The Role of Membrane-Mediated Interactions in the Assembly and Architecture of Chemoreceptor Lattices

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Abstract

In vivo fluorescence microscopy and electron cryo-tomography have revealed that chemoreceptors self-assemble into extended honeycomb lattices of chemoreceptor trimers with a well-defined relative orientation of trimers. The signaling response of the observed chemoreceptor lattices is remarkable for its extreme sensitivity, which relies crucially on cooperative interactions among chemoreceptor trimers. In common with other membrane proteins, chemoreceptor trimers are expected to deform the surrounding lipid bilayer, inducing membrane-mediated anisotropic interactions between neighboring trimers. Here we introduce a biophysical model of bilayer-chemoreceptor interactions, which allows us to quantify the role of membrane-mediated interactions in the assembly and architecture of chemoreceptor lattices. We find that, even in the absence of direct protein-protein interactions, membrane-mediated interactions can yield assembly of chemoreceptor lattices at very dilute trimer concentrations. The model correctly predicts the observed honeycomb architecture of chemoreceptor lattices as well as the observed relative orientation of chemoreceptor trimers, suggests a series of “gateway” states for chemoreceptor lattice assembly, and provides a simple mechanism for the localization of large chemoreceptor lattices to the cell poles. Our model of bilayer-chemoreceptor interactions also helps to explain the observed dependence of chemotactic signaling on lipid bilayer properties. Finally, we consider the possibility that membrane-mediated interactions might contribute to cooperativity among neighboring chemoreceptor trimers.

Introduction

The chemotaxis signal transduction pathway [1] allows bacteria to respond to minute relative changes in chemical concentration over several orders of magnitude in ambient chemical concentration [2], and ranks among the most studied signaling pathways in biology. The extreme sensitivity of the chemotaxis system results from amplification of external signals coupled with adaptation to persistent stimuli [3–7]. Fluorescence resonance energy transfer (FRET) experiments [8,9] have revealed that a crucial step in signal amplification occurs at the level of chemoreceptors: chemoreceptor signal in cooperative teams [8–10]. Indeed, Ising [11–13] and Monod-Wyman-Changeux [9,14–16] models of coupled teams of chemoreceptors both achieve quantitative agreement with the FRET data. The fundamental assumption underlying these models of signaling teams is that chemoreceptors do not respond independently to changes in the external ligand concentration, but rather each receptor influences the collective state of a team of neighboring receptors. Thus, the observed functional characteristics of chemotactic signaling rely on cooperative local interactions among chemoreceptors, and suggest a well-defined spatial organization of chemoreceptors.

From a structural perspective, chemoreceptors are homodimers, which interact strongly to form trimers-of-dimers [17,18]. Recent breakthroughs in in vivo electron cryo-tomography have revealed [19–25] that chemoreceptor trimers form two-dimensional honeycomb lattices in which each trimer has three nearest-neighbors arranged in a face-on orientation. The honeycomb lattice architecture and characteristic lattice constant of 12 nm appear to be universally conserved among bacterial species [22]. Functional complexes require chemoreceptors plus the linker/kinase CheA and the linker protein CheW [19,20], which may mediate cooperative interactions among neighboring trimers [9,10]. Fluorescence experiments have indicated that chemoreceptor lattices can exhibit variable stoichiometries of chemoreceptors, CheA, and CheW [9,26,27], while clustering of chemoreceptors requires neither CheA nor CheW [27,28]. The size of chemoreceptor clusters can range from tens to thousands of receptors, with large chemoreceptor clusters observed predominantly at the cell poles but smaller clusters also found in the midcell regions [21–25,27–30]. Superresolution light microscopy of chemoreceptor lattices has suggested a stochastic model for cluster assembly [28,30] in which self-assembly of chemoreceptor lattices proceeds by nucleation and growth. Such stochastic self-
assembly of chemoreceptor lattices relies on the existence of attractive interactions between chemoreceptor trimers, but in principle does not require direct cytoskeletal involvement or active transport of chemoreceptors.

What are the molecular mechanisms yielding attraction between chemoreceptor trimers and, hence, self-assembly of chemoreceptor lattices? Chemoreceptors are transmembrane proteins localized in the cytoplasmic membrane of bacteria. In general, membrane proteins deform the surrounding lipid bilayer [31,32], which can lead to membrane-mediated interactions between proteins [33,34]. Thus, while chemoreceptors can be coupled by protein-protein interactions [9,10], they may also interact via the cytoplasmic membrane. Here we develop a biophysical model of membrane-mediated interactions between chemoreceptor trimers which shows that membrane-mediated interactions can yield stochastic cluster assembly even at very dilute trimer concentrations. The model correctly predicts the observed face-on orientation of chemoreceptor trimers at small trimer separations [19,20] and suggests a series of “gateway” states for chemoreceptor lattice assembly. We find that the three-fold-symmetric directionality of membrane-mediated interactions between trimers can stabilize the observed honeycomb architecture of chemoreceptor lattices [19,20] even at suboptimal stoichiometries of chemoreceptors, CheA, and CheW [9,26,27]. The model also suggests a simple mechanism by which bilayer-chemoreceptor interactions can localize large chemoreceptor clusters to the cell poles even in the absence of interactions with CheA and CheW [27]. Furthermore, based on the assumption that the chemotactic signaling state impacts the hydrophobic thickness of chemoreceptors, our model allows us to quantify the membrane contribution to chemotactic signaling. In agreement with previous experimental observations [35–37] we find a dependence of chemotactic signaling on lipid bilayer properties. Finally, we examine the possibility of membrane-mediated cooperative signaling among neighboring chemoreceptor trimers.

Models
In our analysis of membrane-mediated interactions between chemoreceptor trimers we follow the standard membrane-mechanical framework [31–33] for describing bilayer-protein interactions, and model chemoreceptor trimers as rigid membrane inclusions inducing elastic deformations in the surrounding lipid bilayer membrane. Such deformations can take the form of thickness deformations (Fig. 1A), which originate from a hydrophobic thickness mismatch between chemoreceptors and the lipid bilayer, and midplane (curvature) deformations (S1 Figure), which may be induced by a conical shape of chemoreceptor trimers resulting from a tilt in the transmembrane helices. To leading order, the elastic energies associated with thickness and midplane deformations decouple from each other, and can therefore be analyzed separately (see S1 Text section 1). We focus here on bilayer-chemoreceptor interactions and, hence, only consider the transmembrane regions of trimers in our model, with the peri- and cytoplasmic regions of trimers in Fig. 1A and S1 Figure only being shown for illustration.

In general, neighboring membrane proteins are expected to induce overlapping deformation fields of the bilayer membrane,
yielding [33,34] membrane-mediated interactions between proteins. Thus, membrane proteins can interact over several nanometers [33,34,30] without being in direct protein-protein contact. Membrane-mediated interactions due to thickness curvature deformations induced by identical proteins are generally expected [33] to be attractive/repulsive at small protein separations, with thickness deformations yielding stronger membrane-mediated interactions than curvature deformations. Based on previous work concerning the far-field limit of membrane-mediated interactions between conical membrane inclusions [39,40], analytic series solutions describing membrane-mediated interactions between proteins of arbitrary symmetry and at arbitrary separation have recently been developed [41]. Here we employ these analytic series solutions to determine the membrane-mediated interactions between chemoreceptor trimers. We find that midplane interaction energies resulting from the conical shape of trimers are typically well below $k_B T$ and more than an order of magnitude smaller than thickness interaction energies (see S1 Text section 2). We therefore focus on membrane-mediated interactions between chemoreceptor trimers due to thickness deformations (Fig. 1A).

