The Rhodopsin family of G-protein–coupled receptors (GPCRs) comprises the targets of nearly a third of all pharmaceuticals. Despite structural water present in GPCR X-ray structures, the physiological relevance of these solvent molecules to rhodopsin signaling remains unknown. Here, we show experimental results consistent with the idea that rhodopsin activation in lipid membranes is coupled to bulk water movements into the protein. To quantify hydration changes, we measured reversible shifting of the metarhodopsin equilibrium due to osmotic stress using an extensive series of polyethylene glycol (PEG) osmolytes. We discovered clear evidence that light activation entails a large influx of bulk water (~80–100 molecules) into the protein, giving insight into GPCR activation mechanisms. Various size polymer osmolytes directly control rhodopsin activation, in which large solutes are excluded from rhodopsin and dehydrate the protein, favoring the inactive state. In contrast, small osmolytes initially forward shift the activation equilibrium until a quantifiable saturation point is reached, similar to gain-of-function protein mutations. For the limit of increasing osmolyte size, a universal response of rhodopsin to osmotic stress is observed, suggesting it adopts a dynamic, hydrated sponge-like state upon photoactivation. Our results demand a rethinking of the role of water dynamics in modulating various intermediates in the GPCR energy landscape. We propose that besides bound water, an influx of bulk water plays a necessary role in establishing the active GPCR conformation that mediates signaling.

G-protein–coupled receptors (GPCRs) are essential cellular signal transducers involved in the progression of countless human diseases, making them attractive drug targets (1–5). In recent years, considerable attention has been devoted to structural studies of active and inactive conformations of GPCRs, occasionally with binding partners, by X-ray crystallography (6, 7) and cryogenic electron microscopy (cryo-EM) (8, 9). Such studies provide valuable information on key residues and structural motifs involved in several stages of GPCR signaling. At the same time, these structural studies are incomplete, reducing the dynamic conformational ensembles occupied by the various receptor states to a single representative structure selected by the crystallographic or cryogenic conditions (10, 11). However, GPCR activation is thought to be more complex than a binary or bimodal switch model (12), wherein the receptor adopts only on or off conformations modulated by the binding of a ligand (13). Alternative activation mechanisms of GPCRs involve a ternary complex model (14, 15) and an ensemble of states (16, 17) as described by an energy landscape model (ELM) (18–22). In addition, more recent studies provide evidence that the soft matter surrounding these receptors, including water and membrane lipids, plays crucial roles in shaping the energy landscape of GPCR activation (23–28).

Because crystallography and cryo-EM are conducted under dehydrating conditions, and because bulk-phase water molecules are inherently disordered, we cannot rely on these techniques alone to accurately depict hydration states of a protein molecule. As an example, we consider the case of bovine rhodopsin, the archetypal GPCR responsible for dim-light vision which adopts a dynamic equilibrium between two metastable intermediates—the inactive metarhodopsin-I (MI) and active metarhodopsin-II (MII)—in response to light (29). Although crystal structures are available for dark (30) and photoactivated MII (31) states of rhodopsin, both of these show very few internal water molecules. Such structures are insufficient to indicate the true rhodopsin-associated water volume and dynamics. We hypothesized that the metarhodopsin equilibrium following isomerization of the retinal ligand is hydration modulated, e.g., due to large-scale helical movements (29, 32) which are coupled to a flood of bulk water into the protein core (33, 34).

Supporting this hypothesis is recent evidence from small-angle and quasi-elastic neutron scattering studies of rhodopsin (35–37), which describe the MII state as solvent-swollen (35). Additional computational (5, 38, 39) and hydroxyl radical methods (40) have suggested internal water pathways extending...
through the rhodopsin interior. Following Frauenfelder et al. (41), we connected water to a hierarchical energy landscape in which the protein dynamics involve transitions among different tiers or substrates that are coupled (slaved) to the hydration shell or bulk water. To this end, we aimed to monitor changes to the metarhodopsin equilibrium upon selective perturbation of the water corresponding to the substrates of the energy landscape. This strategy can be realized using force-based methods—hydrostatic pressure to probe core shell hydration (42, 43), and osmotic pressure to probe the bulk solvent (44, 45). Yet despite significant interest into the role of bound water in GPCR function (5, 46), there has been little such direct experimental evidence detailing the functional role of bulk water in GPCR activation.

