Membrane cholesterol access into a G-protein-coupled receptor

Ramon Guixa-González1,2,*, José L. Albasanz3,*, Ismael Rodriguez-Espigares1, Manuel Pastor1, Ferran Sanz1, Maria Martí-Solano1, Moutusi Manna4, Hector Martinez-Seara4,5, Peter W. Hildebrand2, Mairena Martín3 & Jana Selent1

Cholesterol is a key component of cell membranes with a proven modulatory role on the function and ligand-binding properties of G-protein-coupled receptors (GPCRs). Crystal structures of prototypical GPCRs such as the adenosine A2A receptor (A2AR) have confirmed that cholesterol finds stable binding sites at the receptor surface suggesting an allosteric role of this lipid. Here we combine experimental and computational approaches to show that cholesterol can spontaneously enter the A2AR-binding pocket from the membrane milieu using the same portal gate previously suggested for opsin ligands. We confirm the presence of cholesterol inside the receptor by chemical modification of the A2AR interior in a biotinylation assay. Overall, we show that cholesterol’s impact on A2AR-binding affinity goes beyond pure allosteric modulation and unveils a new interaction mode between cholesterol and the A2AR that could potentially apply to other GPCRs.
G-protein-coupled receptors (GPCRs) are complex signaling machines that are embedded in the cell membrane. They are able to respond to extracellular signalling stimulus by triggering diverse intracellular pathways of high relevance for human biology. Despite tremendous advances in characterizing GPCR structure and activation mechanisms, relatively little is known about the role of the membrane environment or of specific membrane lipid composition in receptor function.

Recent work shows that membrane phospholipids can allosterically modulate the activity\(^1\) and oligomerization\(^2\) of GPCRs. In addition, membrane cholesterol significantly modulates the stability, ligand-binding properties and function of several GPCRs (reviewed in refs 3–6). Specifically, the presence of cholesterol in cell membranes can either enhance\(^7–11\) (that is, positively modulate) or decrease\(^12–15\) (that is, negative modulation) ligand binding and/or functional properties of different GPCRs (see Supplementary Table 1 for a comprehensive summary). A well-known example of this modulation is observed in rod outer segments for the prototypical receptor—rhodopsin, where higher cholesterol concentrations in newly formed basal disks are used by these cells to stabilize the structure of metarhodopsin I (MI), thus hampering the formation of the active intermediate metarhodopsin II (MII)\(^16–19\).

Whether this modulation is exerted through indirect effects\(^20,21\) (that is, changes in membrane properties), direct interactions\(^22–25\) between cholesterol and GPCRs, or both, has for long been a matter of intense debate (see ref. 6 for a recent review on this topic). Specific cholesterol-binding sites have been identified at the surface of different GPCRs\(^26\), suggesting a potential allosteric role of cholesterol in modulating GPCR function. Intriguingly, other studies postulate that closely related cholesterol derivatives can even modulate the function of certain class-A GPCRs from the orthosteric binding pocket thus acting like conventional class-A GPCR ligands. For instance, oxysterol is thought to follow this binding mode at the Epstein–Barr virus-induced G-protein coupled receptor 2 (GPR183)\(^27\) or the chemokine receptor CXCR2 (ref. 28). Similarly, oxysterol derivatives are known allosteric modulators of the oncprotein Smothened\(^29\) (SMO), a class F GPCR. A very recent crystal structure of SMO shows one cholesterol molecule in the binding site of the extracellular domain of this receptor\(^30\).

The adenosine A\(_2\)AR receptor (A\(_2\)AR) is a class-A GPCR that plays a major role in the heart and brain by regulating oxygen consumption and blood flow\(^31\). In fact, in the central nervous system (CNS)\(^32\), the A\(_2\)AR constitutes a potential therapeutic target for the treatment of Alzheimer and Parkinson’s disease\(^33,34\). Cholesterol binding to the A\(_2\)AR at allosteric sites has been previously demonstrated by a high-resolution X-ray crystal structure (PDB ID 3EML)\(^24\). Computational work has further quantified allosteric cholesterol binding to the receptor surface\(^35,36\) and suggested a stabilizing effect on the apo-form of the A\(_2\)AR\(^35\). However, the ability of cholesterol to impact ligand-binding properties at the A\(_2\)AR remains unclear.

For this purpose, in the present study we analysed the influence of cholesterol depletion on ligand binding and studied the dynamics of cholesterol–A\(_2\)AR interaction by extensive long-scale molecular dynamics (MD) simulations. Our simulation data reveals an unexpected mechanism of cholesterol action on ligand binding consisting on the entry of a cholesterol molecule into the receptor transmembrane bundle. Different lipophilic ligands that bind to the orthosteric site of class A GPCRs are suggested to access the protein from the membrane milieu (reviewed in ref. 6). Interestingly, in recent crystal structures of rhodopsin\(^37,38\), a molecule of a commonly used detergent (that is, n-octyl β-d-glucopyranoside) replaced retinal from the ligand-binding pocket. Therefore, as recently discussed by Gimpl\(^6\), it would seem plausible that cholesterol can access the interior of class A GPCRs like the A\(_2\)AR. To validate this mechanism of action, we used a specifically tailored experimental approach to assess cholesterol impact on chemical modification of the A\(_2\)AR interior. Taken together, our combined long-scale MD simulation and experimental results show that cholesterol can compete with orthosteric ligands by entering the receptor interior from the membrane side.