Our biophysical model of membrane-mediated interactions between chemoreceptor trimers is based on the standard framework of membrane mechanics [42–46]. We represent the lipid bilayer within the Monge representation for curved surfaces using the functions $h_0(x,y)$ and $h(x,y)$, which define the heights of the hydrophilic-hydrophobic interfaces at the coordinates $(x,y)$ along the outer and inner membrane leaflets. The thickness deformations $u = u(x,y)$ correspond to

$$u(x,y) = \frac{1}{2} \left[ h_0(x,y) - h(x,y) - 2h_0 \right], \quad (1)$$

where $h_0$ is one-half the hydrophobic thickness of the unperturbed lipid bilayer. Following previous work on bilayer-protein interactions [31–33,38,46–55] we describe the energetic cost of thickness deformations by the functional

$$G_u[u] = \frac{1}{2} \int dxdy \left\{ K_u (\nabla^2 u)^2 + K_t \left( \frac{u}{h_0} \right)^2 + \sigma \left( 2 \frac{u}{h_0} + (\nabla u)^2 \right) \right\}. \quad (2)$$

where $K_u$ is the bending rigidity, $K_t$ is the stiffness associated with thickness deformations, and $\sigma$ is the membrane tension. The term $K_u (\nabla^2 u)^2$ in Eq. (2) captures the energetic cost of membrane bending, while the term $K_t (u/h_0)^2$ provides a simple description of the energetic cost of compressing or expanding the lipid bilayer. Typical measured values of $K_u$ and $K_t$ are $K_u = 20 \ k_B T$ and $K_t = 60 \ k_B T/\text{nm}^2$ [33,48], which we used for all the calculations described here. The term $\sigma (\nabla u)^2$ describes the effect of membrane tension on membrane undulations [42–47,49,50]. For generality we also allow for the term $2\sigma (u/h_0)$ in Eq. (2) [38,43,48], which captures the effect of membrane tension on lipid surface area under conservation of lipid volume. Phenomenological membrane deformation energies of the form in Eq. (2) have been employed to describe protein-induced bilayer thickness deformations in a range of systems [31–33,38,46–55], and can be systematically refined [41,56–68] to provide a more detailed model of bilayer-protein interactions.

The thickness deformation energy in Eq. (2) scales approximately with the square [31–33] of the hydrophobic mismatch which, in turn, is equal to the difference between one-half the trimer hydrophobic thickness, $h_0$, and $h_0$. A typical value of $h_0$ for the E. coli cytoplasmic membrane is $h_0 = 1.7 \ \text{nm}$ [69] while, for example, the approximate hydrophobic thickness of the chemoreceptor Trg is $h_0 = 2.025 \ \text{nm}$ [70]. The resulting hydrophobic mismatch $h_0 - h_0 = 0.325 \ \text{nm}$ yields a thickness deformation energy of the order of $50 \ k_B T$ for a single chemoreceptor trimer, which induces strong membrane-mediated interactions between neighboring trimers (see the Results section). For a given value of $h_t$, the value of $h_0$ and hence the magnitude and sign of the hydrophobic mismatch, can be tuned by changing the membrane composition which, as demonstrated for gramicidin [47,71] and mechanosensitive [49,72] channels, allows for direct experimental tests of membrane-mechanical models of bilayer-protein interactions. We study here membrane-mediated interactions between chemoreceptor trimers as a function of hydrophobic mismatch. Thus, while we use in our calculations a chemoreceptor hydrophobic thickness consistent with Trg, our conclusions can be applied equally to other chemoreceptors.

In the absence of detailed structural information on the transmembrane region of chemoreceptor trimers, we adopt a highly simplified model designed to capture two key features of chemoreceptor trimers: (1) As described above, we assume that chemoreceptors have a hydrophobic mismatch with the lipid bilayer, which induces membrane-mediated interactions between neighboring chemoreceptor trimers (Fig. 1A). (2) In addition, the characteristic three-fold symmetry of chemoreceptor trimers yields directionality in membrane-mediated interactions between chemoreceptor trimers (Fig. 1B). In particular, different relative orientations of neighboring chemoreceptor trimers produce distinct deformations of the bilayer membrane, resulting in a dependence of the energy of membrane-mediated interactions on the relative trimer orientation. Thus, membrane-mediated interactions between chemoreceptor trimers not only depend on the hydrophobic mismatch between chemoreceptors and the bilayer membrane [33], but also on the distinctive three-fold symmetry of chemoreceptor trimers. While the precise size of the transmembrane cross section of trimers is not crucial for our model predictions, we allow for a finite characteristic size of trimers and lipids, which imposes steric constraints on the minimum edge-to-edge separation of neighboring trimers.

Our simple model for the shape of chemoreceptor trimers (Fig. 1B) is consistent with recent electron cryo-tomography studies [19,20]. However, we focus here on the effects of generic aspects of chemoreceptor trimers, such as their symmetry, on membrane-mediated interactions, and our predictions do not rely on the detailed supramolecular shape of trimers. In particular, data obtained from electron cryo-tomography [19,20,73] mostly pertains to the cytoplasmic regions of chemoreceptor trimers, and the transmembrane structure of chemoreceptor trimers remains unknown. Indeed, the chemoreceptor dimers forming a trimer may spread apart within the membrane [19,20,73], with the lipid bilayer infiltrating chemoreceptor trimers. Such lipid-chemoreceptor complexes would imply membrane-mediated interactions between the chemoreceptor dimers forming a trimer. Here we do not consider membrane-mediated interactions within trimers and, instead, focus on membrane-mediated interactions between chemoreceptor trimers. Thus, our model of the transmembrane shape of chemoreceptor trimers (Fig. 1B) may correspond to chemoreceptor trimers composed of only proteins as well as lipid-chemoreceptor complexes. For simplicity, we assume a constant hydrophobic thickness of chemoreceptor trimers. More detailed descriptions would allow for a variation of the hydrophobic thickness along the trimer circumference, which may result from details of the transmembrane structure of chemoreceptors or the formation of lipid-chemoreceptor complexes.
Multiple lines of evidence [74–78] have indicated that chemoreceptor dimers signal the binding of a ligand across the cytoplasmic membrane through a piston-like sliding of one of the four transmembrane helices relative to the other three helices, by approximately 0.16 nm. This suggests that chemotactic signaling perturbs the hydrophobic surface of chemoreceptors and, indeed, it has been found [35–37] that bilayer-chemoreceptor interactions affect chemotactic signaling. Furthermore, the in vivo signaling response of chemoreceptors implies [9,16] that trimers exhibit strong cooperativity, and are either in the fully active or the fully inactive state. We account for these observations by assuming that chemoreceptor trimers can be active or inactive, with active and inactive trimers exhibiting a difference in hydrophobic thickness. For simplicity, we also assume that this difference in hydrophobic thickness is uniform along the trimer circumference, and is approximately equal to 0.16 nm as indicated by the piston model of chemotactic signaling [74–78]. (While consistent with the observed role of the membrane in chemotactic signaling, this working model is highly simplified; more detailed models would allow, for instance, for the possibility of a tilt in the transmembrane helices upon switching [74,79], for variations in the shift in hydrophobic thickness along the trimer circumference, and for possible differences in the hydrophobic surfaces exhibited by distinct chemoreceptors.) The predicted strength of the coupling between bilayer properties and chemotactic signaling depends on model details, but the basic mechanism for membrane-mediated cooperativity among chemoreceptors considered here relies only on a difference in hydrophobic thickness between active and inactive trimer states.

