Energy Landscapes as a Tool to Integrate GPCR Structure, Dynamics, and Function

G protein-coupled receptors (GPCRs) are versatile signaling molecules that mediate the majority of physiological responses to hormones and neurotransmitters. Recent high-resolution structural insights into GPCR structure and dynamics are beginning to shed light on the molecular basis of this versatility. We use energy landscapes to conceptualize the link between structure and function.

G protein-coupled receptors (GPCRs) evolved to be the principal line of communication between cells and tissues in higher organisms. Moreover, they contribute to communication between individuals through the senses of vision and olfaction. These common portals for information transfer have also been hijacked by HIV and other viruses for the purpose of cellular infection. This extraordinary functional versatility is accomplished through a common seven transmembrane structure, and, consequently, GPCRs are also called seven transmembrane receptors and heptahelical receptors.

As a result of their broad influence over human physiology and behavior, GPCRs are promising candidates for the development of new and more effective small molecule therapeutics. However, the development of selective GPCR drugs is challenging for several reasons. First, there is a high degree of homology among many closely related receptor subtypes that can regulate diverse physiological functions. Also, a single GPCR may couple to more than one G protein, signal through G protein-independent pathways, undergo complex regulatory processes, and be allosterically regulated by small molecules and other proteins, including other GPCRs. Moreover, the predominant signaling behavior of a GPCR may differ for different cells or organs. Finally, drugs may preferentially activate or inhibit specific signaling pathways. Although this functional versatility is important for normal physiological signaling, it makes identifying effective therapeutics very challenging.

The goal of this review is to interpret the functional versatility of GPCRs in terms of their structural plasticity and to show how these concepts can be visualized as energy landscapes of individual GPCR protein molecules. Much of what is known about GPCR structure and mechanism of activation comes from work on a relatively few GPCRs including rhodopsin and several monoamine receptors. Although the review will focus on these proteins, it is likely that the concepts will apply to most if not all members of this extensive family.

Evidence for Structural Plasticity

As noted above, individual GPCRs may mediate a spectrum of signaling and regulatory behavior that can be modulated in a ligand-specific manner (23). As an example, FIGURE 1 illustrates the diverse signaling behavior observed for the human $\beta_2$-adrenergic receptor ($\beta_2$AR), a close relative of rhodopsin. The $\beta_2$AR was one of the first GPCR signaling pathways to be characterized, and the first hormone-activated GPCR to be cloned (13) and have its structure determined by crystallography (8, 37, 38). Early studies revealed the predominant signaling pathway to be from agonist-activated $\beta_2$AR to Gs, the stimulatory G protein for adenylyl cyclase. However, it is now known that the $\beta_2$AR couples to the G protein Gi (10, 12), as well as activating G protein-independent pathways through arrestin (2, 41) and possibly other cellular signaling proteins. The activity of the $\beta_2$AR can be modulated in different ways by small molecule ligands. The term efficacy is used to describe the effect of a ligand on the signaling behavior of a receptor (FIGURE 1, INSET). Agonists fully activate the receptor; partial agonists produce a submaximal activity. Antagonists block the binding of other drugs, without altering the basal activity of the receptor. Like many GPCRs, the $\beta_2$AR exhibits agonist-independent activity that can be suppressed by drugs classified as inverse agonists (FIGURE 1, INSET). Further complicating the classification of drugs for a given GPCR is the observation that the efficacy of a panel of drugs may differ for different downstream signaling profiles (23, 24, 48).

The remarkable versatility of $\beta_2$AR signaling is not limited to this particular GPCR but is more representative of the larger family. The most notable exception is the namesake for Family A GPCRs, rhodopsin. In contrast to most GPCRs, rhodopsin
exhibits a more restricted signaling behavior necessary for its highly specific role, the efficient detection of light. Nevertheless, even rhodopsin exhibits dynamic properties and different functional and structural states (21).

This functional versatility of GPCRs cannot be explained by a simple on-off switch model of receptor activation, and is more compatible with dynamic and adaptable structures. In fact, there is a growing body of biochemical and biophysical data showing that GPCRs are flexible, dynamic proteins and that functionally distinct ligands can stabilize specific receptor conformations (25). In the next section, we will discuss these conformations in terms of their thermodynamic energy profile in the context of energy landscapes and consider how physiologically relevant variables can change this energy landscape and thereby the signaling behavior of the receptor.