### Results

#### Effect of cholesterol depletion on A\(_2\)AR ligand binding

To investigate the effect of membrane cholesterol on A\(_2\)AR-binding properties, we removed cholesterol from the membrane and monitored the specific binding of the radioligand [\(^3\)H]ZM241385 (Fig. 1), a selective antagonist of this receptor. We depleted membrane cholesterol by treating C6 glioma cells with methyl-β-cyclodextrin (M\(_\beta\)CD), a specific cholesterol-sequestering agent, for time lengths between 0 and 50 min (Fig. 1). Our data indicate that M\(_\beta\)CD is able to deplete around 70–80% of membrane cholesterol after 30 min. To accurately assess the level of cholesterol depletion, we carried out targeted lipidomics in plasma membranes. Remarkably, 40 min treatment with 5 mM M\(_\beta\)CD according to the described protocol in the method section depletes up to 61% of cholesterol from the membrane (Supplementary Fig. 1). In addition, further radioligand binding assays using the former membrane preparations (Supplementary Fig. 2) confirm the effect we describe in Fig. 1 and Supplementary Fig. 3 using intact cells.

To rule out any cytotoxic effect of M\(_\beta\)CD or WSC treatment, cell viability was determined at 20, 40 and 60 min after M\(_\beta\)CD or WSC treatments using the XTT method (see Methods). As shown in Supplementary Fig. 4A, neither M\(_\beta\)CD nor WSC treatments affected cell viability. Here it is worth to highlight that cells remain viable despite the depletion of more than 60% of their membrane cholesterol using the M\(_\beta\)CD treatment

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**Figure 1 | A\(_2\)AR-specific binding and cholesterol content.** Time course of 5 mM M\(_\beta\)CD addition on [\(^3\)H]ZM241385-specific binding to A\(_2\)AR in intact cells and membrane cholesterol content. This experiment was carried out using a saturating radioligand concentration of 40 nM. Mean ± s.e.m. values obtained from \(n = 3\) separate experiments carried out in triplicate.

\(*P < 0.05, \ **P < 0.01 and \ ***P < 0.001\) significantly different from control value (time 0, \(n = 5\)) according to a Student’s \(t\)-test.
Interestingly, addition of cholesterol significantly decreases and 7) and targeted lipidomic analysis (Supplementary Fig. 1). Fractions by filipin fluorescence staining (Supplementary Figs 6 membrane was monitored in intact cells and plasma membrane using water-soluble cholesterol (WSC) (Fig. 3).

Of this effect, we replenished membranes with cholesterol this inhibitory effect (see Fig. 2). To validate the reversibility using a wide range of radioligand concentrations confirm likely due to a compensatory mechanism to maintain cholesterol

Control (closed circles) and 5 mM MJ(CD) (40 min) (open circles) treated cells were incubated with different concentrations of [3H]ZM241385 as described in the Methods. These results are mean ± s.e.m. values obtained from six separate experiments carried out in duplicate. Kinetic parameters (Bmax and KD) of the corresponding saturation binding curves are indicated at the bottom the figure. **P < 0.01 significantly different from control value according to a Student’s t-test.

Remarkably, the lack of cholesterol increases the specific binding of [3H]ZM241385 to the A2AR by more than 100% (30 min) when compared to non-treated cells (0 min) (Fig. 1 and Supplementary Fig. 3). Longer incubation times with MJ(CD) (that is, 240 min) did not result in higher levels of A2AR specific binding (469.7 ± 13.7 fmol per mg protein, n = 2), likely due to a compensatory mechanism to maintain cholesterol homeostasis in treated cells. Saturation binding experiments using a wide range of radioligand concentrations confirm this inhibitory effect (see Fig. 2). To validate the reversibility of this effect, we replenished membranes with cholesterol using water-soluble cholesterol (WSC) (Fig. 3).

Adequate cholesterol depletion and insertion into the membrane was monitored in intact cells and plasma membrane fractions by filipin fluorescence staining (Supplementary Figs 6 and 7) and targeted lipidomic analysis (Supplementary Fig. 1). Interestingly, addition of cholesterol significantly decreases [3H]ZM241385 binding in cell membranes either untreated (Fig. 3, columns 1, 3 and 4) or previously depleted from cholesterol using MJ(CD) (Fig. 3, columns 2, 5 and 6). This clearly suggests that cholesterol has an inhibitory effect on [3H]ZM241385 binding to the A2AR. This effect was confirmed in membranes from control cells by competitive binding experiments in the presence of increasing WSC concentrations (see Fig. 4 and Supplementary Note 1). To rule out that the former effect is the result of a higher number of A2ARs available due to an inhibition of receptor internalization by MJ(CD), we performed new binding assays in the presence of different inhibitors of endocytosis (see details in Supplementary Note 2). As shown in Supplementary Fig. 8, inhibiting endocytosis does not significantly modulate A2AR specific binding, hence demonstrating that receptor internalization is not involved in the cholesterol-mediated modulation of A2AR-specific binding. While cholesteryl hemisuccinate,
a cholesterol derivative, enhances the stability and activity of detergent-solubilized A2AR, our data clearly shows that naturally occurring cholesterol has an inhibitory effect on [3H]ZM241385 binding to the A2AR in more physiological environments (that is, intact cells and cell membrane preparations). Our findings go along with previous experimental evidence in several class A GPCRs reporting a negative cholesterol-mediated modulation of ligand binding (Supplementary Table 1).

All-atom MD simulations of cholesterol interaction sites. To shed light on the structural basis of the observed cholesterol modulation, we carried out a complete set of atomistic MD simulations of the A2AR (Supplementary Table 2). As described in the Methods section, we used a comprehensive set of lipids of different chains, nature and length to model a native-like lipid bilayer rich in cholesterol (Supplementary Table 3). The crystal structure of the A2AR was embedded in the former membrane and the system was solvated, neutralized, adjusted to an ionic strength of 150 mM Na\(^+\)Cl\(^-\) and equilibrated following standard protocols (see Methods). First, we simulated a set of 4/1 μs replicas to study the frequency and stability of the interaction between cholesterol and the A2AR. The analysis of the accumulated 4/μs shows that on average 12 cholesterol molecules are in contact (below 2.9 Å) with the A2AR throughout the simulation (Supplementary Fig. 9). This value is in agreement with the number of cholesterol molecules required for an ideal conformational stability of the A2AR experimentally observed in cholesterol-rich micelles.