Results

Nucleation and growth of chemoreceptor lattices

We followed the approach developed in Refs. [39–41] to obtain analytic expressions for the energy $G_{\text{int}}$ of membrane-mediated interactions between chemoreceptor trimers due to thickness deformations. The energy $G_{\text{int}}$ is a function of center-to-center distance between trimers, $d$, relative trimer orientation, membrane tension, and hydrophobic mismatch (see S1 Text section 1). A negative value of the energy of membrane-mediated interactions, $G_{\text{int}} < 0$, implies energetically favorable interactions between chemoreceptor trimers. For a hydrophobic mismatch corresponding to chemoreceptors and the cytoplasmic membrane of *E. coli*, we find three regimes of membrane-mediated interactions between chemoreceptor trimers (Fig. 2A): (1) For trimer separations greater than $d \approx 12$ nm membrane-mediated interactions are negligible, yielding energies smaller than $k_BT$. (2) For intermediate trimer separations, from $d \approx 8.5–10.3$ nm (depending on relative trimer orientation) up to $d \approx 12$ nm, interactions are weakly unfavorable. (3) For small trimer separations, $d$ smaller than $d \approx 8.5–10.3$ nm (depending on relative trimer orientation), membrane-mediated interactions are strongly favorable. In particular, we find that for the smallest values of $d$ allowed by steric constraints on lipid size, corresponding to a minimum edge-to-edge separation between trimers of approximately 0.8 nm, membrane-mediated interactions can reduce the thickness deformation energy by more than 15 $k_BT$ compared to noninteracting chemoreceptor trimers.

Fig. 2. Membrane-mediated interactions yield attraction between chemoreceptor trimers. Calculated elastic interaction potentials between chemoreceptor trimers, $G_{\text{int}}$, as a function of center-to-center distance between neighboring trimers and (A) trimer orientation, (B) lipid monolayer hydrophobic thickness, and (C) membrane tension, $\sigma$. The solid lines at $d = 6.9$ nm indicate the face-on trimer separation measured by electron cryo-tomography of chemoreceptor lattices in the presence of CheA and CheW [19,22] and, where appropriate, steric constraints on trimer configurations due to finite trimer and lipid size are indicated by dashed vertical lines marking the end points of interaction potentials. For (A) and (B) we set $\sigma = 0$, and for (A) and (C) we used the monolayer thickness $h_0 = 1.7$ nm corresponding to the *E. coli* cytoplasmic membrane. All interaction potentials were calculated analytically under the approximation that trimers are weakly perturbed cylindrical inclusions.

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The interaction potentials in Fig. 2A show that membrane-mediated interactions yield strong attraction between chemoreceptor trimers over several nanometers, which suggests that membrane-mediated interactions may be sufficient for nucleation and growth of chemoreceptor lattices. Indeed, while chemoreceptors interact with CheA and CheW to form ordered lattices [19,20], clustering of chemoreceptors requires neither CheA nor CheW [27,28]. Furthermore, superresolution light microscopy of chemoreceptor clusters has suggested [29,30] that chemoreceptor lattices self-assemble by stochastic nucleation of small clusters and capture of diffusing receptors by preexisting clusters. Fig. 2A implies that membrane-mediated interactions provide a plausible biophysical mechanism for the efficient self-assembly of chemoreceptor lattices via stochastic nucleation and capture. In particular, based on the statistical mechanics of phase segregation [30,81] the interaction energies in Fig. 2A allow us to estimate the critical trimer concentration for nucleation and growth of chemoreceptor lattices in the *E. coli* cytoplasmic membrane (see S1 Text section 3). We find that the critical trimer concentration for clustering is already reached with approximately 15 chemoreceptor trimers in the cytoplasmic membrane. This means that, even if trimers are very dilute in the cytoplasmic membrane, membrane-mediated interactions can lead to nucleation and growth of chemoreceptor lattices.

Our model predicts that chemoreceptor clustering due to membrane-mediated interactions shows a characteristic dependence on hydrophobic mismatch (Fig. 2B) and membrane thickness (Fig. 2C). In Fig. 2B we consider a range in hydrophobic mismatch which may be realized, for instance, by varying the tail lengths in phosphatidylethanolamine (PE) lipid bilayers from PC10 to PC24 [82], while in Fig. 2C we consider values of membrane tension up to the approximate rupture tension of lipid bilayers [48,82]. Fig. 2B shows that membrane-thickness-mediated interactions between chemoreceptor trimers vanish when the bilayer hydrophobic thickness matches the chemoreceptor hydrophobic thickness, and increase in magnitude with increasing magnitude of hydrophobic mismatch. Fig. 2C predicts that, for a hydrophobic mismatch corresponding to chemoreceptors and the cytoplasmic membrane of *E. coli*, membrane-mediated interactions between chemoreceptor trimers become more pronounced with increasing membrane tension, yielding an increased propensity for chemoreceptor clustering.

The basic qualitative features of the interaction potentials in Fig. 2 can be understood from the thickness deformation field due to a single membrane inclusion. Consider, for simplicity, a cylindrical membrane inclusion with a hydrophobic thickness that exceeds the unperturbed bilayer hydrophobic thickness. The resulting thickness deformation decays approximately exponentially around the membrane inclusion with a characteristic decay length \( \lambda = (\hbar_k R_{K} / K) \approx 1 \) nm [33,48]. The decaying thickness deformation will overshoot [47,52], leading to a zone of compression of the lipid bilayer, before the deformation eventually approaches zero (S3 Figure). The attractive regime of membrane-mediated interactions in Fig. 2 corresponds to edge-to-edge separations of up to approximately \( 4 \lambda \), for which thickness deformations mainly overlap in the region of initial exponential decay and the overall deformation footprint of the two trimers is reduced compared to noninteracting trimers (Fig. 1A). For edge-to-edge separations from approximately \( 4 \lambda \) to \( 6 \lambda \), the compressed and expanded membrane regions induced by the two trimers strongly overlap, which results in frustration of membrane deformations and the repulsive regime in Fig. 2. Finally, the noninteracting regime in Fig. 2 corresponds to edge-to-edge separations greater than approximately \( 6 \lambda \), for which there is only little overlap in the thickness deformations induced by the two trimers. The scale of the maximum interaction energies in Fig. 2 is set by the single-cylinder thickness deformation energy \( G_{\text{single}} \approx \sqrt{2 \pi R (h_l - h_0)^2 / l^3} \approx 30 \) k_BT [33] for a radius \( R = 3.1 \) nm. Also, since \( G_{\text{single}} \propto (h_l - h_0)^2 \), the strength of the attractive and repulsive regimes increases with the magnitude of the hydrophobic mismatch as in Fig. 2B. Moreover, the single-inclusion thickness deformation energy increases with membrane tension if, as is the case for chemoreceptors in the cytoplasmic membrane, the hydrophobic mismatch takes a positive value [48], yielding an increase in the strength of membrane-mediated interactions with increasing membrane tension as in Fig. 2C.

**Gateway to chemoreceptor lattice architecture**

Fig. 2 shows that membrane-mediated interactions between chemoreceptor trimers are strongly directional, and reflect the three-fold symmetry of trimers. We find two dominant trimer configurations as a function of trimer separation: (1) In Fig. 2A, for trimer separations greater than \( d \approx 8.3 \) nm, the tip-on configuration (red inset) is energetically most favorable. (2) For small trimer separations, \( d \) smaller than \( d \approx 7.3 \) nm, the face-on configuration (blue inset) is most favorable. These two regimes occur because the tip-on configuration yields the smallest edge-to-edge separation (and thus the longest-range interactions, Fig. 1B left panel), while the face-on configuration maximizes the membrane area over which trimer-induced thickness deformations can overlap (and thus provides the maximum interaction strength overall, Fig. 1B right panel). We estimate that the energy difference between tip-on and face-on configurations can be more than \( 10 \) k_BT for the minimum trimer separations allowed by steric constraints in the two configurations. In particular, membrane-mediated interactions favor the face-on trimer configuration at the observed separation \( d = 6.9 \) nm (grey vertical line) as measured by electron cryo-tomography of chemoreceptor lattices in *E. coli* as well as other organisms [19,22], in the presence of CheA and CheW. The face-on configuration of trimers predicted by our model for small trimer separations has been observed in chemoreceptor lattices in a variety of different organisms [19,20] and allows the formation of chemoreceptor-CheW-CheA complexes, yielding a well-defined trimer separation due to direct protein-protein interactions.