Energy Landscapes as a Tool to Visualize Protein Activation

Protein structures elucidated through X-ray crystallography reveal a highly specific assembly of smaller domains (alpha helices, beta sheets, connecting loops) that may give the impression that proteins are rigid entities. On the contrary, proteins are dynamic and exhibit small-scale movements at the level of amino acid side chains and larger scale movements between domains, on time scales ranging from nanoseconds to seconds (20). As a result, proteins exist in an ensemble of conformations, each with a distinct energy resulting from intramolecular chemical bonds and ionic interactions. Conformations of lower energy (i.e., more stable) are more populated, following a Boltzmann distribution. In addition, the probability associated with the occupancy of each conformation...
Conformational entropy also plays a role in how the conformational states are populated. Overall, the thermodynamically most favorable conformations (as a balance of internal energy and conformational entropy) predominate in the ensemble. These particularly stable structures correspond to those conformations likely to be captured in X-ray crystallography. Thus our view of protein structure is biased toward these stable conformations. However, there is growing experimental evidence, primarily by NMR, suggesting that conformational states involved in activation occur at equilibrium even in the absence of agonist, yet they represent only a small fraction of the ensemble (19). Thus the less stable (i.e., less populated) conformations are also important for function.

Although the framework to relate structural flexibility and protein function has been developed primarily for enzymes, these concepts also can be applied to GPCRs. These proteins are remarkably versatile signaling molecules due to their flexible and dynamic three-dimensional structure (25). However, although the amount of structural information is increasing rapidly, we still do not understand how ligand binding is translated into the series of conformational changes that lead to activation. As a first approximation, ligand-receptor interplay can be visualized by representing the energy of the receptor conformations along the activation pathway. In their simplest form, these energy landscapes are two-dimensional curves (FIGURE 2). The middle panel shows a very simple
situation of a receptor that, in the absence of ligand, exists in two conformations. The energy minima correspond to (meta)stable conformations, inactive (R) and active (R*), connected through a saddle point of higher energy that represents a particular conformational change. The relative population of the states and the rate of transition between them are determined, respectively, by their difference in energy and by the height of the barrier. In this example, the lower energy of R results in a higher population (FIGURE 2, MIDDLE RIGHT, where size is proportional to population). In addition, in this case, receptors that achieve the state R* (responsible for constitutive activity) revert back rapidly to the state R, due to the low energy barrier of the R* → R transition.

These simplified energy landscapes can be used to illustrate the difference between receptor activation by an induced-fit mechanism or by conformational selection. In an induced-fit scenario (FIGURE 2, BOTTOM), the ligand binds to the conformation R to create the R-L complex, which possesses a different energy landscape (red line) than the "empty" receptor (broken line). In this case, ligand binding accelerates activation by lowering the energy of the conformational transition to R*-L. In addition, ligand-receptor interactions stabilize the complex, resulting in a lower conformational energy of R*-L. Ligand binding also provides the energy to induce the conformational changes that allow crossing the energy barrier to R* (black arrows). These effects are ultimately translated in an increase of the population of the R*-L conformation (FIGURE 2, BOTTOM RIGHT). On the other hand, in conformational selection (FIGURE 2, TOP), the ligand binds to and stabilizes (i.e., lowers the energy) the active conformation R*, resulting in a change of the energy landscape (blue line) for the R*-L complex. In the new landscape, the transition to the inactive form R-L becomes unfavorable due to the higher energy of the R-L complex and a higher energy barrier for this transition. Thus alteration of the shape of the energy landscape results in a population shift of the receptor conformations toward the R*-L active state (FIGURE 2, TOP RIGHT).

Once an equilibrium state has been reached between ligand, receptor, and the complex, activation by induced-fit or by conformational selection may result in completely indistinguishable states. The fundamental difference between these mechanisms resides in the first stages of ligand-receptor interaction and activation. The great structural diversity of the natural GPCR ligands, ranging from ions and small organic molecules to peptides and glycoprotein hormones, suggests important differences in those first stages, which may be related, to some extent, to different ligands activating the receptor either by induced-fit or conformational selection. For instance, although rhodopsin (6) and the angiotensin AT1 receptor (22) appear to operate primarily by an induced-fit mechanism, β2AR ligands may function by selecting specific receptor substates (25).