While our simulations show several transient cholesterol–A2AR interactions (light grey bars, Fig. 5 right), certain cholesterol molecules establish permanent binding interactions to the receptor over nearly 100% of the simulation time (dark grey bars, Fig. 5 right). To spot preferred cholesterol interaction sites at the A2AR surface, we used a 3D volumetric map to depict the density of cholesterol molecules in the simulation (Fig. 5 left and Supplementary Fig. 10). In agreement with Lee et al., our data show that transmembrane helices 2 and 3 (TM2–3), TM3-4-5 and TM7-1 are preferred interaction areas of cholesterol at the A2AR surface. Notably, one of the predicted interaction sites, namely TM2–3, overlaps with one of the cholesterol binding sites shown in the experimental high-resolution structure of A2AR in complex with ZM241385 (ref. 24; Fig. 5 left).

Cholesterol accesses the A2AR interior in MD simulations. Strikingly, the volumetric analysis also shows high cholesterol density inside the transmembrane bundle of the A2AR (Fig. 5, left) indicating that cholesterol entered the A2AR from the membrane milieu. A visual inspection of the individual trajectories shows that one cholesterol molecule spontaneously accesses the interior of the protein from the extracellular leaflet through helices TM5–6 occupying a key area of the orthosteric binding pocket (see replica 1 in Fig. 5 and Supplementary Movies 2 and 4). Although cholesterol has been shown to occupy deeply buried sites in other membrane proteins, this is the first dynamic view of membrane cholesterol spontaneously invading the orthosteric binding pocket of a GPCR.

To exclude simulation artefacts and better explore the cholesterol entry pathway, we performed new simulations and studied the tendency of cholesterol to access the A2AR. First, we selected four representative snapshots from the original cholesterol entrance trajectory (that is, replica 1) prior to the complete invasion of the receptor (Fig. 6). Then, each starting point was used to re-spawn 10 new trajectories of 100 ns each (that is, 10 × 4 × 100). To quantify cholesterol progression towards the interior of the protein, we monitored the distance between cholesterol and residue E1.39 (Fig. 6). In most of these short trajectories, cholesterol does not back away from the receptor but it stays bound or progresses towards the interior of the receptor. As shown in Fig. 6 and Supplementary Table 4, cholesterol progression is much faster once cholesterol slightly tilts down (Fig. 6a,b) adopting a favored position to enter the receptor. Inclusion of intracellular loop 3 (ICL3) do not
have a significant effect (Supplementary Fig. 11). These data clearly suggest that cholesterol can spontaneously access the A2AR through a portal gate between TM5 and 6.

The nature of the membrane environment could be one of the driving forces behind the spontaneous cholesterol entrance into the A2AR. As shown in Supplementary Movie 3, an unsaturated phospholipid (in yellow) along with a cluster of four cholesterol molecules seem to influence cholesterol entrance by preventing it from diffusing back to the membrane bulk.

To study the impact of membrane composition, we substituted the compact membrane environment used so far by a pure 1-palmitoyl-2-oleyl-sn-glycero-3-phosphocholine (POPC) bilayer (see Methods) leaving intact both the target cholesterol and the A2AR (see Methods). Interestingly, in the absence of a more compact and thicker membrane (Supplementary Fig. 12), cholesterol progression towards the interior of the protein is significantly diminished after 100 ns (see Supplementary Fig. 13 and Supplementary Table 4). Therefore, our simulations suggest that the ability of cholesterol to access the interior of the A2AR can be modulated by the nature of the membrane environment.

The sequence of events during cholesterol entrance is depicted in Fig. 7. Initially, cholesterol is interacting with other membrane lipids (not shown) and water molecules (Fig. 7a) until it leaves this preferred position plunging its polar head into the hydrophobic core of the membrane (Fig. 7a,b). Next, cholesterol descends along TM5–6 guided by the formation of a hydrogen bond with the hydroxyl group of Y5.411 side chain (Fig. 7b). At this point, TM5 and TM6 are tightly packed involving an aromatic cluster of staggered residues F5.45, F4.44, H6.52 and W6.48 (red arrows, Fig. 7b). Thereafter, cholesterol tilts 90° and pushes aside the aromatic side chains of residues F5.45 and F4.44–H6.52, see red crosses in Fig. 7c,d), thus likely hampering the reverse progression of cholesterol towards the membrane bulk.

To better characterize the behaviour of cholesterol inside the receptor, we extended three of the original 100 ns replicas (namely replicas 1, 35 and 38 in Fig. 6) up to 10 μs (3 × 10 μs). Once inside the receptor, cholesterol molecules explore the interior of the protein by establishing transient interactions rather than adopting a stable binding pose. Interestingly, our simulations show that cholesterol highly populates a specific area of the A2AR binding pocket (yellow surface, Fig. 8a and Supplementary Movie 4) that highly overlaps with the classical orthosteric binding site and the position of the ZM241385 ligand.

**Figure 6 | Short simulation replicas of cholesterol entrance.** Boxplots display the distance between the centre of mass of cholesterol and residue E1.39 for a set of 40 replicate simulations of 100 ns. Four different starting positions (a–d) re-spawned from the original cholesterol entrance trajectory were used to run each 10 replicates (that is, 1–10, 11–20, 21–30, 31–40). The distance at the beginning of the simulation as measured from the snapshot used to re-spawn each set of 10 trajectories (that is, a–d) is represented as one single red horizontal line in each of the graphs. Average distance for each set of replicates is reported in Supplementary Table 4. Inset figures show the initial structure of the A2AR (in blue) and cholesterol residue (in orange) used to start each set of simulations. E1.39 residue is displayed as van der Waals spheres. The BENDIX plugin for VMD was used to depict protein helices. Protein loops were omitted for clarity.
bound to the A<sub>2A</sub>R crystal structure (PDB:3EML) (red sticks, Fig. 8a). Consequently, our data indicate that, if cholesterol reaches the interior of the protein, it will likely hamper the binding of A<sub>2A</sub>R ligands such as ZM241385 to the orthosteric binding site. This finding represents one plausible mechanism behind the observed increase in [<sup>3</sup>H]ZM241385 binding upon cholesterol depletion.

