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Karhu, Lasse

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Determinants of Orexin Receptor Binding and Activation—A Molecular Dynamics Study

Lasse Karhu,† Aniket Magarkar,‡ Alex Bunker,‡ and Henri Xhaard*,†

†Division of Pharmaceutical Chemistry and Technology, Faculty of Pharmacy, University of Helsinki, POB 56, FI-00014 Helsinki, Finland
‡Division of Pharmaceutical Biosciences, Faculty of Pharmacy, University of Helsinki, POB 56, FI-00014 Helsinki, Finland

Supporting Information

ABSTRACT: We assess the stability of two previously suggested binding modes for the neuropeptide orexin-A in the OX2 receptor through extensive molecular dynamics simulations. As the activation determinants of the receptor remain unknown, we simulated an unliganded receptor and two small-molecular ligands, the antagonist suvorexant and the agonist Nag26 for comparison. Each system was simulated in pure POPC membrane as well as in the 25% cholesterol−POPC membrane. In total, we carried out 36 μs of simulations. Through this set of simulations, we report a stable binding mode for the C-terminus of orexin-A. In addition, we suggest interactions that would promote orexin receptor activation, as well as others that would stabilize the inactive state.

INTRODUCTION

Orexin System. Peptide Binding. The orexin receptors are key regulators in several neurological processes. The main physiological role of the orexinergic system is the regulation of sleep, and it also plays a role in energy homeostasis and the reward system.¹ Orexin receptors are prominent drug targets for novel hypnotics,²,³ and hopefully, in the future, narcolepsy could be alleviated through orexin receptor activation.⁴,⁵ Orexin receptors are also putative targets for cancer treatment, as orexinergic signaling directs certain cancer cell lines to apoptosis.⁶ However, the treatment of both narcolepsy and cancer through orexin receptors would require agonistic ligands, whereas the pharmaceutical industry has concentrated on antagonists. A few potent small-molecular agonists have been reported and characterized,⁷–⁹ but the poor understanding of the receptor activation determinants hampers the search for new agonists.

The orexinergic system comprises two peptide ligands, orexin-A and -B, and two G protein-coupled receptors (GPCR) OX1 and OX2.¹⁰ In aqueous solution, the orexin peptides adopt conformations comprising helical segments I and II (and III for orexin-A), with a structural hinge motif between the helices I and II (Figure 1).¹¹–¹³ Orexin-B, which is a linear 28-amino-acid amidated peptide, shows a 90° bend between the helices I and II, at N20−A22, i.e., 1.5 turns from the C-terminal. Orexin-A, a 33-amino-acid amidated peptide with two intramolecular disulfide bridges, is often observed with a similar bend at A23−N25, half a turn further from the C-terminus, resulting in bending in the opposite direction in comparison to orexin-B (Figure 1). In addition to the bent conformations, orexin-A appears in some NMR-models in a straight α-helical conformation, except for the disulfide-bridge-stabilized hook at the N-terminus. For orexin-B, such conformation has not been reported.

Site-directed mutagenesis (SDM) on both receptor subtypes and both peptides has highlighted several key residues for...
peptide—receptor interaction. The most notable observations is that the C-terminus of both orexin peptides is vital for bioactivity, whereas the N-terminus can be mutated or even deleted without the loss of activity, which has given rise to the assumption that the C-terminus enters the canonical class A GPCR binding cavity. With this information, molecular modelers have provided few suggestions on the binding mode for the orexin peptides, with the assumption that the C-terminus binds in the α-helical conformation observed in solution, but validation of these suggestions has been challenging. Recently, we suggested that orexin peptides locked in an α-helical conformation can indeed activate the orexin receptors, but as the method included an introduction of a bulky hydrocarbon linker between two helical turns and severely impaired the peptide’s potency, no firm conclusions could be drawn.

**GPCR Activation. Simulations of GPCRs.** GPCR activation is a process where ligand binding stabilizes a receptor conformation that allows for G protein (or other effector protein) binding, activation and downstream signaling. As GPCR ligands differ greatly, also the key interactions at the binding site must differ. Among the class A GPCRs, the activation cascade that links events at the binding site with the intracellular G protein binding site appears to converge into defined interaction patterns, and large-scale movements such as the outswing of the TM6 (the sixth transmembrane helix), which opens up the G protein binding site.