In the following sections, we discuss the differences between these two models of activation in terms of energy landscapes by using rhodopsin and the β2AR as model systems.

**Energy Landscapes of Rhodopsin Activation**

Most of our knowledge of GPCR activation has originated from rhodopsin. A variety of techniques, particularly spectroscopic methods, have revealed several metastable intermediates in the rhodopsin activation pathway (21). In addition, the two-dimensional energy landscape of this mechanism has been delineated by calorimetric and reaction kinetic experiments (9, 42). Rhodopsin activation implies a linear pathway along this energy landscape, which is schematically depicted in FIGURE 3, TOP. Inactive (dark) rhodopsin is trapped in a deep energy well (FIGURE 3A, black line). The stability of this conformation, which cannot overcome the energy barrier (i.e., change conformation) to populate active states, results in the absence of constitutive activity of rhodopsin. Retinal isomerization on light absorption transforms the 11-cis retinal inverse agonist into the all-trans retinal agonist, generating a series of ligand-receptor steric clashes that increase the energy of the system and change the shape of the energy landscape (FIGURE 3A, red line). As the receptor structure relaxes, the intermediate active states (Batho, Lumi, and Meta I) become energetically available and get populated. The crystal structures of Batho (32) and Lumi (33) show only small local structural changes in the binding pocket, and the electron cryomicroscopy density map of Meta I (39) reveals only a modest rearrangement in TM6 at the level of the binding site. Also, infrared spectroscopy on azido-labeled rhodopsin suggests a small rotation of TM6 and movement of the cytoplasmic side of TM5 at the Meta I stage (53). The ionic interaction between Arg135(3.50) and Glu247(6.30) [the Ballesteros-Weinstein general number (4) of residues is given in parenthesis], which has been suggested to stabilize the inactive conformation of GPCRs (3), is, in fact, relatively weak (49) and may become partially opened already in Meta I (53). These early intermediates are formed in microseconds due to the low energy barriers between them, and the reaction displaces almost completely to the lower energy Meta I conformation. Subsequent transition to the Meta II states constitutes a rate-limiting step due to a
higher energy barrier. Activation proceeds by sequential formation of the Meta IIa, Meta IIb (by rigid-body movement of TM6), and Meta IIbH+ [by neutralization of Glu134(3.49) in the ERY motif of TM3] intermediates. These Meta species exist in a G protein-dependent equilibrium, reached after a few milliseconds. G protein binding to Meta IIb/ Meta IIbH+ lowers the energy of the ternary complex (FIGURE 3B, blue line) and displaces this equilibrium toward these forms (see Ref. 21 for a comprehensive review on this process). In summary, retinal isomerization provides the energy for the receptor to “jump” the high initial activation barrier and proceed through activation along the energy landscape through small structural changes to the Meta I state. Subsequently, larger-scale conformational rearrangements and changes in prototiation states lead to formation of an equilibrium between the Meta forms. Finally, G protein binding further changes the energy landscape, displacing the equilibrium to the active form of the receptor, capable of catalyzing the GDP-GTP exchange in the G protein. Other works (5) show that there are additional rhodopsin conformations (as Meta III) that create “bifurcations” in the activation pathway. The graphical representation of such a process as an energy landscape would require more than two dimensions (see below).

**FIGURE 3.** Energy landscapes of rhodopsin and β2AR activation

A: inactive (dark) rhodopsin is trapped in a deep energy well (black line). Retinal isomerization provides the energy for the receptor to overcome the high initial activation barrier and proceed along the energy landscape through small conformational changes (Batho and Lumi) to the Meta I state (MI). Subsequently, activation proceeds by formation of the Meta IIa (MIIa), Meta IIb (MIIb; by rigid-body movement of TM6), and Meta IIbH+ (MIIbH+; by neutralization of Glu134(3.49) in the ERY motif of TM3) intermediates, and establishment of an equilibrium between the Meta forms.

B: G protein binding further changes the energy landscape (blue line), displacing the equilibrium to the active ternary complex, capable of catalyzing the GDP-GTP exchange in the G protein.