**Experimental validation of cholesterol occupying A<sub>2A</sub>R.**
Providing experimental evidence for a native membrane lipid occupying the interior of a GPCR is a challenging mission. Cholesterol has been shown to bind the A<sub>2A</sub>R surface in recent X-ray data<sup>24</sup>. However, there is currently no structural hint or experimental evidence pointing towards cholesterol ability to occupy the interior of this receptor. In this study, in an
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Figure 9 | Biotinylation experiments. (a) Effect of the MTSEA-B reagent and cholesterol depletion on $[^3H]$ZM241385-specific binding (mean ± s.e.m.) at a saturating radioligand concentration of 40 nM. (b) Scheme of cholesterol influence on receptor biotinylation according to the experimental results presented in a. Bar 1 (control, n = 11): ZMA binding in untreated (cholesterol-containing) control cells yields low ZMA binding due to cholesterol shielding effect. Bar 2 (n = 5): ZMA binding in MβCD-treated cell membranes (cholesterol-depleted) results in highest ZMA binding due to an empty binding pocket. Bar 3 (n = 4): biotinylation of a cholesterol-depleted binding pocket results in a high degree of biotinylation in the orthosteric binding pocket. As the binding pocket is occupied by the biotinylated side chain of C3.30, ZMA binding is reduced. Bar 4 (n = 4): low biotinylation degree due to the shielding effect of cholesterol in the binding pocket. Subsequent removal of the shielding agent cholesterol with MβCD results in an empty binding pocket that allows for high ZMA binding. Bar 5 (n = 4): lowest ZMA binding due to the accumulated hindering effect of biotinylation and cholesterol. CHL, cholesterol (yellow); BIO, biotinylation (green); MβCD, methyl-β-cyclodextrin (blue); ZMA, ZM241385 ligand (red). In a ***P < 0.001 and **P < 0.01 significantly different from control 1, and *P < 0.05 significantly different between them (bars 3 and 4) according to a Student’s t-test.

attempt to provide experimental evidence supporting our computational findings, we adapted a biochemical assay originally used in GPCRs by Javitch et al.42 for scanning residues exposed to the binding-site crevice in the dopamine D₂ receptor. This assay takes advantage of the fact that thiol groups of cysteine residues facing the binding-site crevice of a GPCR will selectively react with hydrophilic and sulfhydryl-specific reagents such as derivatives of methanethiosulfonate (MTS) (detailed in Supplementary Note 3). As shown by the A₂AR crystal structure (PDB: 3EML)24 (see Supplementary Fig. 14 and Supplementary Table 5), water-accessible cysteine residues with a free sulfhydryl group are only found in the interior of the receptor. In our experiments, we chemically modified by biotinylation these water-exposed cysteines using β-n-biotinylaminoethyl-methanethiosulfonate (MTSEA-B), a positively charged MTS derivative similar in size and molecular weight to the ZM241385 ligand (see Methods). Figure 8c illustrates that biotinylation of residue C3.30 (green sticks) results in a modified side chain that occupies a large region of the orthosteric binding site that overlaps with the preferred binding site of cholesterol as well as of ZM241385 (yellow and red sticks, respectively, Fig. 8b). As shown in Supplementary Fig. 15, chemically modified cysteine residues C5.46 and C6.56 yield a similar overlap with the ZM241385 ligand. Accordingly, if cholesterol is able to invade the A₂AR-binding pocket as proposed by our simulations, cysteine biotinylation should be hindered in cholesterol-rich membranes compared to cholesterol-depleted membranes.

Based on this conceptual framework, we designed a new set of radioligand binding experiments using MTSEA-B to covalently modify cysteine residues in the A₂AR interior and MβCD to deplete membrane cholesterol (Fig. 9). Experiments are schematically depicted in Fig. 9b. First, we used two control experiments where we measured $[^3H]$ZM241385 in untreated (Bar 1) and cholesterol-depleted cells (Bar 2). These control experiments corroborated that cholesterol depletion by MβCD favours specific binding of $[^3H]$ZM241385 (Bar 2) by three fold when compared with untreated conditions (Bar 1). Then, we assessed if cysteine residues in the A₂AR interior are susceptible to MTSEA-B biotinylation in cholesterol-depleted conditions, where no competition between cholesterol and $[^3H]$ZM241385 is expected. We observe that after cholesterol depletion, biotinylation reduces $[^3H]$ZM241385 binding by ~40% (Bar 3) when compared with treatment with MβCD alone (Bar 2).

This marked reduction in specific binding strongly indicates that at least one of the cysteine residue in the A₂AR interior (Supplementary Note 3) is susceptible to biotinylation and confirms that its chemically modified side chain occupies the orthosteric-binding site (Fig. 9c). In the next measurement, we tested the influence of cholesterol binding in cysteine biotinylation (Bar 4). Remarkably, $[^3H]$ZM241385 binding increases by about 25% (Fig. 9, Bars 3–4) when receptors are treated with MTSEA-B prior to cholesterol depletion. This suggests that cholesterol has a shielding effect by protecting cysteine residues from biotinylation inside the receptor. Thus, this observation supports the presence of a cholesterol molecule inside the A₂AR transmembrane bundle with the ability to block access of MTSEA-B to water-accessible cysteine residues. Finally, we observe that if the system is biotinylated in the presence of cholesterol and cholesterol is not removed afterwards (Bar 5), there is an additive effect of cholesterol and biotinylation that hampers binding of $[^3H]$ZM241385, yielding less specific binding than in control conditions (Bar 1).
Cholesterol entry/exit pathways. Our data strongly suggest that a cholesterol molecule can penetrate the A2AR from the membrane side. In fact, membrane access of molecules to class-A GPCRs has been reported for few other cases such as the lipid receptor S1P1 (ref. 43), the cannabinoid CB2 receptor44 and the opsin receptor45. Among them, retinal uptake into the opsin receptor is probably the best-studied case where two potential gateways for ligand uptake or release have been described45. The larger opening (gateway 1) between helices 5 and 6 has initially been proposed to allow for retinal loading, whereas a smaller gate between helices 1 and 7 (gateway 2) allows for ligand exit46. In contrast, more recent studies point to a retinal entrance/exit in the opposite sense47,48. The complete tunnel pathway for retinal into the binding pocket of opsin including gateways 1 and 2 can be detected using the Caver software49 (red surface, Fig. 10a).

