
in terms of the number of water molecules per lipid

$$
\Pi = \phi \left( \frac{RT}{\nu_w} \right) \frac{1}{N_w} = \phi \left( \frac{k_B \Pi}{\nu_w} \right) \frac{1}{N_w},
$$

(36)

in which $\nu_w = \Pi / N_A$ and $R = k_B N_A$ is the gas constant. In the above
formula $\phi$ is the osmotic coefficient [121] which is defined in terms of
the solvent (water) mole fraction $X_w$ by:

$$
\phi = \frac{(\mu_w^* - \mu_w^*)}{RT} \ln X_w.
$$

(37)

Here $\mu_w^*$ and $\mu_w$ are the chemical potentials of pure water and the
aqueous solvent in the solution, and $X_w$ is the solvent (water) mole fraction
for either the stressing polymer solution or the multilamellar lipid
dispersion. In Eq. (36), the osmotic coefficient $\phi$ is a measure of the
nonideality of the aqueous solvent, where $\phi = 1$ represents the limit
for osmolytes with purely colligative behavior. The above equation of
state, Eq. (36), has been tested experimentally [3] and the applied
osmotic pressure $\Pi$ is found to scale with $1/N_w - 1/n_w$ for the lipid
systems studied.
It can also be shown that the osmotic coefficient is the ratio of the separation work to thermal energy via Eqs. (29) and (36):

$$\phi = \frac{\Gamma v w N_w}{k_B T} = \frac{P_w v_w}{k_B T} = \frac{F_{Dw/2}}{k_B T}.$$  \hspace{1cm} (38)

For completely disassociated molecules, the thermal energy results from their kinetic motion. Attractive forces between the solute molecules (either in the case of polymer solutions or multilamellar lipids) and the aqueous solvent reduce the osmotic coefficient. Conversely, nonideal repulsive forces between the repelling bilayers give a larger osmotic coefficient. We are now in a position to ask how the thermodynamic formalism can be connected to the changes in bilayer observables studied by solid-state $^2$H NMR spectroscopy.

5. Remodeling and elasticity of membranes viewed by solid-state NMR spectroscopy

Let us now return to the question of how the atomistic results of solid-state $^2$H NMR are connected with membrane structure and the associated intermolecular forces. In this section, we explain how $^2$H NMR spectroscopy allows one to investigate the possibility of membrane deformation due to osmotic stress [47,49,69,122]. Our aim is to address how changes in thermodynamic state variables correspond to restructuring or remodeling of biomembranes, and how these effects can be quantified. We then turn to how knowledge of such state variables—as they emerge from atomistic level interactions—leads us to an enhanced comprehension of lipid–protein interactions in relation to the actions of membrane proteins, such as ion channels or G-protein-coupled receptors (GPCRs).

5.1. Correspondence of dehydration and osmotic stress of membrane lipids

Fig. 5 shows the striking changes in the solid-state $^2$H NMR spectra and the corresponding C–$^2$H bond order parameter profiles observed for DMPC-<em>ds</em> samples [3] due to applying osmotic stress. Deconvoluted (de-Paked) $^2$H NMR spectra are shown at the left of Fig. 5(a) for DMPC-<em>ds</em> samples in the liquid-crystalline state, where the water–to-lipid mass ratio is varied gravimetrically. Removal of water begins to stress the membrane noticeably, as revealed by changes in the observed quadrupolar splittings. A continuous increase is evident from 30 wt.% H_2O (<em>N_w</em> = 18) until 3.1 wt.% H_2O (<em>N_w</em> = 1.5). Moreover, in Fig. 5(b) at the left we see that similarly striking changes are evident in the $^2$H NMR spectra of DMPC-<em>ds</em> upon exposure to stressing polymer solutions. Osmotic stress is introduced by controlling the water activity through exposure to polymer solutions containing polyethylene glycol of molar mass M_0 = 1500 (PEG 1500). For the de-Paked $^2$H NMR spectra corresponding to DMPC-<em>ds</em> samples with different PEG 1500 mass ratios, there is a striking increase of the RQCs as the concentration of osmolyte increases, or equivalently as the osmotic pressure increases from 0% PEG 1500 (excess hydration) to 87.6% PEG 1500 (<em>N_w</em> ≈ 1.3). For either gravimetric dehydration or osmolyte addition, the spectral changes are due to varying the water activity of the samples.

