

# Predictions for Cholesterol Interaction Sites on the $\mathsf{A}_{\mathsf{2A}}$ Adenosine Receptor

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**Supporting Information** 

**ABSTRACT:** Molecular dynamics simulations of the  $A_{2A}$  adenosine receptor totaling 1.4  $\mu$ s show clear evidence for specific sites mediating interactions between adenosinebound  $A_{2A}$  and cholesterol. The strongest evidence is for three binding sites. Two are in the extracellular leaflet, with one site interacting with helices VII and I, and the other with helices II and III. One site is located in the intracellular leaflet, interacting with helices III and IV. One of our three predicted binding sites is confirmed by a just-published high-resolution structure of  $A_{2A}$  cocrystal-lized with an antagonist.

As our understanding of the complexity of the cell membrane grows, so too does the recognition that the membrane plays an active role in regulating integral membrane proteins (IMPs).<sup>1</sup> Cholesterol—present in fractions of up to 30 mol% in mammalian cell membranes—is a central figure in the regulation of IMPs. Cholesterol modulates the miscibility of membrane components<sup>2</sup> and the material properties of bilayers.<sup>3</sup> There is also crystallographic evidence in several cases for specific interactions between cholesterol and IMPs.<sup>4–8</sup> Generally, identification of such interactions is very challenging, but is essential in order to link such interactions with their biochemical consequences.

The G-protein-coupled receptors (GPCRs) are IMPs, and are the largest superfamily of proteins in the human genome, including roughly 800 members that transduce signals in every imaginable physiological context. GPCRs represent roughly half of all drug targets.<sup>9,10</sup> As integral membrane proteins, GPCR function is sensitive to the local environment of proteins and lipids. Cholesterol in particular is an indispensable component of cell membranes,<sup>11</sup> and is known to play an important role in GPCR structure and function.<sup>12–16</sup> Direct interactions between cholesterol and GPCRs have been observed by X-ray crystallography of the  $\beta_2$ -adrenergic receptor ( $\beta_2 AR$ )<sup>5,6</sup> and  $A_{2A}$ ,<sup>8</sup> and by electron microscopy<sup>4</sup> and molecular dynamics simulation of rhodopsin.<sup>17</sup> Specific sequences of amino acids that are thought to mediate cholesterol interactions have also been identified.<sup>6,18,31</sup>

Functional consequences dependent on cholesterol have also been reported. Bulk cholesterol has been shown to improve the stability of  $\beta_2 AR^{6,19}$  and to mediate receptor–receptor interaction as a necessary component for crystallization.<sup>5</sup> Ligand binding has been shown to be cholesterol dependent in intact cells for the oxytocin and cholecystokinin receptors.<sup>20</sup> Several recently published reports have focused on  $A_{2A}$ cholesterol interactions. Based on the  $\beta_2AR$  crystal structure,<sup>6</sup> Lyman et al. proposed that specific binding in the cleft between helices I and III stabilizes helix II in the apo state.<sup>21</sup> Nearly simultaneously, functional studies on micelle-solubilized  $A_{2A}$ showed that a precise titration of sterol derivatives is essential in order to recover native-like ligand binding.<sup>12,13</sup> Several publications have reported X-ray crystal structures of  $A_{2A}$  with<sup>8</sup> and without cholesterol,<sup>22–26</sup> providing a unique opportunity to test the predictive validity of molecular dynamics in the context of GPCR–lipid interactions.

We present blind predictions of preferred cholesterol interaction sites on  $A_{2A}$  by molecular dynamics simulations, one of which coincides with an interaction site identified by a very recently published X-ray structure.<sup>8</sup> The protein was simulated in a palmitoyloleoyl phosphatidylcholine (POPC) bilayer with 30 mol% cholesterol, with the initial positions of the cholesterols chosen at random. The cholesterols, by virtue of their diffusive wandering, identify the interaction sites in an unbiased and blind way. We report three specific interactions (shown in Figure 1) that are identified in both independent 800 ns trajectories. The data indicate that even cholesterols in well-



**Figure 1.** Trajectories of representative cholesterols projected onto the membrane plane. The trajectories connect the center of mass positions of six cholesterols in each simulation snapshot. The trajectories of three cholesterols at IS1, IS2, and IS3 (as marked) are shown in red. The trajectories of three other cholesterols are shown in green, blue, and magenta. In each trajectory, the initial position is shown as a large ball with the same color as the trajectory.  $A_{2A}$  is also shown, viewed from the extracellular side, with H1 (red), H2 (green), H3 (blue), H4 (purple), H5 (cyan), H6 (gray), and H7 (orange).