In comparison, Caver computation for the simulated A2AR (150 ns) reveals three gateways 1 to 3 (yellow surface, Fig. 10b). The last largest gateway 3 is the classical channel for ligand entrance from the extracellular side into the receptor. The smallest gateway 2 is directed towards helices 1 and 7 whereas the intermediate gateway 1 is located between helices 5 and 6 (yellow surface, Fig. 10b). Remarkably, we find that gateway 1 is flanked by two residues, namely V5.40 and F5.44, which are conserved between the A2AR and opsin (Fig. 10a,b). Hence, our data indicate that the opening that cholesterol uses to penetrate into the A2AR is identical to the opening that retinal takes to enter/exit the opsin receptor.

Unveiling the putative exit pathway of cholesterol from the A2AR is currently beyond the reach of all-atom unbiased simulations. If cholesterol behaved in a similar way to classical GPCR ligands, this entry/exit pathway could occur via the extracellular domain, as suggested elsewhere28,50. However, due to the large hydrophobic moiety of the cholesterol molecule, it is not likely that cholesterol abandoning the A2AR-binding pocket via the extracellular aqueous phase is energetically favoured. To determine the energetic cost of such exit route, we carried out a set of biased simulations (see Methods) where we computed the free energy profile (PMF) of extracting cholesterol from the binding site into the extracellular water phase. These simulations confirm that the energetic cost of a complete cholesterol exit from its observed binding site to the water phase is as high as 120 kJ mol⁻¹ (Supplementary Fig. 16). Hence, it is more likely that cholesterol leaves the A2AR via the transmembrane helices. In this context, we find that TM2-TM1 or TM7-TM1 could be possible cholesterol exit ways, as detailed in Supplementary Note 4. Notably, TM7-TM1 exit is in line with A2AR channels that have been computed using the Caver software (Fig. 10b) as well as with the described gateway 2 in the crystallized structure of opsin (Fig. 10a)47,48.

Discussion

In the present work, we have studied the effect of membrane cholesterol on the ligand-binding properties of the A2AR. Our in vitro experiments demonstrate that cholesterol significantly decreases the binding of the antagonist [³H]ZM241385 to the A2AR. Molecular insights obtained in this and other studies point to numerous allosteric binding sites at the A2AR surface that could potentially be involved in reducing ZM241385 binding. However, our work reveals an additional and unexpected mode of cholesterol action. We have observed that cholesterol modulates orthosteric ligand binding at the A2AR after entering from the membrane side. These results have been validated combining both computational and specifically tailored experimental approaches.
Our study suggests that the gateway that cholesterol uses to enter the A<sub>2A</sub>R is evolutionarily conserved and identical to the retinal gateway present in the opsin receptor. Since molecules of a similar size and amphiphilic nature have been shown to access the retinal-binding pocket<sup>37</sup>, it is reasonable to speculate that the negative modulation exerted by cholesterol on rhodopsin function<sup>16–19</sup> could be partly mediated by the new mechanism proposed here. Taken all together, cholesterol is likely to modulate A<sub>2A</sub>R ligand-binding properties through a mixed mode of action, namely both orthosteric- and allosterically. Nevertheless, further research shall be aimed to unveil the exact contribution of each mode of cholesterol-mediated modulation at this or other GPCRs.

All in all, our results provide an important advance in the understanding of the interplay between cholesterol and the A<sub>2A</sub>R. Ultimately, our work highlights the importance of considering the influence of different levels of membrane cholesterol in GPCR function and stresses the importance of accounting for these particular cholesterol effects in the study of cardiovascular and CNS disorders. From a broader perspective, our work opens the door to new studies on the effects of cholesterol and other endogenous lipids in GPCRs modulation. This knowledge paves new roads for exploring potential therapeutic uses of membrane sterols or sterol-mimetic molecules in GPCR drug discovery.

Methods

**Cholesterol and MTS E B experiments.** Materials—[3H]ZM241385 ([2-<sup>3</sup>H]-1-[4-(2-[7-amino-2-(2-furyl)-1-[1,2,4] triazolo [2,3-x] [1,3,5] triazin-5-ylamino)]ethyl)phenol 27.4 Ci mmol −1) was purchased from ARC (St Louis, MO, USA). The reagents MjCJd, WSC, MTS E B and Thephoplyline were acquired from Sigma Aldrich (Madrid, Spain). Dynasore (3-Hydroxynaphthalene-2-carboxylic acid, 3,4-dihydroxybenzylidenehydrazone) and Pitstop 2 [N-[5-(4-bromobenzylidene)-4-oxo-4,5-dihydro-1,3-thiazol-2-yl]naphthalene-1-sulfonamide] were from Abcam Biochemicals (Cambridge, UK). Liquid scintillation cocktails were supplied by Perkin Elmer (Boston, MA, USA). Additional reagents were of analytical grade.

**Cell culture.** C6 cells from rat glioma were obtained from the American Type Culture Collection (ATCC). Dulbecco’s modified Eagle’s medium was supplemented with 2 mM glutamine, 10% fetal calf serum, 1% nonessential amino acids and antibiotics. Cells were maintained at 37°C in a humidified atmosphere of 95% air and 5% CO<sub>2</sub> (ref. 51).