Molecular dynamics (MD) simulations have shed light on different parts of the receptor function, such as ligand binding, receptor transitions both between and within active and inactive states, and allosteric modulation. The opening of the G protein binding site has not been observed in classical MD simulations reaching up to tens of microseconds, but the reverse event has been reported. In contrast, the binding site and the so-called core triad can adopt active-like conformations within a microsecond-scale simulation.

In the present study, we set out to simulate the OX2 receptor with two previously reported alternative orexin-A binding modes, a small molecular agonist Nag26 and a small molecular antagonist suvorexant. In addition, we simulated an unliganded receptor. Our main point of interest was the ligand binding interactions and their stability. As our simulations did not include G proteins or mimetics thereof, we anticipated that the transition of the receptor into its active state was likely beyond our reach, but we hoped to observe such differences between the ligands that could indicate key interactions for agonism.

## METHODS

**Overview.** We carried out and analyzed a total of 36 μs of molecular dynamics simulations. We simulated OX2 receptor without a ligand, with small molecules suvorexant and Nag26 and with two alternative binding modes of orexin-A. Each simulation was run both in a pure POPC (1-palmitoyl-2-oleoylphosphatidylycholine) membrane and in a POPC–cholesterol (25%) membrane for 3 μs. For Nag26 and orexin-A simulations, we also produced 1 μs replicas. The simulation conditions are summarized in the Table 1, and the simulations are referred to by their numbers thereafter. For the system setup and analysis we employed Gromacs 5.1, and for the MD simulations Gromacs 4.6.7.

**Amino Acid Numbering.** For clarity, we use one-letter amino acid codes for the orexin-A, and three-letter codes for the OX2 receptor. For the receptor, we use the GPCRdb numbering scheme, in which the most conserved amino acid in each transmembrane helix is denoted Na50, where N is the number of the respective helix. Amino acids up- and downstream are numbered consecutively based on their overlap with an ideal α helix; if a residue is “missing”, the corresponding number is skipped, and in the case of bulges, the residue furthest from the helical axis is denoted NαYY, where NαYY is the number for the previous residue.

**System Setup. The OX2 Receptor.** The OX2 receptor crystal (PDB id: 4S0V) is missing the N-terminus upstream of Pro50, residues 160–163 (the second intracellular loop, ICL2), 198–202 (top of the hairpin in the second extracellular loop, ECL2), 255–293 (ICL3), 336–337 (ECL3), and the C-terminus downstream of Cys381. Few side-chain atoms are also missing near the unresolved sections. To fill these gaps, we used Modeller 9.14 with default settings to build 30 homology models of the human OX2 (residues 50–254 and 294–381) using the crystal structure 4S0V as a template, and selected the model with the lowest RMSD in comparison to the template. We left out the receptor termini and the ICL3 since long stretches such as these do not produce reliable models without suitable templates, which were not available at the time. We set all arginine, lysine, glutamic acid, and aspartic acid residues as charged, except for the Asp102 which we set to be protonated. We allowed Gromacs to decide the protonation states for histidine residues based on the optimal hydrogen bonding conformation with default settings.

**Peptide Docking.** Orexin-A2–33 in the straight α-helical conformation14 was docked to the receptor model with ZDOCK35 and docking poses were refined with RDOCK36 via Discovery Studio 4.5. As pyroglutamate is not parameterized in these docking algorithms, we were forced to omit the first residue. It was added manually prior to simulation. From top-scoring docking poses, we selected manually two poses that were close matches to our previously published binding modes, which were named the TMS-mode and the TM7-mode based on the orientation of H26.

**Small Molecule Placement.** Suvorexant was copied from the OX2 crystal structure 4S0V. Nag26 was docked to the crystal structure as described previously. In short, Nag26 was docked with Glide induced-fit protocol, retaining the binding site water molecules. Shape and interaction similarities to
suvorexant directed the pose selection. The selected pose (but not the water molecules) was then copied into the binding cavity of the homology model with no further refinement.