Here, we describe an experimental approach that provides complementary and essential new insights into the surprising role of bulk water in GPCR activation. To observe large-scale hydration-coupling interactions that are typically invisible to structural techniques such as X-ray crystallography (Fig. 1 A and B) or cryo-EM, we used an osmotic stress approach that directly shows a bulk influx of water stabilizing the active rhodopsin conformation (34, 47). A universal osmotic response is demonstrated for a range of excluded osmolytes, revealing that approximately 80–100 water molecules hydrate rhodopsin during MII formation, forcing the intracellular side of the receptor to open for transducin (G) binding (Fig. 1 C and D). We detail how our results may be reconciled with the alternate perspective of crowding and the parallel approach of using hydrostatic pressure. A picture is proposed where rhodopsin becomes solvent-swollen during activation (Fig. 1 E–G) to recruit G-protein binding. Finally, we point out that the activated state of the GPCR is not unlike a dynamic, hydrated sponge with substantial penetration of waters into the protein core.

Results

Biophysical approach to measuring hydration changes in GPCRs. We selected polyethylene glycol (PEG) as the primary osmolyte due to its relative inertness and ability to induce high osmotic pressures. Rhodopsin, the class-A namesake GPCR, provides a convenient case study due to its photoactive nature (SI Appendix, Fig. S1). Its photoactivation is typically portrayed by the following general reaction scheme (29, 48): Rho + hν → MI = MII = MI + H₂O⁺ = MIH₂⁺, that is, photon absorption triggers isomerization of the 11-cis retinal inverse agonist in dark-state rhodopsin (Rho) to all-trans, culminating in formation of the photoactivated intermediates MI and MII (29). Protonation of the conserved E(D)RY motif stabilizes the active MII conformation, to which the G-protein transducin can bind and which is differentiated from inactive MI by the deprotonated retinylidene Schiff base (31).

To directly monitor the fraction of deprotonated Schiff base corresponding to the MII state of rhodopsin in various osmotic conditions (49), we used ultraviolet (UV)—visible spectroscopy of rhodopsin in native retinal disk membranes (RDM) (48). Unlike osmotic pressure which governs movement of water into or out of a cell, we consider here the osmotic stress across a virtual membrane that defines the protein-water interface from which the osmotically active solute is excluded. Our experiment directly assays the difference in associated water volume of the rhodopsin molecule in various conformations versus the bulk solution (44). Quantification of relative MI and MII fractions relied on the technique of basis spectral fitting (48). Using this method, UV-visible difference spectra (light-activated minus dark state) for rhodopsin under MI and MII-favoring conditions were collected as basis spectra (Fig. 2 A and B). Linear combinations were fit to experimental difference spectra to determine the apparent equilibrium constant $K = [\text{MII}]/[\text{MI}]$ and fraction of rhodopsin in the MII state $\theta$ in various equilibrated PEG environments (Fig. 2 C and D and SI Appendix, Fig. S2).

Thermodynamic formulation of osmotic pressure in terms of protein hydration. Osmotic pressure $\Pi$ is expected to behave similarly to hydrostatic pressure $P$ by altering the molar Gibbs free energy $G$ associated with a protein conformation where $(\partial G/\partial \Pi)_T = \bar{V}$. Because the molar hydration volume $\bar{V}$ is itself dependent on osmotic pressure, we may express it as the virial expansion $\bar{V} = (RT/\Pi)(1 + B\Pi + C\Pi^2 + \ldots)$ for constant temperature $T$, where $B$, $C$, etc. are virial coefficients characteristic of a specific hydrated protein state. Upon integrating with respect to $\Pi$ and considering the MI to MII transition, we can derive the standard change in Gibbs free energy perturbed by osmotic pressure, $\Delta G^\circ(\Pi) = \Delta G^\circ + \Pi \Delta \bar{V}^\circ - \frac{1}{2}RT\Pi^2\Delta C + \ldots$, where $\Delta G^\circ$ is the standard free energy change under zero osmotic pressure (see SI Appendix). Notably, the first-order term gives the change in protein
hydration volume $\Delta V^o$ between MI and MII under standard-state (zero osmotic pressure) conditions. We may then relate the change in free energy to the apparent equilibrium constant $K = [\text{MII}]/[\text{MI}]$ by $\Delta G^o = -RT\ln K$, so that the logarithm of the equilibrium constant becomes a second-degree polynomial of $\Pi$:

$$\ln K = \ln K^o - \left(\frac{\Delta V^o}{RT}\right) \Pi + \left(\frac{1}{2} \Delta C\right) \Pi^2, \quad [1]$$

where $K^o$ is the unperturbed equilibrium constant. Experimentally, we observe that large molar mass polymer osmolytes universally back shift the metarhodopsin equilibrium to the MI state, where the linear dependency is proportional to the MII volume change in accordance with Eq. 1. Curvature in the empirical relationship between osmotic pressure and $\ln K$ may indicate a change in compressibility of the hydrated protein volume, or increasing quinary interactions of the polymers with the hydrophilic domains of the receptor during the MI to MII transition (Fig. 2D) (50). Fitting the data with a quadratic function according to Eq. 1 yields the number of hydrating water molecules per mole rhodopsin $N_W$ under standard-state conditions, based on the partial molar volume of water $V_W^p$, and using the relation $\Delta V^o \approx N_W V_W^p$ for the MI–MII transition.

For each large PEG osmolyte with relative molar mass ($M_i$) between 1,000 and 6,000 Da, an increase of ~80–100 hydrating water molecules was calculated for the MI–MII transition from the initial decrease of $\ln K$ with osmotic pressure (Figs. 1C and 2D). This number is much greater than the few structural waters typically seen in protein crystallography (51). Similarly, based on the curvature of the osmotic stress dependencies for each size osmolyte (Figs. 1D and 2D), we calculate virial coefficient changes $\Delta C$ on the order of $0.1 \text{ MPa}^{-2}$, estimated to be equivalent to changes in osmotic compressibility of ~0.01 MPa$^{-1}$. The universal response of the metarhodopsin equilibrium to applied osmotic pressure obtained for various large PEGs (Fig. 3A) strongly implies the effects are colligative and not due to specific interactions with the protein. Although PEG osmolytes also increase sample viscosity, this property of solutions is a descriptor of kinetics, whereas here we consider the thermodynamic equilibrium which appears stable over the course of the experiment. Furthermore, if viscosity were the primary factor at work, we should not expect such drastically different results between small PEGs and large PEGs, which are similarly effective viscosgens. From the osmolyte response, we determine a total number of ~80–100 water molecules entering the rhodopsin interior under standard-state conditions. These results are consistent with Fourier-transform infrared data suggesting increased water accessibility in the MII state (52, 53). Similarly, they correspond to the ~70 water molecules modeled by μ-scale molecular dynamics simulations that flood rhodopsin following induced retinal isomerization from the dark state (38), exhibiting microscopic reversibility (54) with no approximations of additional disorder (55).

**Osmolyte effects on rhodopsin depend on exclusion from the hydrated protein core.** Polymers of differing molar mass face varying degrees of entropic exclusion from a protein cavity. Larger polymers have greater conformational entropy, making their confinement within a tightly closed cavity highly unfavorable in terms of the entropy reduction. The interface between the receptor and surroundings acts as a virtual Gibbs dividing surface analogous to a semipermeable membrane, allowing the movement of water but preventing the penetration of large polymer osmolytes. However, for smaller osmolytes, a different effect is expected. Due to their lower conformational entropy in solution, these osmolytes face a decreased entropic penalty for penetrating the hydrated protein interior. Hence, the Gibbs dividing surface is permeable to small osmolytes, allowing an equal chemical potential of water on both sides of the interface. By consequence, nonsmotic effects should prevail and...
modulate the receptor equilibrium for low concentrations of small PEGs. These trends have been observed previously in analogous experiments with small osmolytes, which give only a partial view of colligative effects on rhodopsin (56). Recent static structures of rhodopsin in complex with transducin or Gγ solved by cryo-EM (8, 57) indicate a Gγ-C-terminal volume of ~1,500 Å³, equivalent to ~5 molecules of PEG 200, while not even a single molecule of PEG 1500 (volume > 2,000 Å³) can be accommodated neglecting entropic con