**Plasma membrane isolation.** Cells were homogenized on ice-cold isolation buffer (50 mM Tris-HCl pH 7.4, 10 mM MgCl<sub>2</sub>-containing protease inhibitors) and centrifuged at 4°C for 5 min at 1,000g in a Beckman JA 21 centrifuge. The supernatant was centrifuged at 4°C for 20 min at 27,000g and the pellet was resuspended in isolation buffer<sup>51</sup>.

**Radioligand binding assays.** Specific binding to A<sub>2A</sub>R was assayed in intact C6 cells in the conditions described above. Intact cells grown in 24-well plates were incubated with 40 nM [3H]ZM241385 (that is, saturating ligand concentration, based on our previous work<sup>52</sup>). Adenosine deaminase (ADA, 5 U ml<sup>−1</sup>) was used to remove endogenous adenosine. [3H]ZM241385 ranging from 1.25 nM to 40 nM was utilized in saturation binding assays, where 5 mM theophylline was used to obtain non-specific binding. After 2h at 25°C, cells were washed with ice-cold culture medium, lysed with 0.2% SDS, and transferred to vials to count radioactivity. Two wells from each plate were employed for protein content measurement.

When binding assays were performed in plasma membranes, fifty to one hundred micrograms of protein were pre-incubated with 5 U ml<sup>−1</sup> ADA for 30 min at RT and maintained for 2h at 25°C in the presence of 20 or 40 nM [3H]ZM241385. Non-specific binding was also obtained with 5 mM theophylline, as in previous work<sup>52</sup>. For competition binding experiments, different (1 μM – 3 mM) concentrations of WSC were used to displace total binding of 20 nM [3H]ZM241385. Binding to plasma membranes was finished by rapid filtration through Whatman GF/B filters, which were immediately washed and counted. Radioactivity measurements in vials or filters were performed in a Microbeta Trilux liquid scintillation counter (Wallac).

**Cholesterol analysis in intact cells.** Cholesterol content in intact C6 cells was measured with a Cholesterol Quantiation kit (MAK043) from Sigma (Madrid, Spain), following manufacturer’s instructions. Briefly, samples (106 cells) were extracted with 200 μl of chloroform/isopropanol/GEFAL CA-630 (7:1:0.1) in a micro-homogenizer. After centrifugation at 13,000g for 10 min to remove insoluble material, the organic phase of samples (160 μl) was transferred to a new tube and dried at 50°C in a SpeedVac for 30 min to remove chloroform. Dried lipids were then dissolved with 200 μl of the Cholesterol Assay Buffer. Fifty microliters of samples and standards (1–5 μg) were added to 50 μl of reaction mixture and absorbance at 570 nm measured after 60 min incubation at 37°C.

**Filtrin fluorescence staining.** A cell-based Cholesterol Assay Kit from Abcam (Cambridge, UK) was performed in intact C6 cells in order to visualize and measure cholesterol by using Filipin III as a fluorescence probe of cholesterol. Briefly, after removal of culture medium from wells, cells were fixed for 10 min and washed (3 × 5 min). Filipin III solution was added to each well assayed and maintained in the dark for 45 min at room temperature. After washing of cells (2 × 5 min) fluorescence images were obtained with a digital camera (Leica DFC350FX), attached to a Leica DMi6000B (Leica Microsystems, Wetzlar, Germany) fluorescent microscope using ×20 HCX PL FLUOTAR objective.

**Depletion and readding of cholesterol in living C6 cells.** To test whether cholesterol could enter the cells with 5 mM MjCJd for indicated period of time at 37°C. Cholesterol enrichment of the cells was started with 1 mM WSC for indicated period of time at 37°C. Finally, chemical modification of the A<sub>2A</sub>R-binding pocket by biotinylation was performed with N-biotinylaminomethylmethanethiosulfonate (MTSEA-B). This compound was dissolved in DMSO, and aliquots of 100 mM stock solution were thawed just prior to use. When needed, cells were incubated in 500 μM MTS E B for 5 min at 37°C. In all cases, treatment was performed in serum-free medium.

**Lipidomic analysis.** Plasma membrane preparations were isolated from control (n=3), 5 mM MjCJd 40 min (n=2) and 1 mM WSC 50 min (n=2) treated cells. Membrane samples containing deuterated cholesterol D7 as internal standard were mixed with an equal volume of methanol and two volumes (0.2 ml) of chloroform. After each addition tubes were vortexed for 10 s. Chloroform phase (lower) was transferred to a glass tube after centrifugation for 15 min at 4°C and 1,000g. This last step was repeated twice. The chloroform phase was evaporated in a Speed Vac (Thermo Fisher Scientific, Barcelona, Spain) and resuspended in 50 μl of methanol/chloroform (3:1).<sup>53</sup> These lipid extracts (2 μl) were analysed by means of mass-spectrometry using a HPLC 1290 series coupled to an ESI-Q-TOF MS/MS 6520 (Agilent Technologies, Barcelona, Spain). LC/MS analysis required an XBridge BEH C18 shield column (100 mm × 2.1 mm ID × 1.7 μm) from Waters (Milford, MA, USA) kept at 80°C. Mobile phases (0.5 ml min<sup>−1</sup>) consisted of 20 mM ammonium formate (pH 5) (A) and methanol (B). The gradient profile was: 50–70% B in 14 min, 70–90% B in 50 min, isocratic separation of 90% B during 15 min, 90–100% B in 5 min, and maintained so for an additional 5 min.<sup>55</sup> This protocol allowed the orthogonal characterization of lipids based on exact mass (<10 p.p.m.) and on retention time features. Collection of data was achieved in both positive and negative electrospray ionization time-of-flight modes and performed in full-scan mode at 100–3,000 m/z in an extended mass range (2 GHz), using N2 (51 min<sup>−1</sup> , 300°C) as nebulizer gas. The capillary voltage was 3,500 V (1 scan per s). Data were recorded and analysed by MassHunter Data/Qualitative analysis software (Agilent Technologies, Barcelona, Spain) to obtain the molecular features of the samples.<sup>56</sup>

**Inhibition of endocytosis in living C6 cells.** To inhibit endocytosis, C6 cells were incubated with 80 μM Dynasore, a cell-permeable dynamin inhibitor, or 25 μM Pitstop 2, for 20 or 40 min at 37°C (see Supplementary Note 3 for more details).