**Membrane.** We employed CHARMM-GUI\textsuperscript{39--42} to generate two hexagonal membranes solvated in TIP3P\textsuperscript{43} water to result in a 12 nm total height. We used a pure POPC membrane of 150 lipids per leaflet, and a POPC + CHOL (25%) membrane of 126 + 42 lipids per leaflet. The membrane size was selected to ensure at least 50 Å between the periodic images of the protein in the membrane plane, and 25 Å in the direction perpendicular to the membrane. Both membrane models employed here are simplified representations of reality. For computational feasibility, we selected only one phospholipid for the study. POPC membranes are frequently used in MD simulations as the lipid is commonly present in biomembranes.\textsuperscript{44} The comparison of simulations, especially in the field of GPCRs, is facilitated by employing membranes similar to other studies.\textsuperscript{28,45--50} While most MD simulations with GPCRs employ single-component membranes, we additionally replicated all simulations with cholesterol-containing membranes to probe for the effects of the membrane composition. Cholesterol is known to be present in the cellular membrane, and its effects on the membrane thickness and packing are clear.\textsuperscript{51--54} Also, several interaction sites for cholesterol have been mapped onto the membrane-facing interface of GPCRs.\textsuperscript{55}

As the starting positions for the cholesterol in contact with the receptor can have a significant impact on their distribution through the simulation, we carried out coarse-grained MARTINI\textsuperscript{56} simulations to define suitable starting locations. We used CHARMM-GUI\textsuperscript{39--42,57} to embed the OX2 receptor in POPC–cholesterol bilayers with 10%, 25% or 50% of cholesterol, and to derive necessary topologies for the simulation. All systems were energy-minimized for 5000 steps and subsequently equilibrated for 100 ns (NVE ensemble). Then, each system was simulated for 10 μs in triplicate under NPT conditions. A time step of 20 fs was used in the simulations. Temperature of 310 K and pressure of 1.0 bar were maintained with V-rescale thermostat and Berendsen barostat with coupling times of 1.0 ps. Long-range electrostatics were calculated with the reaction field method and the van der Waals interaction was cut off at 1.1 nm. All backbone atoms of the receptor were position restrained with a force constant of 1000 kJ/mol × nm\textsuperscript{2}. The first 3 μs of each simulation were treated as equilibration, and the final 7 μs were subjected to analysis of cholesterol density around the protein. This revealed 8 sites for cholesterol molecules, and 15 sites for POPC molecules. These were back-mapped to an all-atom system.

**Construction of Simulation Systems.** We inserted the receptor complexes into the membranes with the Gromacs tool *membed* with default settings except for a 0.1 resize factor in the xy-dimension. To the POPC membrane, the protein complex was embedded as such; this resulted in a deletion of 3 + 3 POPC molecules, except for the peptide simulations, where the intracellular leaflet lost an additional lipid. For the POPC–cholesterol membrane, we first combined the receptor with the lipids identified by the coarse-grained simulation. Then, to ensure a symmetrical membrane with the intended 25% cholesterol content, we selected manually a suitable location for the insertion into the generated membrane and created a circular hole by deleting 18 POPC molecules per leaflet and 6 + 4 cholesterol molecules from extra- and intracellular leaflets respectively (as 5 out of 8 identified cholesterol were on the extracellular leaflet). The receptor and the close lipids were then inserted into the pregenerated hole.

In each system, we added NaCl to neutralize the charge for the receptor complex and to reach a salt concentration of 100 mmol/L. The resulting systems contained 293–294 POPC lipids or 246 POPC + 82 CHOL lipids, and ~25 000 water molecules in addition to necessary ions and the receptor complex.