Experimental, the metarhodopsin equilibrium is found to behave differently in the presence of small PEGs (200–600 Da) as compared to large PEGs. Initial addition of small PEGs shifts ln K linearly to the MII state up to a critical saturation point (Fig. 3B and SI Appendix, Figs. S3–S5). Midsized PEGs achieve this critical value at lower concentrations and a lower maximum MII fraction compared to the smallest-M₃ PEGs. In the higher-concentration regime above this critical value, the metarhodopsin equilibrium more closely resembles the large-M₃ PEGs. Fitting this latter tendency of the osmotic pressure curve as the second component of a piecewise function, we calculated associated thermodynamic values for the MI–MII transition (i.e., the magnitude of the hydration event and change in the system virial coefficient). Initially, small osmolytes penetrate the transducin binding cavity of rhodopsin and stabilize the MII state through nonosmotic, chemical (quinary) interactions. As more osmolytes crowd the transducin binding cleft, they preclude the entry of additional polymer osmolytes as seen for large PEGs. Hence, osmotic effects become the dominant equilibrium-modulating forces beyond receptor saturation. The critical saturation point varies with polymer size, because smaller osmolytes are capable of greater packing inside rhodopsin and reach saturation at higher concentrations.

Our results further show that the apparent thermodynamic parameters of the MI–MII transition have an expected property of osmolyte saturation. With smaller PEGs, a slightly lower hydration volume is measured, combined with a substantially smaller virial coefficient change ΔC. Because small PEG replaces protein-associated water molecules in the low-concentration range, fewer waters are available for withdrawal from the protein in the osmotic regime, requiring higher osmotic pressures to generate similar volume reductions. Consequently, we find a less negative first-order term in the osmotic regime, and thus lower apparent hydration volume for the MI–MII transition for the small PEGs versus the better excluded large PEGs (Fig. 1C). In addition, the stabilizing effect of PEG penetration is observed in the smaller virial coefficient change ΔC for small PEGs: these small osmolytes penetrate and stabilize the protein in the lower-concentration range, decreasing the variability in properties such as osmotic compressibility for the high-concentration regime (Fig. 1D). Strongly interacting PEGs should experience fewer variations in number within rhodopsin compared to water, thereby reducing the stochastic volume fluctuations that may give a higher compressibility, or virial coefficient C, in the MII state.

**Rhodopsin activation due to conserved ERY motif protonation is shifted depending on osmolyte size.** Another striking influence of PEG is its effect on the apparent pKₐ of Glu¹³⁴, following deprotonation of the Schiff base and breakage of its ionic lock with Glu¹¹³ (31, 48, 59). Protonation of Glu¹³⁴ within the conserved ERY motif couples to disrupting another ionic lock involving Arg¹³² and Glu¹⁴⁷, which stabilizes the outward-tilted TM6 conformation of MII, or MIIM⁺ in the protonated case (48). Accordingly, the pH-dependent equilibrium between MI with protonated Schiff base (PSB) and the MII state with a deprotonated Schiff base (SB) is described by an extended titration curve. The apparent pKₐ of this curve is given by Glu¹³⁴ (Fig. 4A) and is thought to be modulated by ionic strength and membrane surface potential (60) plus specific ion binding effects (61). Selectively stabilizing either the closed MI or open MII states impacts this pKₐ as well as the protein environment around the Schiff base (e.g., hydration within the protein). Apart from the pKₐ describing the rhodopsin activation, the pH titration of rhodopsin also demonstrates a nonzero alkaline endpoint at higher temperatures due to thermally activated MII substates (48). In an extended mechanism, the MII state
corresponds to MIIₐ and MIIₐₚ substates, where both have a deprotonated Schiff base and absorb maximally at 380 nm even at high pH (48, 62). Another factor is the additional ρₓ_M under acidic conditions that reflects the protonation of the Schiff base in MIIH⁺ (63). Below this ρₓ_M, the apparent MII fraction determined by UV-visible spectroscopy decreases (Fig. 4A). The phenomenological pH-titration curve thus consists of two ρₓ_M values and an alkaline endpoint, which we model by an extended Henderson-Hasselbalch equation for the MII fraction θ (see SI Appendix for derivation):

$$\theta(\text{pH}) = \frac{\theta_{\text{alk}} + 10^{\rho_{\text{K,Glut}} - \text{pH}}}{1 + (1 + 10^{\rho_{\text{K,Glut}} - \text{pH}})10^{\rho_{\text{K,Glut}} - \text{pH}}}.$$  \[2\]