Fig. 2 implies a scenario for the assembly of chemoreceptor lattices in which the tip-on trimer configuration is a gateway state yielding attraction between chemoreceptor trimers over several nanometers, with the directionality of membrane-mediated interactions ensuring that, at small separations, trimers are arranged in the face-on orientation allowing further stabilization through direct protein interactions mediated by CheA and CheW [19,20]. In particular, the interaction potentials in Fig. 2 suggest that the face-on trimer configuration found in chemoreceptor lattices [19,20] could be achieved through the sequence of gateway states shown in Fig. 3. For large \( d \), the tip-on configuration is strongly favored (for ease of visualization, the tip-on configuration is set as the zero of \( G_{\text{int}} \) in Fig. 3). As the trimer separation shrinks below the steric constraint on the tip-on configuration, the membrane deformation energy can be lowered further by a symmetric rotation of the chemoreceptor trimers (S1 Video), ultimately yielding the observed face-on trimer configuration [19,20] as the lowest-energy configuration, thus ensuring correct assembly of chemoreceptor lattices. Consistent with the results in Fig. 2, we find that the membrane-mediated interactions stabilizing the sequence of gateway states in Fig. 3 vanish for lipid bilayers matching the chemoreceptor hydrophobic thickness and increase with the magnitude of the hydrophobic mismatch (Fig. 3A).
Stabilization of chemoreceptor lattice architecture

A simple arrangement of trimers in chemoreceptor lattices would be a close-packed hexagonal lattice structure (Fig. 4 grey insets, S4A Figure) in which each trimer has six nearest-neighbors and, hence, the number of nearest-neighbor interactions is maximized. However, electron cryo-tomography has shown [19,20] that chemoreceptor trimers are not closely packed in chemoreceptor lattices but rather form a honeycomb lattice in which each trimer has three nearest-neighbors arranged in the face-on orientation (Fig. 4 blue insets, S4B Figure), which allows formation of an extended lattice composed of chemoreceptor trimers, CheA, and CheW. To elucidate the stability of the observed face-on honeycomb-lattice architecture we calculated the energy per chemoreceptor trimer resulting from membrane-mediated interactions due to thickness deformations, $G_{\text{lat}}$, in face-on honeycomb, tip-on honeycomb (Fig. 4 red insets, S4C Figure), and hexagonal lattices. We find that, while tip-on honeycomb and hexagonal lattices can be energetically favorable for large lattice spacings, both these structures are unstable to the formation of a face-on honeycomb lattice with small lattice spacing, which provides the minimum-energy lattice architecture (Fig. 4A). This conclusion is robust with respect to variations in hydrophobic mismatch (Fig. 4B) and membrane tension (Fig. 4B inset). In contrast, cylindrical membrane inclusions, which do not exhibit directional interactions, would yield the hexagonal lattice as the minimum-energy structure. Thus, the directionality of membrane-mediated interactions stabilizes the observed face-on honeycomb lattice architecture against the tip-on honeycomb and hexagonal lattice structures. Specifically, the three-fold symmetry of trimers allows honeycomb ordering of chemoreceptor lattices, and thus further stabilization of a well-defined lattice constant through direct protein interactions with CheA and CheW [19,20].

Fig. 4B predicts that, for the lattice spacings indicated by arrows in Fig. 4A, the strength of favorable interactions between chemoreceptor trimers in face-on honeycomb, tip-on honeycomb, and hexagonal lattices grows monotonically with increasing hydrophobic mismatch between lipid bilayer and chemoreceptors, as well as with increasing membrane tension. For the lattice spacings in Fig. 4A yielding a crossover from favorable ($G_{\text{lat}}<0$) to unfavorable ($G_{\text{lat}}>0$) lattice energies we obtain a more complex dependence of the lattice energy on bilayer hydrophobic thickness and membrane tension (S5 Figure). In particular, for such crossover lattice spacings our model predicts favorable lattice energies for bilayer hydrophobic thicknesses exceeding the chemoreceptor hydrophobic thickness, with unfavorable lattice energies for bilayer hydrophobic thicknesses smaller than the chemoreceptor hydrophobic thickness. This can be understood by noting that the decay length $\lambda$ increases with $h_0$, thus shifting membrane-mediated interactions into the attractive regime if $h_0$ increases beyond $h_c$, and vice versa. Finally, we note that the lattice energy due to membrane-mediated interactions between chemoreceptor trimers is dominated by nearest-neighbor interactions, with longer-range interactions only yielding minor shifts in the lattice energy (S6 Figure).

Transition in chemoreceptor lattice architecture

Our calculations imply that close-packed hexagonal lattices of chemoreceptor trimers are metastable in the sense that the hexagonal lattice structure is only a local minimum of the membrane deformation energy, with the global minimum provided by the face-on honeycomb lattice (Fig. 4). However, the membrane area per trimer in honeycomb lattices is greater than the membrane area per trimer in hexagonal lattices—by 50% if both lattice structures have the same trimer separation and by 15% for the trimer separations indicated by arrows in Fig. 4. Thus, in situations where the clustering of chemoreceptor trimers is strongly constrained by the available membrane area, membrane-mediated interactions may yield hexagonal chemoreceptor lattices. On the basis of electron microscopy it has indeed been observed [83–85] that overexpression of chemoreceptors results in hexagonal lattices of trimers in the cytoplasmic...
membrane. The observed two-dimensional hexagonal lattices were distinct from the “zippered” cluster structures [83] also found in overexpression experiments, which strongly bend the membrane and form interdigitated protein contacts. In agreement with our model, in the case of overexpression the clustering of trimers, the stability of the lattice, and the two-dimensional hexagonal lattice architecture did not rely on the presence of CheA and CheW, although the presence of CheA and CheW yielded more ordered lattice structures and modified the lattice spacing [84,85]. The trimer orientation in the observed two-dimensional hexagonal lattices [84,85] is consistent with the hexagonal lattice architecture of trimers shown in Fig. 4 (grey insets).

**Fig. 4. Membrane-mediated interactions yield the observed architecture of chemoreceptor lattices.** Calculated elastic interaction energy per trimer, $G_{\text{nat}}$, in face-on honeycomb (blue), tip-on honeycomb (red), and hexagonal (black) lattices as a function of (A) center-to-center distance between neighboring trimers and (B) monolayer hydrophobic thickness and membrane tension (inset) at $d = 6.3$ nm (face-on honeycomb lattice; blue curve), 7.2 nm (hexagonal lattice; black curve), and 8.3 nm (tip-on honeycomb lattice; red curve), corresponding to the lattice spacings indicated by arrows in (A). The solid vertical line at $d = 6.9$ nm in (A) shows the trimer separation observed in face-on chemoreceptor lattices [19,22], and dashed vertical lines in (A) indicate steric constraints on tip-on honeycomb (red) and hexagonal (black) lattice configurations. For (A) we assumed a membrane with $\sigma = 0$ and a monolayer thickness $h_0 = 1.7$ nm corresponding to the *E. coli* cytoplasmic membrane (shown by a dashed vertical line in (B)). For the main panel in (B) we set $\sigma = 0$ and the curves in the inset were obtained with $h_0 = 1.7$ nm. All lattice energies were calculated from pairwise nearest-neighbor interaction potentials as shown in Fig. 2. doi:10.1371/journal.pcbi.1003932.g004

**Localization of chemoreceptor lattices to cell poles**

As noted in the Models section, chemoreceptor trimers induce midplane deformations in addition to thickness deformations. While midplane interaction energies are typically negligible compared to thickness interaction energies (see S1 Text section 2), midplane deformations provide a simple mechanism for segregation of chemoreceptor trimers to the cell poles [86]. In particular, the energetic cost of trimer-induced curvature deformations depends on the interplay between the conical shape of chemoreceptor trimers [87] and the preferred curvature of the surrounding lipid bilayer: Since the average membrane radius of curvature at the poles of *E. coli* is approximately twice that of the midcell region, and both have the same sign as the radius of curvature of chemoreceptor trimers, midplane deformations may act as curvature sensors mediating localization of chemoreceptor trimers to the cell poles.