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defined interaction sites are quite mobile when associated with the protein, and for this reason we use the terminology "interaction site" rather than "binding site".

We simulated the adenosine-bound structure (PDB code 2YDO),<sup>24</sup> with four thermostabilizing mutations replaced by the native sequence, and residues not resolved in the crystal structure built as described in a recent publication.<sup>27</sup> The POPC+30 mol% Chol membrane was generated with the CharmmGUI.<sup>28</sup> The simulation protocol is identical to our recently published work;<sup>27</sup> simulations were performed with Desmond v. 30110<sup>29</sup> under conditions of semi-isotropic constant pressure and constant temperature by the Marty-nya–Tobias–Klein method.<sup>30</sup> Two independent trajectories of 800 ns each were obtained, requiring roughly 50 days of computation on 150 cores of our local cluster.

Figure 1 shows the trajectories of six cholesterols (out of a total of 110) over the course of the two independent simulations. Three cholesterols that interact most strongly are shown in red; the others are selected to represent the range of observed cholesterol mean-squared displacements (MSDs). Two of the selected cholesterols form no contacts with  $A_{2A}$  for the entire simulation time. A range of MSDs is observed for such non-interacting cholesterols, from relatively compact trajectories (blue) to wide-ranging (magenta). Transient interactions are also observed-the cholesterol shown in green makes contact with A2A for some time, but on the basis of the MSD we conclude that it does not form a specific interaction. (Taken in total, the trajectories of all cholesterols over both trajectories provide exhaustive sampling of the membrane-facing surface of the receptor, as shown in Figure S1.) On the other hand, three traces (shown in red) both have very compact trajectories and interact with the protein. These two criteria are our basis for the identification of specific interaction sites.

Figure 2 presents histograms of the averaged in-plane distances and root-mean-squared deviations (rmsd's) of the



Figure 2. Distribution of distance and rmsd. Histograms of (a) the averaged distance of cholesterol center of mass from  $A_{2A}$  and (b) the rmsd of the cholesterol centers of mass in the membrane plane. Solid (dashed) line corresponds to the first (second) trajectory.

positions for all cholesterols from  $A_{2A}$ . Cholesterols with both small distances and small rmsd's form contacts with  $A_{2A}$ , and those contacts remain stable for the duration of the simulation. In the first trajectory (solid line in Figure 2), there are seven cholesterols in the distance range of 15–25 Å (distance measured from the center of mass of  $A_{2A}$ ); of those seven, five have rmsd's in the range of 0–10 Å. In the second trajectory (dashed line in Figure 2), seven of the nine cholesterols with average distances of 15–25 Å also have rmsd's between 0 and 5 Å. In other words, we find five and seven cholesterols that have both small rmsd and interaction with the protein in the first and second trajectories, respectively. Among these, three interactions are with the same location on the protein. These reproducible, tight interactions we identify as cholesterol $-A_{2A}$  interaction sites.

Figure 3 shows the interaction sites common to both the trajectories, labeled IS1, IS2, and IS3, as well as three



**Figure 3.** Predicted cholesterol interaction sites. (a) Top view, cholesterols in the extracellular leaflet and (b) bottom view, cholesterols the intracellular leaflet. The images are structures at 250 ns, when all the cholesterols form contacts with  $A_{2A}$ . The coloring of the helices is the same as in Figure 1. The instantaneous cholesterol positions are shown by red (green) sticks for the first (second) trajectory. The cholesterols resolved in the recently published high-resolution structure of Liu et al.<sup>8</sup> are shown as orange sticks. Side views showing the positions of the cholesterols for each interaction site are shown in the bottom three panels.