Here, θ_{alk} designates the alkaline endpoint (apparent MII fraction at high pH), while ρ_{K,Glut} reflects the MIIH⁺ Schiff base protonation equilibrium and ρ_{K,Glut} reflects the Glu¹³⁴ protonation step (Fig. 4A). Titration data for the apparent MII fraction of rhodopsin in solutions of four different large-Mₘ PEGs were used to determine the model parameters (Fig. 4B and SI Appendix, Fig. S6). These results revealed a striking p_{K,Glut} shift from 7.4 to 5.2 in the presence of 50% (wt/wt) large polymer osmolytes.

The pH titration curves for rhodopsin activation are analogous to pharmacological dose-response curves (39, 48) due to changing ligand binding affinity by 2–3 orders of magnitude. This observed p_{K,Glut} shift implies a model in which large osmolytes dehydrate the receptor to shift its equilibrium to the closed MI state, stabilizing the intact ionic lock and favoring a lower fraction of MII. An apparent MII fraction of unity is never achieved for these conditions, due in part to the MIIH⁺ Schiff base protonation region of the titration curve, which begins to overlap with the principal MI–MIIH⁺ region. Strikingly, large-Mₘ PEGs of different molar mass have identical behavior with a nearly equal p_{K,Glut} shift, thus reinforcing the universal osmotic effect by large polymers. Near the alkaline endpoint of the titration, pH values >9.5 induced base-catalyzed retinal hydrolysis following photoactivation, yielding the apoprotein opsin. A quantifiable effect of PEG on the alkaline endpoint was extrapolated where large polymer osmolytes lowered the alkaline endpoint. Our results indicate that large PEGs destabilize the more-open MII state under high-pH conditions where Glu¹³⁴ is fully deprotonated, consistent with the p_{K,Glut} shift that reflects favorability of the closed MI state under osmotic pressure. For small PEGs, the opposite effect is observed, with forward shifting of the metarhodopsin equilibrium by intermediate concentrations of small PEGs (Fig. 4B). With a concentration of 30% (wt/wt) PEG 200, for example, the titration curve incurs a substantial p_{K,Glut} shift from 7.4 to 8.2 opposite to that observed for large PEGs. In total, there is a p_{K,Glut} difference of 3 units between the large and small PEG titration curves, showing the dramatic influence that osmolyte size poses for receptor hydration and proton transfer (64), with consequent activation. This magnitude of p_{K,Glut} shift is analogous to that observed for constitutive mutations of rhodopsin or retinoid antagonists (39, 65).

Discussion

Our findings recast water as an important player in the activation of GPCRs such as rhodopsin (34, 38). Using osmotic stress from large polymer osmolytes (e.g., PEG 1000–6000), we obtain results consistent with a large influx of ~80–100 water molecules into rhodopsin during formation of the active MII state. These direct experimental results are supported by microsecond-scale molecular dynamics (MD) simulations (38, 39), as well as indirect evidence of small-angle neutron scattering (35) and radiolytic protein footprinting showing reorganization of internal waters during rhodopsin activation (40, 46). The osmotic stress methodology primarily investigates the larger volume bulk water associated with the protein, in contrast to experimental techniques which assay structural or bound water molecules (66). While previous hydroxyl radical footprinting experiments did not observe labeling from bulk water, this is possibly because of limited exchange between bound and bulk waters over the experimental timescale (40). In such experiments, tightly bound structural waters are better poised to label internal protein residues upon hydroxyl radical formation. Hence, we view the results as consistent with and complementary to our proposal that various classes of water affect GPCR activation. According to our interpretation,
osmotic stress is a weak force that mainly acts on bulk water, whereas hydration shell or structural water is affected by hydrostatic pressure perturbation or hydroxyl radical labeling.