The energy of trimer-induced midplane deformations can be estimated using a variety of different approaches [48,50,60,81]. Independent of the particular model formulation, we find that for a single chemoreceptor trimer the difference in midplane deformation energy between the poles and midcell of *E. coli* is well below $k_BT$ (see S1 Text section 2). This suggests that curvature deformations are not able to localize individual chemoreceptor trimers to the cell poles. However, as described above, we also find that strong membrane-mediated interactions due to thickness deformations effectively bind chemoreceptor trimers into chemoreceptor lattices, which may be further stabilized by interactions with CheA and CheW. For a lattice composed of $N$ chemoreceptor trimers we estimate an energy difference

$$
0.055Nk_BT \lesssim \Delta G_{\text{midcell-pole}} \lesssim 0.13Nk_BT
$$

between the midcell and poles of *E. coli* in the regime of weak interactions due to midplane deformations, where the lower and upper bounds correspond to different model formulations (see S1 Text section 2). Thus, bilayer-trimer interactions yield only weak curvature sensitivity for small chemoreceptor lattices but can readily induce localization of large chemoreceptor lattices to convex regions of the cytoplasmic membrane such as the cell poles.

Large chemoreceptor lattices composed of thousands of receptors (for which Eq. (3) yields $\Delta G_{\text{midcell-pole}} \gtrsim 20k_BT$) are indeed observed predominantly at the cell poles, while smaller chemoreceptor lattices are also found in the midcell regions [21–25,27–30].

**Effects of bilayer-chemoreceptor interactions on chemotactic signaling**

Reconstitution of chemoreceptors in bilayer vesicles [36] and nanodiscs [35] has indicated that the signaling properties of chemoreceptors depend on the composition of lipid bilayers. Furthermore, modification of the transmembrane properties of chemoreceptors by site-directed mutagenesis has shown [37] that bilayer-chemoreceptor interactions influence chemotactic signaling. Within the simple “piston” model of chemotactic signaling such a coupling between chemoreceptor function and lipid bilayer properties arises naturally—specifically, the on and off states of chemoreceptors differ in their hydrophobic mismatch with the lipid bilayer and thus in their bilayer deformation energies. Assuming a uniform 0.16 nm difference in hydrophobic thickness between on and off states, our model predicts that for *E. coli* the
membrane contribution to the total free-energy difference between on and off states of a single chemoreceptor trimer is greater than $20 \ k_B T$ in magnitude, and can vary over more than $60 \ k_B T$ with bilayer or trimer hydrophobic thickness (S7 Figure). In agreement with experiments [35–37] we therefore find that shifts in the membrane contribution to the trimer transition energy due to modification of lipid composition or chemoreceptor transmembrane properties can dominate over shifts in the transition energy due to chemoreceptor methylation, which are of the order of $1 \ k_B T$ per methyl group [88]. In addition, our model predicts a dependence of chemotactic signaling on membrane elastic properties such as membrane tension (S7 Figure inset). In particular, we find that variation in membrane tension can shift the membrane contribution to the on-off transition energy by up to $6 \ k_B T$ which, again, is comparable to shifts in the trimer transition energy due to chemoreceptor methylation [88].

We speculate that membrane-mediated interactions between chemoreceptor trimers could contribute to cooperativity among chemoreceptor trimers, complementing the contribution of direct protein interactions mediated by CheA and CheW [9,10,19,20]: Consider a chemoreceptor trimer in the on state, with a neighboring trimer in the off state (Fig. 5A upper panel). Assuming that the two trimers induce distinct thickness deformations due to their different signaling states, membrane-mediated interactions are energetically unfavorable at small trimer separations [33]. If, however, both trimers are in the off state (Fig. 5A lower panel), membrane-mediated interactions are strongly favorable. Thus, the presence of a neighboring trimer in the off state lowers, via membrane-mediated interactions, the free energy of the off state (and similarly a neighbor in the on state lowers the free energy of the on state), potentially yielding membrane-mediated cooperativity among chemoreceptor trimers.

In order to quantify the above mechanism for membrane-mediated cooperativity we calculated the membrane contribution to the free-energy difference between the on and off state of a chemoreceptor trimer, $\Delta G_{\text{on-off}}$, for the trimer orientation [19,20] and separation [19,22] observed in chemoreceptor lattices (Fig. 5B). Consistent with our results for a single trimer (S7 Figure) we find that there is a substantial membrane contribution to the transition energy of trimers in chemoreceptor lattices. Since chemoreceptors are functionally required to operate near zero transition energy [89], this membrane contribution must be compensated by internal protein contributions to the transition energy. However, Fig. 5B also shows that membrane-mediated interactions can lower the transition energy by up to approximately $15 \ k_B T$ depending on the activity state of neighboring trimers, or by approximately $5 \ k_B T$ for each nearest-neighbor trimer in the off state. This cooperative shift in the transition energy is comparable to the shift in the trimer transition energy obtained by methylation of all 24 modification sites on a trimer [88], and may therefore be relevant for the cooperative signaling properties of chemoreceptor lattices. Furthermore, Fig. 5B shows that the strength of the predicted cooperative interactions among chemoreceptor trimers is robust with respect to variations in hydrophobic mismatch (Fig. 5B main panel) and membrane tension (Fig. 5B inset).

**Discussion**

Fluorescence experiments have suggested a stochastic model for chemoreceptor lattice formation [28,30] in which self-assembly of chemoreceptor lattices proceeds by nucleation and growth, without direct cytoskeletal involvement or active transport. Lattices consist of trimers-of-dimers of chemoreceptors and require the linker/kinase CheA and the linker CheW [19,20] for their function. However, clustering of chemoreceptor trimers requires neither CheA nor CheW [27,28]. In common with other membrane proteins [31,32], chemoreceptor trimers are expected...

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**Fig. 5. Membrane-mediated interactions may contribute to cooperative signaling.** (A) We assume the trimer hydrophobic thickness differs by $\delta h = 0.08 \text{ nm}$ between the on state (white trimers, $h_{\text{trimer}} = 2.025 \text{ nm}$) and the off state (grey trimers, $h_{\text{trimer}} + \delta h = 2.105 \text{ nm}$) according to the piston model of chemotactic signaling. For ease of visualization, the magnitude of $\delta h$ is exaggerated in the schematic illustrations of the off state. (B) Calculated difference in the thickness deformation energy associated with the on and off states of a chemoreceptor trimer, $\Delta G_{\text{on-off}}$, in a face-on honeycomb lattice with $n$ nearest neighbors in the off state as a function of membrane hydrophobic thickness and membrane tension (inset) at the trimer spacing $d = 6.9 \text{ nm}$ measured for face-on chemoreceptor lattices [19,22]. For the main panel the membrane tension is $\sigma = 0$ and for the inset the monolayer hydrophobic thickness is $h_0 = 1.7 \text{ nm}$ corresponding to the E. coli cytoplasmic membrane (indicated by a dashed vertical line in the main panel, with the dashed horizontal lines showing $\Delta G_{\text{on-off}}$ for $n = 0.3$ neighboring trimers in the off state). All cooperative interactions were calculated using pairwise trimer-trimer interaction potentials as in Fig. 4.

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to deform the surrounding lipid bilayer, leading to membrane-mediated interactions [33,34] between neighboring trimers. To quantify the role of membrane-mediated interactions in the assembly and architecture of chemoreceptor lattices we have developed a biophysical model of bilayer-chemoreceptor interactions.