Next, the corresponding order parameter profiles for DMPC-<em>ds</em> obtained under conditions of dehydration or osmotic stress are shown at the right in Fig. 5(a) and (b). The order parameters decrease from the upper acyl chain (C2–C4 plateau position) to the terminus near the bilayer center (C14 carbon) [79]. In the liquid-crystalline state, the lipids are effectively tethered to the aqueous interface through their polar head groups. Among the various rotational isomeric states (e.g. trans, gauche<sup>+</sup>, gauche<sup>−</sup>), correlations of the lipid chains favor their extension (travel) away from the aqueous interface. Approaching the bilayer center, there is a progressive drop in segmental order due to the effect of the chain terminations, see Fig. 3(a). The chain ends are statistically distributed, and require greater disorder of the surrounding acyl groups to maintain the hydrocarbon density $\approx$ constant [123]. Formulated as a potential of mean force, the orientational potential energy is greatest for the top part of the chains, closest to the aqueous interface. On the other hand, the hydrocarbon interior experiences the weakest ordering potential of the membrane, resembling a simple liquid paraffin [124]

Our results demonstrate both theoretically and experimentally that significant bilayer deformation occurs with osmotic pressures of 10–100 atm (1–10 MPa), values within the biological range [3]. Moreover, solid-state $^2$H NMR spectroscopy gives us a basis for investigating how the osmotic pressure results can be compared to bilayer deformation induced by hydrostatic pressure [26]. Effectively we use solid-state $^2$H NMR spectroscopy as a secondary osmometer to establish the equivalence of osmotic pressure and hydrostatic pressure. Referring to Fig. 6, we see that osmotic pressure [3] has a far greater effect on membrane deformation than does hydrostatic pressure [26,125,126]. Previously we have proposed that the comparatively small deformations induced by large hydrostatic pressures (1000 atm) are due to squeezing water from the interlamellar space. This process is far less efficient than direct removal of water by dehydration or osmotic stress, and hence the deformation is correspondingly smaller [103].

5.2. Solid-state NMR spectroscopy of membranes under stress

Biological membranes and lipid bilayers in the liquid-crystalline state are known to be laterally compressible [58] materials. Removal of water from the lipid head groups increases the acyl chain ordering, thereby reducing the cross-sectional area per (lipid) hydrocarbon chain. Conversely, increasing temperature causes disordering to occur with a concomitant area expansion [126]. Previous studies using small-angle X-ray scattering (SAXS) in conjunction with the Luzzati method have concluded that lipid bilayers deform appreciably with
osmotic pressures in the range of 0.5–3.0 MPa (5–27 atm) [46,47,103, 112,127]. However, others have concluded from the analysis of electron density profiles of lipid bilayers that essentially negligible deformation occurs [63]. As pointed out by Mallikarjunaiah et al. [3], an alternative approach is needed to decide among these proposals. In this regard, solid-state $^2$H NMR spectroscopy is unparalleled in the level of detailed structural information that it can deliver in the case of phospholipid liquid crystals [79,128].

Fig. 7 demonstrates the remarkable changes observed in the cross-sectional area per lipid for the DMPC membrane system when the osmotic pressure is varied [129]. The mean-torque model allows changes in the average cross-sectional area per lipid ($A'$), bilayer thickness $D_b = 2D_C + 2D_p$, and water spacing $D_w$ to be established [3]. Reduction of interlamellar water from $N_w = 20$ to $N_w = 1.5$ leads to a change of the water spacing from $D_w = 20.1$ Å to 1.8 Å, a substantial range. Part (a) of Fig. 7 shows that boosting the osmotic pressure up to 200 atm (20 MPa) gives a substantial reduction of the area per lipid, with a gain of the volumetric bilayer thickness. According to $^2$H NMR spectroscopy, the cross-sectional area per lipid shrinks from 60.2 Å$^2$ at full hydration ($N_w \approx 20$) to 50.2 Å$^2$ ($N_w \approx 1.5$) for both gravi-

cational and osmolyte samples at 30 °C. Overall, the lipid cross-sectional area deformation is $\Delta A = 10$ Å$^2$ and represents a 17% area contraction.