**Cell viability assay.** Cells were seeded (10<sup>4</sup> cells per well) and grown in 96-well tissue culture plate and incubated with 0.3 mg ml<sup>−1</sup> XTT solution (sodium 3′-[1-(phenylaminocarboxyl)-3,4-tetrazolium]-bis (4-methoxy-6-nitro) benzene sulfonic acid hydrate) for 30 min at 37°C in control, MjCJd- or WSC-treated cells. The cleavage of XTT to form an orange formazan dye by viable cells was monitored by reading absorbance at 475 and 690 nm according to the manufacturer’s protocol (Cell Proliferation Kit II, Roche, Mannheim, Germany).

**Protein determination.** Protein concentration was measured by the Lowry method, using bovine serum albumin as standard.

**Statistical analysis.** The binding data were analysed using Student’s t-test, one-way analysis of variance and nonlinear regression fitting to saturation: 

\[
Y = B_{max} X/(K_d + X)^{-1}
\]

or complement:

\[
Y = 100 \left( 1 + 10^{[(LogEC_{50} - X)/HillSlope]} \right)
\] (2)
binding curves with the GraphPad Prism 5.0 program (GraphPad Software, San Diego, CA, USA). Differences between mean values were considered statistically significant at \( P < 0.05 \).

**Water-accessible cysteine residues in the A2AR.** According to high-resolution crystallography data \(^{10}\), water-accessible cysteine residues with free sulfhydryl groups are only found in the interior of the receptor as all extracellular water-accessible cysteines are engaged in disulfide bonds and therefore cannot react with the biotinyl reaction (see detailed description in Supplementary Note 3). Among all cysteines, C3 is the closest one to the ZMA ligand and is therefore used as representative in this manuscript (Fig. 8c).

**Classical (non-biased) MD simulations.** All non-biased simulations were performed using the ACEMD simulation package \(^{57}\). The CHARMm36 (ref. 58), CHARMm36c (ref. 59) and CHARMm27 \(^{(ref. 60)}\) force fields were used to represent lipids, cholesterol and proteins, respectively. A list of all simulations performed in this work is displayed in Supplementary Table 2.

Set-up. First, a monoclonal lipid membrane with an area of \( A = 100 \times 100 \ \AA^2 \) (in the membrane plane) was built using the CHARMm-GUI membrane builder \(^{61}\). The exact lipid composition is described below. This membrane was equilibrated for 1 \( \mu \)s in the NPT ensemble. Next, the crystal structure of the adenosine A\(_2\)R (PDB:3EML) \(^{35}\) was manually embedded into the equilibrated membrane using VMD1.9 \(^{(ref. 62)}\). The intracellular loop 3 (ICL3) of the A\(_2\)R, not resolved in any of the current GPCR crystal structures, was omitted and all titratable residues were left in the dominant protonation state at pH 7. Next, an NPT equilibration phase was performed to accommodate water and lipid molecules to the protein and to promote further mixing (see below).

Initial production run. Each of the equilibrated replicas above was simulated for 1 \( \mu \)s in the NVT ensemble (simulation set 1 in Supplementary Table 2).

Short replicas. Four different snapshots of the specific trajectory where the cholesterol enters the receptor (see Results section) were randomly selected (Fig. 6). Each of these snapshots served as a starting seed for 10 new replicas (4 \times 10) that were run for 100 ns in the NVT ensemble (simulation set 2 in Supplementary Table 2). To rule out an effect of ICL3 omission, we performed a similar set of simulations where the ICL3 was included (see simulation protocol below and Supplementary Fig. 11). To validate the membrane effect on cholesterol entrance, we substituted all membrane lipids by POPC. First, four new structures were generated for the lipid molecules from the current GPCR crystal structures. The CHARMm-GUI membrane builder \(^{41}\) was then used to embed each structure into a pure POPC bilayer of \( A = 100 \times 100 \ \AA^2 \). Systems were then minimized and equilibrated (see below). Similarly, each seed was used to simulate 100 ns of 4 \times 10 replicas in the NVT ensemble (simulation set 3 in Supplementary Table 2).

Long replicas. To study the behaviour of cholesterol inside the receptor, we chose 3 of the 100 ns trajectories where cholesterol seemed to have an effective progression towards the interior of the receptor (that is, replicas 1, 35 and 38 in Fig. 6). These simulations were run for 100 ns in the NVT ensemble (simulation set 5 in Supplementary Table 2).

**System composition.** We aimed to create a membrane environment as physiologically relevant as possible, by following key general tendencies observed in specific brain post-mortem studies relevant for the adenosine A\(_2\)R receptor \((A_2 \text{R})_4\) oligomerisation kinetics of adenosine A\(_2\)A and dopamine D\(_2\) receptors. As described in the main manuscript, one A\(_2\)R was receptor was then embedded into this membrane using VMD 1.9 (ref. 62) and the system was then solvated, neutralized and the ionic strength was adjusted using the CHARMm-GUI membrane builder \(^{41}\). Thereafter, a VMD script was used to re-hydrate the membrane patch using \( \sim 30 \) water molecules (TIP3P model) per lipid. The exact composition of the simulation systems is detailed in Supplementary Table 1. The lipid composition used yielded a protein-tolipid ratio of 1:3.37.