Although the protein hydration shell typically couples to more localized motions, here we are interested in the global dynamics accompanying GPCR activation (29, 32). Within the unified model of protein dynamics proposed by Frauenfelder et al. (41), these motions are coupled to bulk solvent movements. The molecular picture afforded by MD simulations provides a possible explanation for this functional hydration mechanism, which we refer to as a sponge model. Water penetration into the GPCR has been described as the formation of a continuous internal water channel upon receptor activation (38, 67), whereby bound water movements may be implicated in the GPCR allosteric link between the cytoplasmic face and ligand binding pocket (46, 68). 

In the limit of increasing osmolyte size, GPCR activation undergoes a bulk water inclusions effect, so that osmotic dehydration by large, excluded polymers inhibits GPCR activation. Small osmolytes are more localized motions, here we are interested in the global dynamics, modulating rhodopsin activation. They penetrate inside the receptor and stabilize the active (expanded) receptor state through interactions with the transducin binding cleft until reaching a quantifiable saturation point. In the limit of increasing osmolyte size, GPCR activation follows a universal dependence on osmotic stress, similar to that observed for ion channel opening (44).

Integration of osmotic pressure and hydrostatic pressure effects on metarhodopsin equilibrium. We next show how further insights into GPCR activation are obtained by integrating osmotic pressure data with hydrostatic pressure effects. Both hydrostatic pressure and osmotic pressure backshift the metarhodopsin equilibrium to the preactive MI state, but for different reasons (71, 72). Because hydrostatic pressure entails a thermodynamically closed system, the perturbation detects the change in molar protein volume (i.e., density) of the system, but not changes in water molecule number. For rhodopsin, these changes in density can involve the collapsing or penetration of water into small protein cavities or voids, or alternatively a higher-density solvation shell versus the bulk (73). By contrast, osmotic pressure involves an open system, which is separated from the surroundings by a semipermeable Gibbs dividing surface (44). Hence, osmotic pressure involves changes to protein hydration, so that both methods are complementary to one another (74). By analogy to pressure effects on protein folding reactions (42, 75), MI can be seen as more tightly packed, which minimizes void volume or forces small amounts of solvent into internal cavities (42, 75–78). Conversely, in MII there may be an increase in number of water molecules, giving a solvent-swollen state (35, 37). While the active MII state is less dense (71, 72), it is also more hydrated than its precursor MI state, with greater penetration of water molecules into the protein (38, 54). The MII state is less tightly packed in its formation of a large internal cavity for G-protein binding, which in tandem with the high-density hydration shell accommodates the larger hydration volume. Water molecules previously forced into restricted MI cavities may find greater entropic stabilization in the larger, nascent hydration cavity of MII. The two force-based methods can thus be unified to give a more comprehensive picture of the rhodopsin changes in a hydrated lipid membrane upon light activation (39).

Osmotic pressure informs volumetric fluctuations and cosolute quinary interactions in the active GPCR state. Additional information regarding GPCR activation is obtained from the dependence of the active MII fraction on osmotic pressure. Modeled as changes in the virial coefficient \( \Delta C \), the curvature term may result from an apparent change in compressibility of the hydrated protein (see SI Appendix), related to increasing volume fluctuations in the dynamic, hydrated conformation of the active receptor (37). Analogous compressibility-governed behavior of enzyme-catalyzed equilibria has been found in the case of large hydrostatic pressures (43). For rhodopsin under osmotic stress, we could gather from the sign of the second-order term an increase in osmotic compressibility, \( \kappa_{ii} \equiv -\left( \partial \ln V / \partial \Pi \right)_{\tau} = -(1/V)(\partial V / \partial \Pi)_{\tau} \), which is related to fluctuations of the hydration volume \( V \) of the protein system (79). Hence, a greater osmotic compressibility in the solvent-swollen active MI state indicates greater volumetric fluctuations, consistent with a hydrated sponge-like state having variations in how water packs within the protein. Greater fluctuations in the active-state GPCR would indicate a greater number of available microstates for the entropy-stabilized active receptor, reflecting similar observations by quasi-elastic neutron scattering (37).