Our biophysical model of bilayer-chemoreceptor interactions shows that membrane-mediated interactions yield attractive interactions between chemoreceptor trimers over several nanometers and hence provide a biophysical mechanism for cluster self-assembly. Our model predicts that the tip-on orientation of a pair of chemoreceptor trimers is a “gateway” state during assembly, whereas at smaller trimer separations, membrane-mediated interactions favor the face-on orientation of each trimer pair also observed in the presence of CheA and CheW [19,20]. Furthermore, we predict that membrane-mediated interactions are strong enough to induce cluster formation even for a trimer concentration of only ~15 trimers per E. coli cell. This suggests a scenario for self-assembly in which membrane-mediated interactions produce clusters of chemoreceptors, which are further stabilized and ordered through protein interactions mediated by CheA and CheW. Since the range of membrane-mediated interactions is set by the elastic decay length of thickness deformations, which is a bilayer property, these conclusions do not rely on the detailed size and shape of trimers in our model of chemoreceptor trimers in Fig. 1B. In particular, for a given bilayer membrane the range of membrane-mediated interactions between trimers, measured in terms of the center-to-center distance between trimers, is determined by the edge-to-edge separation of trimers for each trimer configuration, yielding a longer (shorter) range of membrane-mediated interactions for larger (smaller) trimer sizes. In agreement with experimental observations [30], the strongly favorable interactions between trimers at small separations in Figs. 2 and 3 are expected to yield an approximately exponential size distribution of chemoreceptor clusters [30,90].

In vivo electron cryo-tomography has revealed [19–25] that chemoreceptor lattices are not close-packed hexagonal arrays. Instead, chemoreceptor trimers form honeycomb lattices with a trimer at each vertex (S4B Figure), and a well-defined face-on orientation of trimers [19,20]. Our model predicts that membrane-mediated interactions favor this face-on, honeycomb architecture of the lattice. In particular, we find that the three-fold symmetry and directionality of membrane-mediated interactions favor a honeycomb lattice (three neighbors per trimer) over a close-packed hexagonal lattice (six neighbors per trimer). Thus, while interactions with CheA and CheW are expected to determine the observed separation of trimers in chemoreceptor lattices [19,22] and are likely to be adequate to define the observed lattice symmetry [73,91], we find that membrane-mediated interactions can drive the formation of diffuse, less ordered membrane-mediated interactions for larger (smaller) trimer sizes. In agreement with experimental observations [30], the strongly favorable interactions between trimers at small separations in Figs. 2 and 3 are expected to yield an approximately exponential size distribution of chemoreceptor clusters [30,90].

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Our model of bilayer-chemoreceptor interactions suggests that localization of large chemoreceptor lattices to the cell pole is simply a consequence of the conical shape of individual chemoreceptor trimers [37], and neither requires interactions with CheA and CheW [27] nor curvature-mediated interactions among trimers. In agreement with experimental observations [21–25,27–30], our model implies that large chemoreceptor clusters will tend to localize at the cell poles, while smaller chemoreceptor clusters can be distributed throughout the midcell regions. This mechanism for localization of large chemoreceptor lattices due to curvature sensing by individual chemoreceptor trimers is to be contrasted with a previously proposed mechanism [86] which assumes that trimers interact to yield a non-zero global intrinsic curvature of chemoreceptor lattices. A distinguishing difference between the localization mechanism proposed here and in Ref. [86] is that, according to the latter, chemoreceptor clusters should have a finite characteristic size set by the energy balance between short-range attraction and curvature-mediated long-range repulsion between trimers, whereas our model indicates that curvature-mediated interactions are too weak to limit cluster size in the absence of CheA and CheW. Fluorescence experiments [28,30] measuring chemoreceptor cluster size in the absence of CheA and
CheW may be able to distinguish between these two related scenarios for curvature-driven localization of large chemoreceptor lattices.

FRET experiments have revealed [8,9] that chemoreceptors signal in cooperative teams of coupled trimers [9,11–16]. Cooperative interactions among neighboring trimers are believed to be mediated by CheA and CheW [9,10,19,20]. Our model of chemotactic signaling shows that, provided there is a substantial change in chemoreceptor hydrophobic thickness upon signaling, membrane-mediated interactions between chemoreceptor trimers [85] can in principle yield cooperative interaction energies of the order of several $k_BT$. This would be sufficient [92] to account at least in part for the observed cooperative signaling properties of chemoreceptor lattices. Indeed, electron cryo-tomography indicates that honeycomb lattices of chemoreceptor trimers are somewhat disordered, with the degree of disorder being a matter of debate [21–25]. While interactions between chemoreceptor trimers mediated by CheA and CheW [9,10,19,20] rely on a regular lattice structure, membrane-mediated interactions are less sensitive to defects in the chemoreceptor lattice. Thus, membrane-mediated interactions may increase the robustness of cooperative signaling teams, and complement cooperative interactions mediated by CheA and CheW [9,10,19,20]. Consistent with our biophysical model of chemotactic signaling it has been found using homo-FRET [37,93] that the in vivo signaling response of chemoreceptors depends on membrane-mechanical properties such as membrane tension. However, homo-FRET has so far not produced any evidence for cooperativity among chemoreceptor trimers in the absence of CheA and CheW [94]. Chemoreceptor clusters formed in the absence of CheA and CheW are more diffuse than chemoreceptor lattices formed in the presence of CheA and CheW [27,28], which may substantially reduce membrane-mediated cooperativity.

Our model of chemotactic signaling predicts that shifts in the membrane contribution to the total free-energy difference between on and off states of chemoreceptor trimers due to changes in membrane composition or membrane tension can be comparable to shifts in the chemoreceptor transition energy due to receptor methylation [88], and can therefore be functionally relevant. In agreement with these predictions, it has been found that modifying the composition of lipid bilayers [35,36] or bilayer-chemoreceptor interface [37] affects chemotactic signaling. In particular, changes in lipid composition can strongly bias chemoreceptors towards the active or inactive state [35,36], and the baseline signaling state of chemoreceptors can be controlled by site-directed mutagenesis of chemoreceptor transmembrane helices [37]. Thus, in analogy to gramicidin [47,71] and mechanosensitive [49,72] channels, systematic variation of the membrane lipid composition, the chemoreceptor hydrophobic thickness, or membrane-mechanical properties such as membrane tension may allow quantitative experimental tests of our biophysical model of the role of membrane-mediated interactions in the assembly and architecture of chemoreceptor lattices, as well as our speculation of a membrane-mediated contribution to chemotactic signaling and cooperativity.

Supporting Information

S1 Figure  Schematic of midplane deformations induced by chemoreceptor trimers. To complement the model of membrane-mediated interactions in Fig. 1 of the main text, we have estimated the midplane deformations induced by chemoreceptor trimers. Trimmers can deform the bilayer midplane $h(x,y)$ (dashed red line) by an angle $\alpha$ at the bilayer-trimer interface, and membrane-mediated interactions tilt trimers by an angle $\beta$ in the $x$-direction.

S2 Figure  Thickness deformation fields of chemoreceptor trimers. Thickness deformations $u = u(x,y)$ induced by two chemoreceptor trimers in (A) the tip-on and (B) the face-on orientation. Both chemoreceptor trimers are in the on state. (See also Fig. 1B of the main text.)

S3 Figure  Thickness deformation profile around a cylindrical membrane inclusion. The thickness deformation field $u$ is calculated using the same parameter values as in Fig. 2 of the main text, but for a single cylindrical membrane inclusion of radius $R = 3.1$ nm. The variable $r$ denotes the distance from the inclusion boundary and is measured in units of thickness deformation decay length $\delta \approx 1$ nm. The thickness deformation field $u$ is measured in units of hydrophobic mismatch $h_i - h_0$.

S4 Figure  Schematic of chemoreceptor lattice symmetries. (A) Hexagonal lattice. (B) Face-on honeycomb lattice. (C) Tip-on honeycomb lattice.