**Force Field and Protocol.** We used Amber99sb-ildn force field\textsuperscript{58} together with Slipids\textsuperscript{59} parameters for POPC and cholesterol. We added parameters for pyroglutamate manually, and used atom charges from TINKER\textsuperscript{60} parameters. Suvorexant was parametrized with Antechamber,\textsuperscript{61} with Gaussian \textsuperscript{-derived RESP partial charges. Nag26 parameters were combined from Amber99sb-ildn (most bond and angle parameters), OPLS-AA\textsuperscript{63} (bonds and angles whenever native Amber parameters were unavailable, and torsions for the sulfonamide) and Antechamber (torsions excluding the sulfonamide, AM1-BCC charges\textsuperscript{64}). The topologies for the small molecules are available as part of the Supporting Information.

All simulations were run under periodic boundary conditions in hexagonal prism-shaped boxes. For all simulations, the time step was 2 fs, center-of-mass motion was removed every 10 steps for membrane–protein–ligand complex and solvent separately, and Verlet pair-list was updated every 20 steps. Temperature was kept at 310 K with Nose–Hoover thermostat separately for membrane, solvent and the protein complex, and pressure of 1 bar was maintained independently for z- and xy-dimensions with semi-isotropic Parrinello–Rahman barostat. Short-range electrostatics and van der Waals forces were cut off at 1 nm with long-range dispersion corrections for energy and pressure. PME was used for long-range electrostatics with cubic interpolation and Fourier spacing of 0.12 nm. LINCS constraints were used with all bonds.

**Equilibration and Production.** We used an equilibration scheme inspired by a previously presented work.\textsuperscript{65} Each simulation was equilibrated in seven 10 ns phases; in the first phase we imposed a 1000 kJ/mol restraint on protein and ligand heavy atoms, as well as on cholesterol heavy atoms for the simulations in the cholesterol-containing membrane. Through phases 2–5, we tapered off the heavy atom restraints with 200 kJ/mol intervals, except for the cholesterol, where the restraint was removed after phase 1. At phase 6, the 200 kJ/mol restraint was applied to all protein and peptide Cα, as well as to all heavy atoms in the small molecular ligands. At phase 7, the restraint was lifted from those Cα carbons that reside in the receptor loops. In addition to visual examination of the receptor, ligand and membrane, we monitored the equilibration through the stabilization of potential energy, simulation box vectors and receptor Cα-RMSD.

The equilibration was followed by 3 μs production runs. For the Nag26 and orexin-A simulations in both membranes, we started additional 1 μs replicas from the equilibrated systems.

**Analysis.** For analysis, each trajectory was aligned based on the receptor Cα with the Gromacs flag *progressive*; i.e., each frame was aligned to the previous frame to create continuous motions. Analysis was carried out on these full trajectories, except for the analysis involving water and calculation of the preferred locations for cholesterol around the receptor. For these, trimmed trajectories with frames every 3 ns were used.
The analysis of membrane thickness and area per lipid was carried out on unaligned coordinates of the final frame, as the receptor-based alignment tends to twist the membrane slightly out of the xy-plane.

The analysis of receptor and its interactions was conducted with Gromacs utilities in combination with VMD and Matlab. RMSD for protein and peptide was measured from the α carbons, and for the small molecules from the heavy atoms (i.e., non-hydrogen). We refer to RMSD as “external”, when the alignment was based on the receptor Cα, and “internal” when the alignment considered only the ligand Cα (peptides) or heavy atoms (small molecules). Thus, external RMSD contains ligand translation, rotation and conformational changes, whereas internal RMSD describes only the conformational stability. Interactions were mapped residuewise (or by atom groups in small molecules) with gmx mindist with a cutoff of 4 Å, disregarding hydrogens, and hydrogen bonds with the tool gmx hbond with default settings. Pairwise interaction energies were extracted with gmx enermap. Water-bridged hydrogen bonds were scanned with an in-house script by framewise inspection of multiple hydrogen-bonding partners to a single water molecule.

Stable water molecule locations, as well as preferred locations for cholesterol were mapped with VMD Volmap as density with default settings. Cholesterol density across the membrane was mapped using the gmx densmap with grid size of 1 Å. For the density analysis, we aligned the trajectories on the protein, but allowed rotation only around the z-axis to prevent the membrane from tilting, as this would have hindered the separated analysis of the membrane leaflets.