cholesterols resolved in a very recently published high-resolution structure of  $A_{2A}$ .<sup>8</sup> IS1 is between H1 and H7 in the extracellular leaflet; exactly one cholesterol is found in each trajectory. IS2 is between H2 and H3, also in the extracellular leaflet, and coincides with the location of a cholesterol observed by X-ray crystallography.8 This site is occupied by two cholesterols in the first trajectory, and by a single cholesterol in the second trajectory. IS3 is between H3 and H4 in the intracellular leaflet. One cholesterol occupies this site in the first trajectory; two occupy it in the second trajectory. Our data also allow us to rank the interactions in terms of stability, by considering the rmsd and fraction of time each is in contact with the protein. The two cholesterols at IS1 have rmsd's of 4.5 and 5.5 Å, the three cholesterols at IS2 have rmsd's of 3.0, 4.5, and 5.4 Å, and the three cholesterols at IS3 have rmsd's of 2.4, 3.7, and 3.8 Å. (Below we compare the sites more quantitatively, on the basis of up-concentration relative to the average cholesterol concentration.) It is also straightforward to compute the fraction of time that each cholesterol interacts with the protein: (time with at least one contact with  $A_{2A}$ )/ (whole simulation time). Contact formation is defined by a distance between A<sub>2A</sub> and cholesterol of less than 4 Å. From

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this analysis, the two at IS1 are found to be in contact 70 and 79% of the time, the three at IS2 have 72, 83, and 99%, and the three at IS3 have 85, 96, and 100%. The interaction sites with smaller rmsd values naturally have longer interaction times, indicating that IS3 is the most stable, while IS2 and IS1 have similar stabilities.

A different view of  $A_{2A}$ -cholesterol interactions is obtained by mapping the density of cholesterol around the protein. As shown in Figure 4, there are clearly identifiable cholesterol



**Figure 4.** Density map of cholesterols on the membrane plane. The image is obtained by projecting the center of mass positions for all cholesterols onto the membrane midplane, and mapping the density by histogramming the center of mass positions on a fine mesh. The data are normalized by the value that would be obtained for a uniform distribution of cholesterols, corresponding to a value of 1 on the scale bar at right. Thus, at the extreme of the scale the density of cholesterol is ten times the density expected for a uniform, random distribution at 30 mol%. IS1, IS2, and IS3 are marked by arrows;  $A_{2A}$  is shown from the extracellular side.

"hotspots", several of which coincide with IS1, IS2, and IS3. IS1 and IS2 both correspond to a single localized density of cholesterol. The site we identify as IS3 appears to instead consist of three distinct but very tightly grouped interaction sites. Nonetheless, on the basis of the stability of the IS3 interaction, we identify it as a single interaction site. Several other hotspots are clearly visible in Figure 4, but are not identified as interaction sites by our criteria. Some fail the reproducibility test, being present in only one trajectory. Others are not the result of a single cholesterol, but rather two or more cholesterols that transiently visit the same interaction site. These cases fail the test of a combined low average distance and low rmsd of Figure 2. We also considered the convergence of the observed cholesterol data by computing the density map for nonoverlapping 100 ns trajectory segments, shown in Figure S2. Figure S2 demonstrates that the pattern of enhanced density at the interaction sites is fluctuating but stable by the second half of the trajectories, an indication that the simulations are sampling an equilibrium distribution of local cholesterol concentration.

A simple analysis of the observed cholesterol density at the interaction sites provides a means to estimate the free energy of association at each site, relative to the averaged cholesterol concentration in the bulk:

 $\Delta F_i = -RT \ln(\rho_i / \rho_0)$ 

Here, *i* labels the interaction site, *R* is the gas constant, *T* is the temperature in kelvins,  $\rho_i$  is the density of cholesterol at

interaction site *i* averaged over the last 400 ns of the simulations, and  $\rho_0$  is the averaged bulk density of cholesterol. For the three sites we obtain (in units of RT) -1.4 (IS1), -1.2 (IS2), and -1.8 (IS3). These values are consistent with the rmsd's that typify each site, and with the observation that these are relatively weak and dynamic interactions.