C: the β2AR possesses a shallow energy landscape with several conformational states (R, R’, R’’; that differ in small structural changes in TM5 and TM7) separated by relatively low energy barriers (blue line). This translates to an inherent flexibility that allows the ligand-free receptor to explore different conformations. Ligand binding to certain intermediates (R’’ in this example) changes the shape of the energy landscape (green line), and activation proceeds to populate conformations of lower energy (R’’’ and R*). These conformations probably involve a similar set of conformational changes as rhodopsin, i.e., rearrangement of TM6 and neutralization of Asp130(3.49) in the DRY motif of TM3.

D: binding of the G protein to these latter states further changes the energy landscape (blue line), lowering the energy and stabilizing the active ternary complex.
According to this view, rhodopsin activation represents an extreme case of induced-fit: the ligand is covalently bound in an occluded binding site as an inverse agonist that is transformed into a covalently bound agonist on absorption of a photon. The resulting ligand-receptor clashes and subsequent small-scale rearrangements in the receptor backbone and side chains to optimize the agonist-bound form end up stabilizing larger conformational changes, changes in protonation states, and, ultimately, changes in the stabilization of an active state.

### Energy Landscapes of β_{2}AR Activation

There is substantial evidence for a common mechanism of activation between rhodopsin and the rest of class A GPCRs. Members of this large and diverse family activate one or more members of the relatively small family of highly homologous G proteins. Accordingly, GPCRs display a higher sequence similarity in the cytoplasmic side of the transmembrane bundle (31), which contains the residues involved in receptor activation and in G protein coupling (28). A diversity of biophysical and biochemical studies in different families of GPCRs (see Ref. 11 for a review) provides direct evidence of similar conformational changes underlying GPCR activation. Thus the considerable amount of information about rhodopsin activation, and its representation in terms of energy landscapes, can serve as a framework to discuss activation of other GPCRs using the β_{2}AR as a model system.

Unlike rhodopsin, the β_{2}AR is not trapped in a fully inactive conformation in the absence of agonist but possesses an inherent conformational flexibility that allows the ligand-free receptor to explore different conformations (17, 36). This suggests a shallow energy landscape, with several conformational states separated by relatively low energy barriers (FIGURE 3C, blue line). On the other hand, the remarkable diversity of ligands and the low sequence conservation in the ligand-binding domains indicate the existence of profound differences in the process of ligand binding and ligand-receptor interactions among GPCRs. For a rhodopsin-like activation by induced fit, the agonist would have to bind to the receptor with high affinity and use this high binding energy to initiate conformational changes (i.e., “jump” the first energy barrier, as in retinal isomerization). However, the relatively low affinity and rapid dissociation rates observed for β_{2}AR agonists (16) are not consistent with induced fit and suggest conformational selection. In this scenario, an agonist would bind to a similar extent diverse conformations in the receptor ensemble. Most of the binding events are transient, and the ligand can escape back to the solvent. However, when the agonist binds to certain conformational intermediates (R” in FIGURE 3C), ligand-receptor interactions are more favorable and the complex is stabilized (i.e., lowers its energy). In this case, the binding event changes the shape of the energy landscape (green line), and the ligand-receptor complex “proceeds forward” and populates conformations of lower energy (R^{*} and R^{*}_{2}). Similarly to rhodopsin, binding of the G protein to these states further changes the energy landscape (FIGURE 3D), stabilizing (i.e., lowering the energy) of the ternary complex, which becomes the predominant conformation. The latter states in β_{2}AR activation, which would be analogous to the Meta II states in rhodopsin activation, involve a similar set of conformational changes, i.e., rearrangement of TM6 and neutralization of Asp130(3.49) in the DRY motif of TM3 (11). In this case, the ionic interaction between TM3 and TM6, mediated by Arg131(3.50) and Glu268(6.30), is even weaker than in rhodopsin. The crystal structures of β_{1}AR (50) and β_{2}AR (38) show that this interaction is labile even in inactive states. In addition, molecular dynamics studies on the β_{2}AR (14) suggest that this interaction exists in equilibrium between formed and broken states.

According to this view of activation, partial agonism can be visualized in different ways. On one hand, it has been shown that the duration of the binding event is correlated to agonist efficacy, i.e., partial agonists dissociate faster than full agonists, and, as a result, not all binding events last long enough to promote activation of the G protein cycle (46). Another non-excluding possibility is that partial agonists stabilize different intermediate conformations (17, 45, 52) that lead to alternate activation pathways and to non-optimal G protein activation (see below).