Alternatively, the change in virial coefficient may be explained by competing quinary interactions of the large polymers with the active MII state (80). Similar to the interactions with low concentrations of small PEGs that drive the equilibrium forward to MII (39, 47), large PEGs may also experience chemical, or quinary interactions with the protein, albeit to a lesser extent (80). As the concentration of polymer osmolytes in the surrounding bulk solvent nears its saturating limit, the entropic penalty for osmolyte penetration and interaction with the receptor is reduced, yielding polymer stabilization of the active GPCR that cancels the osmotic effect. A combination of both specific quinary interactions and compressibility changes between the two states may likely be involved in the \( \Delta C \) apparent curvature term. As a second-order approximation, our model of the deviation from linearity may exclude higher-order terms describing either the changes in compressibility or quinary interactions with the polymer osmolyte (see SI Appendix). Although not trivial to mathematically separate these terms, both types of contributions qualitatively confirm the essential behavior expected of a dynamic and solvent-swollen active receptor conformation.

Osmotic stress effects are a direct assay of hydration changes and are compatible with cosolute crowding models. Thus far, we have considered the introduction of polymer
cosolutes purely from the viewpoint of osmotic stress, which is a consequence of the Gibbs-Duhem equation for systems in equilibrium and directly assesses hydration states (45). Naturally arising osmotic effects of cosolutes on macromolecules can also be recovered from popular alternative perspectives such as crowding. In this framework, large cosolutes exert steric forces on macromolecules to restrict their volume occupancy (81, 82). Except to exclusion from the macromolecular environment, these particles create depletion forces that attract multiple proteins or structural motifs together (83, 84), which are themselves a consequence of osmotic stress (84). Steric interactions that confine volume in crowded environments have a first-order effect on the molar free energy of the macromolecule according to \( dG = RTV dM \), where \( M \) denotes the molar concentration of perfectly excluded polymer solutes and \( V \) is the molar hydration volume as before (82). Using the van’t Hoff law of ideal solutions \( \Pi = MRT \), a first-order approximation for osmotic pressures, we can recast this expression to the equivalent osmotic stress relation, \( (\partial G/\partial M)_T = \Pi \). Thus, crowding and osmotic stress effects are mathematically equivalent to first order. It is likely that higher-order terms in either picture can be reconciled, although computationally challenging to demonstrate due to significant overlap of the two approaches (45). While either framework may be applicable to a particular situation, we show here the utility of the osmotic stress approach toward directly measuring hydration changes of rhodopsin as an archetypal GPCR.

Evidence for a hydration-modulated energy landscape model of GPCR activation. Our discoveries enlarge the scope of the role that soft matter plays in key biological phenomena. Already having detailed the influences of membrane curvature stress in GPCR function (25, 28), we now describe an equally powerful biological influence in the form of the bulk solvent (85). It is important to appreciate the fundamental distinction between these two: the osmotic stress effect must be a result of protein dehydration apart from osmotic dehydration of the lipid bilayer. Membrane dehydration alone would produce an effect opposite to that observed in this study, by decreasing the area per lipid and increasing the membrane thickness, which are known to favor active MII (23, 28, 63). Chemical interactions of PEG with lipids are thought to be relatively weak or nonspecific (86) and they promote negative spontaneous curvature of lipid monolayers (87), thereby also tending to favor MII unlike the current results. Consequently, both effects must be weaker than the osmotic stress effect on the protein itself that we observe experimentally. Our findings offer insights into howGPCRs, like rhodopsin, produce their high-fidelity rapid signal amplification. To achieve this biological outcome, there must be ways to allosterically modulate the binding affinity of the G-protein at different stages of its cycle (88–90). For rhodopsin-based signaling, the occurrence of a hydrated, sponge-like active state is consistent with an energy landscape mechanism (ELM) for receptor activation. The thermodynamic parameters are in accord with a partial unfolding of the receptor protein in analogy to hydrostatic-pressure effects on globular proteins. None of these conclusions can be deduced from X-ray crystal structures or cryo-EM structures alone, thus requiring physicochemical studies such as force-based methods as described herein. The hydration-activation mechanism has natural implications in GPCR pharmacology and drug design.