Our data predict interactions between particular residues on A<sub>2A</sub> and cholesterol. Taking into consideration both trajectories, we examine interactions between the two cholesterols at IS1 and the three cholesterols at IS2 and IS3 which are shown in Figure 3. We calculated the fraction of time that individual residues make contact with cholesterol, averaged over both trajectories. (A value of 1.0 means a residue has contact all the time, i.e., for the full 1.4  $\mu$ s of trajectories 1 and 2.) The two cholesterols at IS1 form contacts with the residues Val-8, Tyr-9, and Val-12 on H1 and Leu-272 on H7, with values ranging from 0.3 to 0.4. For the three cholesterols at IS2, the residues Val-57, Leu-58, Pro-61, Phe-62, and Thr-65 on H2 form contacts with values of 0.5 to 0.6, the residue Phe-70 on the extracellular loop 1 has a value of 0.5, and the residues on H3 (Phe-79, Ile-80, Phe-83) all have values less than 0.3. For the three cholesterols at IS3, the residues Phe-93, Ile-100, and Ile-104 on H3 and Ile-124 on H4 have ratios 0.6-0.7, while another group consisting of Leu-96, Ala-97, Tyr-103, and Arg-107 on H3, Cys-128 and Leu-131 on H4, Leu-192 on H5, and Leu-115 on intracellular loop 2 have values from 0.3 to 0.4. As the most stable interaction site as discussed earlier, IS3 forms many contacts that are formed for a significant fraction of the observation time.

To consider whether the observed sites are specific to cholesterol, we compared "fingerprints" of the cholesterol interactions reported here to interactions with oleoyl and palmitoyl chains in two control simulations totaling more than 2  $\mu$ s at low (1 mol%) cholesterol concentration. No reproducible, specific interactions are observed between either lipid chain and the protein at IS1 or IS3. A reproducible interaction between a lipid and the protein that is similar to IS2 is observed in the control simulations (see Figure S3). While that interaction is rather weak in the first of the two control trajectories, it is nonetheless possible that IS2 may promiscuously bind different components of the lipid matrix depending on the local membrane composition.

Overall, our data indicate that cholesterol interacts with specific locations on the membrane-facing surface of  $A_{2A}$ , but that interaction of even the most localized cholesterols is dynamic. The cholesterols do not adopt a rigidly defined orientation and position with respect to the protein, but there is rather some heterogeneity in their interaction with the protein. It is noteworthy that none of our strongest predictions correspond to the hypothesized interaction site between H1 and H3 in the intracellular leaflet, observed in an X-ray crystal structure<sup>6</sup> of  $\beta_2$ AR, expanded on the basis of evolutionary analysis,<sup>31</sup> and studied by molecular dynamics in the context of  $A_{2A}$ .<sup>21</sup> That interaction site is visited by several cholesterols in our simulations, but they do not linger, having rmsd's of 13 Å and above.

In summary, we observe reproducible, localized interaction sites between  $A_{2A}$  and cholesterol by long time scale molecular dynamics simulations. One of our interaction sites is confirmed by a recently published, high-resolution crystal structure of  $A_{2A}^{8}$  two others beg for experimental confirmation. However, we do not observe significant cholesterol density at a different site, also identified in the recent crystal structure, a contra-

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diction that will be the subject of further investigation. Though a growing body of work demonstrates that GPCR function is modulated by the membrane microenvironment, direct observations of GPCR cholesterol interactions are few. Identification of specifically interacting sites is essential in order to understand the thermodynamic forces that drive such associations and to rationalize their functional mechanism. Our data and previous work<sup>17</sup> demonstrate that molecular dynamics offers a straightforward way to identify specific lipid-protein interactions for GPCRs. Together with bioinformatic analysis,<sup>31</sup> MD offers a way to narrow the experimental focus to key areas on integral membrane proteins. Continued characterization of such interactions by experiment and simulation will deepen our understanding of the interplay of GPCRs and the lipid matrix. If our predictions are validated by further experimental work, molecular dynamics offers a valuable approach to studying specific lipid-GPCR interactions.

#### ASSOCIATED CONTENT

#### **S** Supporting Information

Figures S1–S3 described in the text. This material is available free of charge via the Internet at http://pubs.acs.org.

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#### Notes

The authors declare no competing financial interest.

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