**Simulation protocol.** As described in the Methods section of the main manuscript, simulation sets 1 and 2 (Supplementary Table 1) share a common building phase, whereas the set-up of simulation sets 3 and 4 involved a complete substitution of the membrane environment or the inclusion of the ICL3, respectively. At the beginning of the equilibration phase for simulations sets 1 and 2 (Supplementary Table 2), harmonic positional restraints were applied to the \( C_6 \) atoms of the protein and the system was simulated for 10 ns. Such constraints were gradually released from the receptor over 5 ns and the system was further equilibrated for 100 ns. In simulations set 3 (that is, POPC), harmonic positional restraints were applied to all atoms of both protein and cholesterol and the system was equilibrated for 20 ns in the NPT ensemble. In simulation set 4 (that is, ICL3 included), after 10,000 steps of minimization, a gradual release of different applied harmonic constraints was used during 40 ns phase in the NPT ensemble. Harmonic restraints were first applied during 10 ns to all atoms of the system except for the intracellular polar head region of all membrane lipids. Constraints were then released for all water and ion atoms and the system was simulated for another 10 ns. Subsequently, the ICL3 region was released and the system (that is, only protein and target cholesterol restrained) was further equilibrated for 20 ns. NPT simulations were carried out at 310 K and 1 bar using the Berendsen barostat \(^{59}\) with a relaxation time of 400 and 2 fs integration time step. NVT simulations were run at 310 K, using the Langevin thermostator \(^{70}\) with a damping coefficient of 5 ps \(^{-1}\) and 4 fs integration time step. All along the simulations, van der Waals and short-range electrostatic interactions were cut off at 9 \( \AA \) and the particle mesh Ewald method \(^{71}\) was used to compute the long-range electrostatic interactions.

**Inclusion of ICL3.** The ICL3 structure was taken from PDB:3PWH and included in the A\(_2\)R structure using MODELLER v9.10 (ref. 72) and VMD \(^{59}\). The system was first minimized and the ICL3 subsequently equilibrated and relaxed for 50 ns in the NPT ensemble (see simulation protocol above).

**In silico model and conformational analysis for MTSEA-B chemical modification.** In a first step, the side chain of cysteines (C3.30, C5.46 and C6.56) of the A\(_2\)R (PDB:3EML) was chemically modified by attaching MTSEA-B using the builder tool of the MOE package (version 2016.08). In a second step, the conformational space of such chemical modification was explored using the LowModeMD method in the MOE package with: Rejection Limit 100, Iteration Limit 100, RMS Gradient 0.1, MM Iteration Limit 500, RMSD Limit 0.5, Energy Window 100, Conformation Limit 1000 and applying the Amber10:EHT force field. As a result of this search, we obtained 98 different conformers for the chemically modified residues C3.30 and C5.46 and 98 conformers for the chemically modified residue C6.56, as shown in Supplementary Fig. 15.

**Umbrella sampling simulations and free-energy calculations.** Biased simulations were used to compute the free energy of cholesterol extraction from the inside of the A\(_2\)R. To this end, we used the simulation engine Gromacs v5 (ref. 73) in combination with the CHARMm36 force field \(^{58}\). The recommended settings including the usage of Verlet lists, PME with 1.2 nm cutoff for electrostatics and a cutoff function for the Van der Waals at 1.2 nm with a force-switch starting at 1.0 nm were used \(^{58}\). The composition of the system is detailed in Supplementary Table 3. The potential of mean force was computed along the direction perpendicular to the membrane (that is, \( z \) axis). No bias was introduced at the \( x \) axis, that is, molecules could freely diffuse in the membrane plane (\( xy \) plane). The biased umbrella potential was defined on a coordinate \( z \) axis between the centre of mass of the A\(_2\)R backbone and the oxygen atom of cholesterol. In total, 35 different windows were probed. Harmonic restraints with a force constant of 2,000 kJ mol \(^{-1}\) nm\(^{-2}\) were imposed along the reaction coordinate. To ensure an adequate overlap between neighbouring windows, each window is separated by 0.1 \( \AA \) along the reaction coordinate. Based on the quality of the data and the presence of large energy barriers, each window was simulated within 200–600 ns. The accumulated simulation exceeds 8 \( \mu \)s. The reaction coordinate was monitored every 10 fs within each window. The last frame of the 200 ns simulation of the previous window was used as the starting configuration of the next one. Although this protocol cannot be used to simulate all windows in parallel, it drastically improves the equilibration time for each window. The first window corresponds to the cholesterol deeply buried in the ligand cavity. In the last windows, cholesterol moves freely in the water solution. We used the g_wham tool distributed with gromacs v5 (ref. 73) to compute the PMF (Supplementary Fig. 16) disregarding the first 10% of each trajectory to ensure an adequate overlap between the pathways. Autocorrelation time of the data was used to minimize correlation effects and to select the appropriated gathering frequency in the PMF curves. Bootstrapping of 1,000 samples was used to estimate the standard deviation of the PMF results.

**Tunnel pathway calculations.** Tunnel pathways were computed using the Caver \(^{39}\) software. The starting point coordinates were set to the centre of the binding pocket of opsin and A\(_2\)R. Computations were carried out using a shell radius 3 \( \AA \), shell depth 4 \( \AA \), a probe radius of 1.1 \( \AA \) for the opsin receptor and 1.4 \( \AA \) for the A\(_2\)R. Obtained results were plotted using the VMD software.

**Data availability.** The authors declare that all data necessary to support the findings of this study are available within the paper and its Supplementary Information Files. Additional data that were omitted from the paper are available from the corresponding authors upon reasonable request. Crystal structure coordinates from the Protein Data Bank (3EML, 3PWH, 4EIT) were used in this study.

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**Author contributions**

R.G. designed and performed the classical MD simulations with contribution from J.S., I.R.-E. and P.W.H. M.Man. and H.M.-S. performed the umbrella sampling simulations. H.M.-S. performed the free energy calculations. J.L.A. and M.Mar. performed cholesterol and MTSEA-B experiments. J.S. performed tunnel pathway calculations. P.W.H., M.P. and F.S. gave technical support and conceptual advice. R.G.G. and J.S. wrote the paper with contributions from J.-L.A., M.Mar., M.M.-S., H.M.-S. and comments from all authors.

**Additional information**

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