### Extension to 3D Energy Landscapes

Two-dimensional energy landscapes, where energy is represented as a function of a single reaction coordinate (FIGURES 2 AND 3), are relatively easy to interpret. However, they are not suitable to consider complex situations, such as receptors with rich pharmacological properties (synthetic inverse agonists, antagonists, partial agonists and superagonists, in addition to natural full agonists), or dissection of the reaction pathway in discrete nonsequential conformational changes. In these cases, consideration of an additional reaction coordinate, i.e., the extension to three-dimensional (3D) energy landscapes, provides an additional degree of freedom that allows a more detailed study of the activation process. Specifically, the
reaction coordinate (x-axis in FIGURES 2 AND 3) can be decomposed into two sets of conformational changes (e.g., activation pathways) that, in turn, can be used to visualize alternate routes of activation along the energy landscape. These sets of conformational changes can be extracted from the activation pathways depicted in FIGURE 4 that aggregate a series of smaller local conformational changes into specific activation pathways. For instance, in the 3D energy landscapes depicted in FIGURE 5, we represent energy in the z-axis (color) as a function of the conformational changes in TM7 and TM6 (in the x-axis), and of the conformational changes in TM5 and neutralization of Asp130(3.49) in the DRY motif of TM3 (in the y-axis). Activation follows the path traced by the lines in the 3D landscapes. Slicing the 3D surface along these paths results into the simpler 2D landscapes depicted in the right panels.

Conformational changes of TM6 on activation of different receptors have been measured from EPR and fluorescence spectroscopy. Also, structural changes in TM5 are evident in the opsin, adrenergic, and adenosine structures. However, the basis and extent of the structural changes in TM7 on activation have not been studied in detail yet. In rhodopsin, residues K296(7.43)-A299(7.46) form a short 3_10 helical segment that results in a strong local tightening of the helix, maintained by a complex inter-helical hydrogen bond network that involves water molecules (26). A detailed analysis of the crystal structures of opsin reveals that the tightening in TM7 due to the 3_10 segment is significantly reduced. Although only five water molecules

**FIGURE 4. Pathways in the activation mechanism of β_2AR**

This figure represents epinephrine (shown as spheres) bound to the β_2AR (TM5, green; TM6, blue; TM7/helix 8, cyan). Receptor activation comprises a series of conformational changes that aggregate into specific activation pathways. Agonists interact with a group of Ser residues on the extracellular side of TM5 [Ser203(5.42), Ser204(5.43), and Ser207(5.46)] involved in ligand binding and receptor activation (44, 45). This interaction disrupts the TM3/TM5 interface at the binding site (35), and these local changes are transmitted to the cytoplasmic side of the helix through the rearrangement of the Pro-induced distortion of TM5 (green arrow) (Sansuk et al., unpublished observations; Ref. 47). Agonists also induce/stabilize the rigid body movement of TM6 (blue arrow) through their action on Trp(6.48). A complex hydrogen bond network between highly conserved residues and a cluster of water molecules in the TM2-TM6-TM7 interface seems responsible for stabilizing the 3_10-helix segment observed in the inactive state of rhodopsin (26). This network is most likely conserved in other Class A GPCRs (1) and has a structural and functional role (34). Specifically, we suggest that agonist binding induces changes in this network, which results in local structural changes in the NPxxY motif and in a relocation of the cytoplasmic side of TM7 (cyan arrow). These pathways involve highly conserved residues and are probably conserved in other Family A GPCRs. This figure has been created with PyMOL (DeLano, WL. The PyMOL Molecular Graphics System (2002) at http://www.pymol.org).
per monomer were modeled in the refinement of the opsin structure due to weak electron density, and none of them in the TM2-TM3-TM6-TM7 pocket, the partial recovery of the local alpha-helical structure must result in the displacement of some of the waters observed in the structure of rhodopsin. These local changes ultimately result in the partial relocation of the side chain of Y306(7.53), part of the NPxxY motif, which gets closer to center of the transmembrane bundle. If the structures of dark rhodopsin and opsin are used to model the beginning and end of the activation process, we can hypothesize that plasticity in the water cluster around the NPxxY motif of TM7 results in conformational changes in the cytoplasmic side of this helix, which represents an additional pathway in rhodopsin activation (FIGURE 4, cyan arrow). Interestingly, in the currently available non-rhodopsin structures, which correspond to inactive states, the cytoplasmic sides of TM7 resemble the structure of opsin, resulting in Tyr(7.53) facing the protein core. Thus the corresponding region in rhodopsin (with Tyr306(7.53) engaged in an aromatic-aromatic interaction with Phe313 in helix 8) may be partially responsible for locking rhodopsin in a totally inactive state that cannot proceed further in the energy landscape to populate meta-activating conformations.