S5 Figure  Membrane-mediated interactions in chemoreceptor lattices. Calculated elastic interaction energy per trimer, $G_{\text{latt}}$, in face-on honeycomb (blue), tip-on honeycomb (red), and hexagonal (black) lattices as a function of (A) center-to-center distance between neighboring trimers (data as in Fig. 4A of the main text and shown here for completeness) and (B,C,D) monolayer hydrophobic thickness and membrane tension (insets) at $d = 7.2$ nm, $8.5$ nm, and $10.5$ nm, as indicated by arrows in (A). The solid vertical line at $d = 6.9$ nm in (A) shows the trimer separation observed in face-on honeycomb chemoreceptor lattices [19,22], and dashed vertical lines in (A) indicate steric constraints on lattice configurations. As in Fig. 4A of the main text, we assumed for (A) a membrane with $\sigma = 0$ and a monolayer thickness $h_0 = 1.7$ nm corresponding to the E. coli cytoplasmic membrane (shown by dashed vertical lines in (B–D)). For the main panels in (B–D) we set $\sigma = 0$ and the insets in (B–D) were obtained with $h_0 = 1.7$ nm. All lattice energies were calculated from pairwise nearest-neighbor interaction potentials as shown in Fig. 2 of the main text.

S6 Figure  Effect of higher-order interactions on lattice energies. Lattice energies for face-on honeycomb, tip-on honeycomb, and hexagonal lattices of chemoreceptor trimers allowing for up to nearest-neighbor (solid curves; as in Fig. 4A of the main text), next-nearest neighbor (dashed curves), and next-next-nearest neighbor (dotted-dashed curves) interactions. For each lattice symmetry, nearest neighbors, next-nearest neighbors, and next-next-nearest neighbors are indicated by the color coding in the inset, with $G_{\text{latt}}$ corresponding to the central trimer with white filling. All lattice energies were calculated from pairwise interaction potentials as shown in Fig. 2 of the main text.

S7 Figure  Membrane contribution to the transition energy of a single chemoreceptor trimer. For the main panel we set $\sigma = 0$ and for the inset we used the value $h_0 = 1.7$ nm corresponding to the E. coli cytoplasmic membrane.

S1 Video  Video of gateway to assembly of face-on trimer configuration. Gateway states in Fig. 5 of the main text.
References

1. Berg HC (2000) Motile behavior of bacteria. Physics Today 53: 24–29.
2. Segall JE, Block SM, Berg HC (1986) Temporal comparisons in bacterial chemotaxis. Proc Natl Acad Sci U S A 83: 8907–8911.
3. Bray D, Levin MD, Morton-Firth CJ (1996) Receptor clustering as a cellular mechanism to control sensitivity. Nature 383: 45–50.
4. Duke TA, Bray D (1999) Heightened sensitivity of a lattice of membrane receptors. Proc Natl Acad Sci U S A 96: 10104–10108.
5. Bray D (2002) Bacterial chemotaxis and the question of gain. Proc Natl Acad Sci U S A 99: 7176–7178.
6. Bray D, Duke TA (2004) Conformational spread: The propagation of allotropic states in large multiprotein complexes. Annu Rev Biophys Biomol Struct 33: 53–73.
7. Lan G, Schultmeier S, Sourjik V, Tu Y (2011) Adapting locally and acting globally: strategy to maintain high chemoreceptor sensitivity in complex environments. Mol Sys Biol 7: 475.
8. Sourjik V, Berg HC (2002) Receptor sensitivity in bacterial chemotaxis. Proc Natl Acad Sci U S A 99: 123–127.
9. Sourjik V, Berg HC (2004) Functional interactions between receptors in bacterial chemotaxis. Nature 428: 437–441.
10. Sourjik V (2004) Receptor clustering and signal processing in E. coli chemotaxis. Trends Microbiol 12: 569–576.
11. Mello BA, Tu Y (2003) Quantitative modeling of sensitivity in bacterial chemotaxis: The role of coupling among different chemoreceptor species. Proc Natl Acad Sci U S A 100: 8023–8028.
12. Shimizu TS, Aksenov SV, Bray D (2003) A spatially extended stochastic model of the bacterial chemotaxis signalling pathway. J Mol Biol 329: 291–309.
13. Mello BA, Shaw L, Tu Y (2004) Effects of receptor interaction in bacterial chemotaxis. Biophys J 87: 1578–1595.
14. Mello BA, Tu Y (2005) An allosteric model for heterogeneous receptor complexes: Understanding bacterial chemotaxis responses to multiple stimuli. Proc Natl Acad Sci U S A 102: 17354–17359.
15. Klemm JE, Endres RG, Skoge M, Meir Y, Wingreen NS (2006) Chemosensing in Escherichia coli: two regimes of two-state receptors. Proc Natl Acad Sci U S A 103: 17266–17271.
16. Hansen CH, Sourjik V, Wingreen NS (2010) A dynamic-signaling-team model for chemotaxis receptors in Escherichia coli. Proc Natl Acad Sci U S A 107: 17170–17175.
17. Kim KK, Yokota H, Kim SH (1999) Four-helical bundle structure of the cytoplasmic domain of a serine chemotaxis receptor. Nature 400: 787–792.
18. Studdert CA, Parkinson JS (2004) Crosslinking snapshots of bacterial chemoreceptor squads. Proc Natl Acad Sci U S A 101: 2117–2122.
19. Boal D (2002) Mechanics of the Cell. Cambridge: Cambridge University Press.
20. Haselwandter CA, Phillips R (2013) Directional interactions and cooperativity between mechanosensitive membrane proteins. EPL 101: 68002.
21. Boal D (2002) Mechanics of the Cell. Cambridge: Cambridge University Press.
22. Phillips R, Koudiev J, Thiot J, Garcia HG (2015) Physical Biology of the Cell. London and New York: Garland Science, 2nd edition.
23. Szilassi U (1997) Configurations of fluid membranes and vesicles. Adv Phys 46: 13–117.
24. Salvan S (2003) Statistical Thermodynamics of Surfaces, Interfaces, and Membranes. Boulder: Westview Press.
25. Fourrier JB (1999) Microscopic membrane elasticity and interactions among membrane inclusions: Interplay between the shape, dilation, tilt, and tilt difference modes. Eur Phys J B 11: 261–272.
26. Huang HV (1986) Deformation free energy of bilayer membrane and its effect on gramicidin channel lifetime. Biophys J 56: 1061–1070.
27. Ursell T, Koudiev J, Reeves D, Wiggins P, Phillips R (2008) The role of lipid bilayer mechanics in mechanosensation. In: Kamikin A, Kiselev I, editors, Mechanosensitivity in Cells and Tissues 1: Mechanosensitive Ion Channels. New York: Springer Press, pp. 37–70.
28. Wiggins P, Phillips R (2004) Analytic models for mechanotransduction: Gating a mechanosensitive channel. Proc Natl Acad Sci U S A 101: 4071–4076.
29. Wiggins P, Phillips R (2005) Membrane-protein interactions in mechanosensitive channels. Biophys J 88: 900–902.
30. Landheer JA (2006) Regulation of membrane protein function by lipid bilayer elasticity—a single molecular technology to measure the bilayer properties experienced by an embedded protein. J Phys: Condens Matter 18: S1305–S1314.
31. Dan N, Pincus P, Safran SA (1993) Membrane-induced interactions between inclusions. Langmuir 9: 2768–2771.
32. Nielsen C, Goulas M, Andersen OS (1998) Energetics of inclusion-induced bilayer deformations. Biophys J 74: 1966–1983.
33. Dan N, Safran SA (1999) Effect of lipid characteristics on the structure of transmembrane proteins. Biophys J 75: 1410–1414.
34. Aranda-Espinosa H, Berman A, Dan N, Pincus P, Safran S (1996) Interaction between inclusions embedded in membranes. Biophys J 71: 648–656.
56. Partenskii MB, Jordan PC (2002) Membrane deformation and the elastic energy of insertion: Perturbation of membrane elastic constants due to peptide insertion. J Chem Phys 117: 107681–10776.

57. Partenskii MB, Miloshevsky GV, Jordan PC (2004) Membrane insertions as coupled harmonic oscillators: Effects due to anisotropic membrane slope relaxation. J Chem Phys 120: 2183–2193.

58. Braanigan G, Brown FLH (2006) A consistent model for thermal fluctuations and protein-induced deformations in lipid bilayers. Biophys J 90: 1501–1520.