Correspondingly, the volumetric bilayer thickness $D_b$ expands from 43.6 Å ($N_w \approx 20$) to 48.8 Å ($N_w \approx 1.5$). The resulting bilayer thickness deformation is $\Delta D_b = 5.2$ Å giving a 20% swelling of the hydrocarbon thickness ($2D_C$). Such large bilayer deformations have significant implications for hydrophobic matching to proteins. It should also be noted that these osmotic pressures far exceed those than could be practically achieved by applying hydrostatic pressure.

Last of all, in part (b) of Fig. 7 the logarithm of the average cross-

sectional area per lipid is plotted as a function of osmotic pressure. The elastic area compressibility modulus ($K^e$) is calculated as $142 \pm 30$ mJ m$^{-2}$ from the initial slope of the plot of average cross-sectional area against osmotic pressure in accord with Eq. (34). The measured value of $K^e$ is in close agreement with the values reported independently by Koenig et al. [48] ($136 \pm 15$ mJ m$^{-2}$) and by Petrache et al. [47] ($108 \pm 35$ mJ m$^{-2}$), using SAXS and/or solid-state $^2$H NMR measurements. However, our measurements cover a much greater range of osmotic pressure [3], and enable the theory in the preceding sections of this article to be more accurately tested. By comparing the material properties studied with $^2$H NMR to the results of micromechanical studies and SANS measurements, we are able to investigate how the mesoscopic (Hookian) elastic behavior emerges from atomistic interactions due to bilayer interactions with water.

6. Membrane deformation in cellular function

Caught in the debate of whether lipids or proteins are more important [4], one can easily overlook the ubiquitous role of water. Indeed, biological membranes interact strongly with water—that much is at least clear [3]. It is quite improvident to focus on membrane proteins at the expense of the other components, e.g. the lipids [3,130], water [112,113,131], and carbohydrates [30]. Absent water, biological function—indeed life itself—ceases as in the case of anhydrobiosis. Bulk water has also been found to play an important role in lipid-mediated GPCR activation [2] and other membrane protein functions [113,131–133]. Evidently the bilayer deformation due to the lipids alone can influence how osmotic stress affects membrane protein activity. For fluid membranes, the thickness compression is equivalent...
to changing the bilayer thickness by roughly four methylene carbon segments—large enough for changes in protein activity due to hydrophobic matching [20,134]. Altering the lipid hydrophobic thickness by 4 Å incurs an energy penalty of about 0.3 RT per mol lipid, assuming a value of 1.5 RT for the free energy of transfer of methylene groups from hydrocarbon to water [135]. Because the equilibrium constant \( K = \exp(-\Delta G/RT) \), a standard free energy difference of just a few RT is sufficient to \( \approx \) completely shift a protein conformational equilibrium from initial to final states [136]. Bilayer deformation can readily affect membrane proteins, such as ion channels [137] and G-protein-coupled receptors (GPCRs) [4]. The moduli of compressibility and bending rigidity obtained from the atomistic solid-state \( ^1H \) NMR studies can be also compared with micro- or nanomechanics based methods [105, 138,139] like atomic force microscopy.

Knowledge of membrane elasticity at the atomic level as revealed by NMR is necessary to treat the energies involved in protein conformational changes. Proper accounting for lipid forces in biological mechanisms rests upon the quantitative analysis of protein–lipid interactions in membranes. The driving force for inserting proteins into membranes is quantified by the well-established hydrophobicity scales for amino acids [72,140]. The question is then: once inserted into the membrane, how do proteins carry out the work of conformational changes, and interact with the membrane lipid bilayer? In this context, solid-state NMR spectroscopy continues to pay a major role with regard to the interaction with the membrane lipid bilayer? In this context, solid-state NMR spectroscopy will allow us to move beyond immobile structures toward a dynamic vision of biomembranes founded on magnetic resonance spectroscopy.

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References