According to this convention, FIGURE 5 depicts a number of theoretical 3D energy landscapes for the activation of \(\beta_2\text{AR} \). FIGURE 5A shows the energy landscapes of the receptor in the absence of ligand. The receptor population fluctuates between the low energy conformations \(R\), \(R'\), and \(R''\), separated by low energy barriers. In terms of structure, conformational changes in the TM3/TM5 interface and plasticity in the water cluster around the NPxxY motif result in conformational changes in the cytoplasmic sides of TM5 and TM7 that stabilize different conformations. If the energy barrier that leads to \(R''\) (related to movement of TM6) is low enough, the conformation \(R''\) is kinetically available and can get populated. The “open” cytoplasmic side facilitates neutralization of Asp130(3.49) in the DRY motif of TM3, which stabilizes the \(R^*\) conformation, and is able to bind to and activate the G protein. In this case, the receptor will show constitutive activity. FIGURE 5B depicts the effect of an inverse agonist on the receptor energy landscape. The crystal structure of the \(\beta_2\text{AR} \) bound to carazolol (38) shows that this ligand stabilizes a conformation where TM5 and TM7 have rearranged compared with the crystal structure of rhodopsin, which we use as a reference of a GPCR “locked” in a purely inactive state. In our theoretical energy landscape, carazolol would bind (i.e., select) and stabilize conformation \(R''\) and change the overall energy landscape of the receptor (compare FIGURE 5A AND 5B). As a result, the energy barriers to escape this conformation are higher, and the receptor gets trapped. This is a way to visualize the stabilizing effect of inverse agonists in receptor conformation. In this particular case, ligand binding precludes the movement of TM6, which is required to reach the active states. FIGURE 5C shows the possible effect of a full agonist on the \(\beta_2\text{AR} \) energy landscape. An agonist can also bind to and select the \(R''\) conformation. However, the optimized agonist-receptor interactions change the energy landscape in such a way that the barrier to reach \(R'''\) (i.e., to move TM6) is lowered. The stabilized open crevice in the cytoplasmic side favors the neutralization of Asp130(3.49) in the DRY motif so Arg131(3.50) can reach the extended conformation needed for binding of the G protein, as seen in the structure of activated opsin (40). FIGURE 5D shows a possible mechanism for partial agonism in terms of energy landscapes. A partial agonist can also bind to and stabilize \(R''\) to induce a change in the energy landscape. In this case, a conformation \(R'''\) different from the \(R''\) conformation stabilized by agonists, becomes available. In terms of structure, this could represent, for instance, a slightly different relocation of TM6. In this new energy landscape (compare FIGURE 5C AND 5D), the lower energy conformation corresponds to \(R_{20}''\), which can be somewhat different from \(R''\), but with enough overlapping so that it is also able to bind to and activate the G protein. Thus

**FIGURE 5.** Theoretical energy landscapes and activation pathways for the interaction between the \(\beta_2\text{AR} \) and ligands of different efficacies.

Energy in the z-axis (color) as a function of the conformational changes in TM7 and TM6 (in the x-axis), and of the conformational changes in TM5 and neutralization of Asp130(3.49) in the DRY motif (in the y-axis) is shown at left. Activation follows the path traced by the lines in the 3D landscapes. Slicing the 3D surface along these paths results in the simpler 2D landscapes depicted at right.

A: absence of ligand, small conformational changes in the TM3/TM5 interface and plasticity in the water cluster around the NPxxY motif result in a receptor population that fluctuates between the \(R\), \(R'\), and \(R''\) conformations. The conformations \(R''\) and \(R^*\) can get populated by movement of TM6 and neutralization of Asp130(3.49) in the DRY motif of TM3. The \(R^*\) conformation is able to bind to and activate the G protein, resulting in certain constitutive activity.