59. Braanigan G, Brown FLH (2007) Contributions of Gaussian curvature and nonconstant lipid volume to protein deformation of lipid bilayers. Biophys J 92: 864–876.

60. West B, Brown FLH, Schmid F (2009) Membrane-protein interactions in a generic coarse-grained model for lipid bilayers. Biophys J 96: 101–115.

61. Watson MC, Penev ES, Welch PM, Brown FLH (2011) Thermal fluctuations in shape, thickness, and molecular orientation in lipid bilayers. J Chem Phys 135: 244703.

62. Mondal S, Khelashvili G, Shan J, Andersen OS, Weinstein H (2011) Quantitative modeling of membrane deformations by multihelical membrane proteins: Application to G-protein coupled receptors. Biophys J 101: 2092–2103.

63. Haselwander CA, Phillips R (2013) Connection between oligomeric state and gating characteristics of mechanosensitive ion channels. PLoS Comput Biol 9: e1003055.

64. Watson MC, Morriss-Andrews A, Welch PM, Brown FLH (2013) Thermal fluctuations in shape, thickness, and molecular orientation in lipid bilayers. II. Finite surface tensions. J Chem Phys 139: 084706.

65. Bibol AF, Constantin D, Fournier JB (2012) Bilayer elasticity at the nanoscale: The need for new terms. PLoS One 7: e46306.

66. Kim T, Lee KL, Morris P, Pastor RW, Andersen OS, et al. (2012) Influence of hydrophobic mismatch on structures and dynamics of gramicidin A and lipid bilayers. Biophys J 102: 1551–1560.

67. Kim CL, Steigmann DJ (2014) Dispersion-induced gradient capillarity in lipid membranes. Cont Mech Thermod: 1–13.

68. Mondal S, Khelashvili G, Weinstein H (2014) Not just an oil slick: How the energetics of protein-membrane interactions impacts the function and organization of transmembrane proteins. Biophys J 106: 2305–2316.

69. Mira K, Ubarretxena-Belanda I, Taguchi T, Warren G, Engelmann DM (2004) Modulation of the bilayer thickness of exocytic pathway membranes by membrane proteins rather than cholesterol. Proc Natl Acad Sci U S A 101: 4003–4008.

70. Boldog T, Hazelbauer GL (2004) Accessibility of introduced cysteines in chemoreceptor transmembrane helices reveals boundaries interior to bracketing charged residues. Prot Sci 13: 1466–1475.

71. Lundback JA, Koegel RE II, Andersen OS (2010) Amphiphile regulation of ion channel function by changes in the bilayer spring constant. Proc Natl Acad Sci U S A 107: 15427–15430.

72. Peruzzo E, Klauda A, Coors DM, Martinac B (2002) Physical principles underlying the transduction of bilayer deformation forces during mechanosensitive channel gating. Nat Struct Biol 9: 606–603.

73. Briegel A, Wong ML, Hodges HL, Oikonomou CM, Piasta KN, et al. (2014) New insights into bacterial chemoreceptor array structure and assembly from electron cryomicroscopy. Biochemistry 53: 1575–1585.

74. Chevite SA, Falke JJ (1996) Molecular mechanism of transmembrane signaling by the aspartate receptor: A model. Proc Natl Acad Sci U S A 93: 2545–2550.

75. Hughson AG, Hazelbauer GL (1996) Detecting the conformational change of transmembrane signaling in a bacterial chemoreceptor by measuring effects on disulfide cross-linking in vivo. Proc Natl Acad Sci U S A 93: 11546–11551.

76. Miller AS, Falke JJ (2004) Side chains at the membrane-water interface modulate the signaling state of a transmembrane receptor. Biochemistry 43: 1765–1770.

77. Ottemann KM, Xiao W, Shin YK, Koshland DE (1999) A piston model for transmembrane signaling of the aspartate receptor. Science 285: 1751–1754.

78. Falke JJ, Ezhre AH (2009) The piston rises again. Structure 17: 1149–1151.

79. Park H, Im W, Seok C (2011) Transmembrane signaling of chemotaxis receptor Tar: Insights from molecular dynamics simulation studies. Biophys J 100: 2955–2963.

80. Huang KC, Mukhopadhyay R, Wingreen NS (2006) A curvature-mediated mechanism for localization of lipids to bacterial poles. PLoS Comput Biol 2: e151.

81. Mukhopadhyay R, Huang KC, Wingreen NS (2008) Lipid localization in bacterial cells through curvature-mediated microphase separation. Biophys J 95: 1034–1049.

82. Rawicz W, Olbrich KC, McIntosh T, Needham D, Evans F (2000) Effect of chain length and unsaturation on elasticity of lipid bilayers. Biophys J 79: 328–339.

83. Weis RM, Hirai T, Chalal A, Kessel M, Peters PJ, et al. (2003) Electron microscopic analysis of bacterial chemoreceptor arrays formed by the bacterial chemotaxis receptor Tar. J Bacteriol 185: 3636–3643.

84. McAndrew RS, Ellis EA, Manson M, Holzenburg A (2004) TEM analysis of chemoreceptor arrays in native membranes of E. coli. Microsc Microanal 10: 416–417.

85. Manson MD (2010) Sensory Mechanisms in Bacteria: Molecular Aspects of Signal Recognition, Norfolk, UK: Caister Academic Press: pp. 107–150.

86. Enges RS (2009) Polar chemoreceptor clustering by coupled trimers of dimers. Biophys J 96: 453–463.

87. Valknin A, Berg HC (2006) Osmotic stress mechanically perturbs chemoreceptors in Escherichia coli. Proc Natl Acad Sci U S A 103: 592–596.

88. Shimizu TS, Tu Y, Berg HC (2010) A modular gradient-sensing network for chemotaxis in Escherichia coli revealed by responses to time-varying stimuli. Mol Sys Biol 6: 1–14.

89. Enges RS, Wingreen NS (2006) Precise adaptation in bacterial chemotaxis through “assistance neighborhoods”. Proc Natl Acad Sci U S A 103: 13040–13044.

90. Wung H, Wingreen NS, Mukhopadhyay R (2008) Self-organized periodicity of protein clusters in growing bacteria. Phys Rev Lett 101: 218101.

91. Briegel A, Ladinsky MS, Oikonomou C, Jones CW, Harris MJ, et al. (2014) Structure of bacterial cytoplasmic chemoreceptor arrays and implications for chemotactic signaling. eLife 3: e02151.

92. Skoge M, Meir Y, Wingreen NS (2011) Dynamics of cooperativity in chemical sensing among cell-surface receptors. Phys Rev Lett 107: 178101.

93. Frank V, Koder M, Furst S, Valknin A (2011) The physical and functional sensitivity of bacterial chemoreceptors. J Mol Biol 411: 534–566.

94. Valknin A, Berg HC (2007) Physical responses of bacterial chemoreceptors. J Mol Biol 366: 1416–1423.

95. Harroun TA, Helfer WT, Weiss TM, Yang L, Huang HW (1999) Theoretical analysis of hydrophobic matching and membrane-mediated interactions in lipid bilayers containing gramicidin. Biophys J 76: 3176–3185.

96. Govil M, Brunsma R, Pincus P (1995) Long-range forces in heterogeneous fluid membranes. EPL 33: 155.

97. Kim KS, Neu JC, Oster GF (1997) Many-body forces between membrane inclusions: A new pattern-formation mechanism. EPL 48: 99–105.

98. Kim KS, Neu J, Oster G (1998) Curvature-mediated interactions between membrane proteins. Biophys J 75: 2274–2291.

99. Kim KS, Neu J, Oster G (2000) Effect of protein shape on multibody interactions between membrane inclusions. Phys Rev E 61: 4201–4205.

100. Chou T, Kim KS, Oster G (2001) Statistical thermodynamics of membrane bending-mediated protein-protein attractions. Biophys J 80: 1075–1087.

101. Kim KS, Chou T, Rusnick J (2008) Degenerate ground-state lattices of membrane inclusions. Phys Rev E 78: 011901.