To summarize, we have experimentally shown the existence and functional role of a large hydration event coupled to activation of the GPCR rhodopsin. Accordingly, we propose a hydration-mediated sponge-like mechanism with potential implications for biological signal amplification. The GPCR activation is but one event in a sequence of signal transduction, the next being the binding of the G-protein and receptor-catalyzed exchange of GTP for GDP. Interestingly, hydration plays a key role in lipid bilayer dynamics that could possibly also have crucial implications for GPCRs like rhodopsin. Cooperativity of membrane lipids and hydration can potentially affect the energy landscape of GPCR activation. Further studies of hydration effects on GTP-GDP exchange may present important clues as to the complete mechanism of receptor-catalyzed G-protein binding and release. Whether the water interacts synergistically with lipids to affect receptor function, and whether it regulates the binding and unbinding of the G-protein transducin, remain as intriguing questions.

Materials and Methods

Retinal disk membrane purification. Retinal disk membranes (RDM) were isolated from bovine retinas and characterized by UV-visible spectroscopy as previously described (SI Appendix, Fig. S1) (91). The RDM were suspended with concentrations of 10 \( \mu \text{M} \) rhodopsin in 67 mM bis-Tris propane (BTP) buffer with 130 mM NaCl and 2 mM MgCl\(_2\) at pH 7.5. The osmolar contributions of the buffer and salts were assumed to be negligible in comparison to the PEG osmotic pressure. PEG was added in 0–60 wt% concentrations and lightly sonicated with the RDM for equilibration (SI Appendix, Fig. S2).

UV-visible spectroscopy and data analysis. Absorption spectra for suspensions of RDM containing 10 \( \mu \text{M} \) of rhodopsin plus the osmolyte were recorded with a Varian Cary-50 UV-visible spectrophotometer (Palo Alto, CA). Throughout the experiment, the temperature was kept constant at 15° C. A 1-mm pathlength cuvette placed close to the detector was used to minimize light scattering. The UV-visible absorption spectra were collected from 270 to 650 nm for both the dark-state protein and rhodopsin exposed to a green LED (528 nm) actinic light source for 7 s. A scattering correction was performed using the equation \( \Delta A_{\text{corr}} = \Delta A - a/\lambda^2 + b \) to account for scattering changes after bleaching. Here, \( \Delta A_{\text{corr}} \) is the corrected amplitude of the difference spectrum, \( \Delta A \) is the experimental difference amplitude, \( a \) is the fitting constant, \( \lambda \) is the wavelength, and \( b \) is an additive constant to correct for baseline shifts (92). The fraction of photoprotected protein in the MII state (\( \theta \)) was calculated by fitting the experimental difference spectrum by a linear combination of MII and MII basis spectra using least-squares regression. Here, the coefficients \( c_1 \) and \( c_2 \) of each basis vector component yield the relative concentrations of either species. The MII basis spectrum was recorded for RDM suspended in pH 9.2 buffer at 5° C, while the MII basis spectrum was recorded for RDM suspended in pH 5.0 buffer at 15° C. Spectral lineshapes of the basis spectra were confirmed mathematically through singular value decomposition of pH titration spectra in multiple osmotic environments.

Calculation of number of water molecules. Osmotic pressure values were calculated from universal fitting curves to model the experimental osmotic pressure data (93) as reported in Cohen et al. (49). The number of waters involved in the rhodopsin activation process was calculated from the dependency of \( \ln K \) on osmotic pressure (44). A second-order polynomial (Eq. 1) was fit to the osmotic regimes of the data by a nonlinear least-squares algorithm using MATLAB R2018a (for small PEGs, including those curves in the decreasing regions above the identified saturation point). A similar approach was used in fitting the extended Henderson-Hasselbalch curve (Eq. 2) describing the pH dependence of the rhodopsin activation.

Data Availability. All study data are included in the article and/or SI Appendix. Experimental data can be accessed at the University of Arizona Research Data Repository (DOI: 10.25422/au.data.19686555).

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