B: binding of an inverse agonist stabilizes the \(R''\) conformation, where TM5 and TM7 are slightly rearranged, resulting in a change in the receptor energy landscape (compare A and B). As a result, the receptor gets trapped in the \(R''\) conformation. C: an agonist binds to and selects the \(R^*\) conformation and changes the energy landscape in such a way that the barrier to reach \(R'''\) (i.e., to move TM6) is lowered. The stabilized open crevice in the cytoplasmic side favors the neutralization of Asp130(3.49) in the DRY motif so Arg131(3.50) can reach the extended conformation needed for binding of the G protein. D: a partial agonist binds to and stabilizes \(R''\) to induce a change in the energy landscape. In this case, a conformation \(R_{20}''\) different from the \(R'''\) conformation stabilized by agonists, becomes available. In this new energy landscape (compare C and D), the lower energy conformation corresponds to \(R_{20}''\), which can be somewhat different from \(R''\) but with enough overlapping so that it is also able to bind to and activate the G protein. Thus the stabilization of a new energy landscape implies that the receptor must follow a different activation pathway (compare the black and gray lines in C and D).
the stabilization of a new energy landscape implies that the receptor must follow a different activation pathway (compare the black and gray lines in Figure 5C AND 5D). This last example shows how the use of 3D landscapes results in an easier way to depict activation by different ligands, resulting in specific conformational changes that can be shown as new pathways in altered energy landscapes.

**Conclusions**

Activation of GPCRs occurs through a series of conformational intermediates that can be visualized as wells in an energy landscape. Ligands influence GPCR function by shifting this conformational equilibrium and changing the shape of the landscapes.

In addition to ligand binding, there are many elements that can influence energy landscapes (FIGURE 1). One of the major factors is protein-protein interactions, either within the membrane (oligomerization and interaction with other membrane proteins) or with extracellular and intracellular partners. For instance, the SII peptide is able to trigger arrestin-specific activation pathways in the angiotensin II receptor (51), possibly by stabilizing certain conformations in a SII-specific energy landscape. The environment also influences GPCR function. For instance, protonation of Glu134(3.49) in the key (E/D)RY motif is a thermodynamic prerequisite for full receptor activation in membranes but not in detergent (27). Moreover, it has been shown that changes in cell membrane tension and fluidity (7) and in membrane voltage (30) may modulate GPCR activation by affecting conformational dynamics and altering their ability to couple to the G protein. In addition, it has been extensively shown that GPCR activation is pH dependent (18, 29). Finally, posttranslational modifications, such as glycosilation, phosphorylation, or acylation also have a role. For instance, phosphorylation of GPCRs by kinases stabilizes conformations that can be recognized by arrestins (43). These factors are key to translating the dynamic behavior of the protein into ordered function.

To characterize the energy landscapes of GPCR activation, we need to identify the local energy minima in a given reaction pathway, i.e., the intermediate (meta)stable GPCR conformations along the activation process. These intermediates can be identified through X-ray crystallography (e.g., by crystallizing GPCR-effector complexes), electron microscopy (39), or, at a lower resolution, fluorescence spectroscopy (25). In addition, we also need to determine the free energy of activation from the rates of interconversion as well as relative free energies of the substates from the relative populations. These kinetic and thermodynamic parameters can be obtained by Carr-Purcell-Meiboom-Gill (CPMG) nuclear magnetic resonance spectroscopy experiments (15). In addition, CPMG dispersions can detect transient conformers representing only 0.5% of the ensemble population. These rare conformational states are key steps in the activation pathways of GPCRs, but their structure is impossible to access by crystallography.

In summary, energy landscapes are a useful tool to connect GPCR structure, dynamics, and function. Understanding how ligands stabilize different functional states on the energy landscape is key to understanding the structural basis of efficacy. Moreover, characterizing the reaction pathways and identifying intermediate conformational states may facilitate the development of more selective drugs capable of modulating a specific signaling pathway, thereby improving therapeutic activity and minimizing undesirable side effects.

We thank Dr. Soren Rasmussen for providing Figure 1. This work was supported by the Ministerio de Educación y Ciencia (Spain) through the Ramon y Cajal Program (to X. Deupi) and the National Institute of Neurological Disorders and Stroke Grant N5-028471 and the Mathers Charitable Foundation (to B. K. Koblika). Present address of X. Deupi: Condensed Matter Theory Group and Laboratory of Biomolecular Research, Paul Scherrer Institut, Switzerland (e-mail: xavier.deupi@psi.ch).

No conflicts of interest, financial or otherwise, are declared by the author(s).

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