Order parameters for the POPC lipids were calculated with gmx order, and the double bond was treated as described by Pluhackova et al. The area per lipid and bilayer thickness were calculated with GridMAT-MD. For the analysis, the simulation box was surrounded by periodic images in the xy-plane, while calculations included only the lipids from the original box. The average membrane thickness was estimated as the area-weighted mean of the headgroup P–P distance at individual lipids.

\[
\text{thickness} = \sum Z_i \times \frac{A_i}{A_{\text{total}}}
\]

Z: head group P–P distance normal to the membrane plane

**RESULTS AND DISCUSSION**

**Membrane.** First, we analyzed the membranes to verify that our simulations replicate the expected membrane properties. At the end of the simulations, the pure POPC membranes showed an average area per lipid (APL) of 60.9 ± 0.8 Å², and membrane thickness of 36.9 ± 0.4 Å (see Supporting Information, Table S1). These values are in line with experimental data. The unsymmetrical insertion of some peptide–receptor complexes did not result in marked differences in APL. The cholesterol containing membranes showed clear effects of cholesterol condensing with POPC APL of 51.1 ± 0.6 Å² and membrane thickness of 41.4 ± 0.5 Å. Cholesterol APL was 37.5 ± 0.9 Å². The ordering parameters of the POPC tails closely replicate experimental values for both pure and cholesterol-containing membrane; see Supporting Information, Figure S1. Despite the clear effect of cholesterol on the membrane properties, we observe no consistent differences in ligand binding interactions or receptor mobility between pure POPC membrane and the cholesterol-containing membrane. This might reflect the orexin receptors’ indifference to the membrane composition, or more likely catching such effects would require more replicas or extended simulation periods.

The class A GPCRs have been postulated to feature specific cholesterol-binding sites facing the membrane milieu. On the extracellular leaflet, crystal structures of GPCRs have shown cocrystallized cholesterol molecules next to the TM3, at the TM4–5 interface and along the ECL3-side of the receptor, where the position at the TM6–7 interface has been most frequently populated. On the intracellular leaflet, the most notable site is the suggested “Cholesterol Consensus Motif” (CCM) located at the cleft formed by the TM1–4, which provides a binding site for one or two cholesterol molecules at least in the β2-adrenoceptor. As these sites offer a possible route for cholesterol-mediated effects, we sought to find and populate them through the coarse-grained simulations. The simulations suggested five sites on the extracellular leaflet (one facing the TM5, one facing the TM4–5 and one facing the TM6–7) and three sites on the intracellular leaflet (two next to the TM5, 1, and 2, and one facing the TM3–5) (Figure 2). Many of the sites discovered by the coarse-grained simulations match the cholesterol binding observed in the crystal structures; both sides of the extracellular end of the TM7, at the extracellular TM4–5 interface, and at the intracellular TM3–5 interface. Two cholesterol molecules were located close to the CCM site enclosed by the TMs 1–4, but not completely within the cavity. Of the starting sites, only the extracellular leaflet positions on both sides of the TM7 and next to the TM4–5 were stable through equilibration, and only the two positions flanking the TM7 were consistently stable in the production simulations (Figure 2, Supporting Information, Figure S2). Individual simulations show also other binding sites, most often in the extracellular leaflet, but these are not consistent across the pool of simulations (Supporting Information, Figure S2). Interestingly, the TM7-flanking sites, which were the most stable, are among those observed through X-ray crystallog-
Pro235 and Val142 remained in the starting (inactive) conformation. Val75, Trp345, Phe348, Trp351, and Tyr53 appeared to retain their closed, inactive conformation. RMSF showed that most of the core triad comprises Phe313, Phe346, Pro235, and Val142 remained in the starting (inactive) conformation, except for the simulations 7 (TM7-mode) and 10 (suvorexant). The observed reorganization was not replicated in other simulations with these ligands, and not reflected by the large-scale conformation of the TM6, which remained in the inactive conformation.

Receptor Conformation. GPCR activation is heralded by the outswing of the intracellular end of the TM6, which is accompanied by the reorganization of interhelical interactions at the intracellular side of the receptor. The interaction patterns have been analyzed for all GPCRs that have been crystalized both in the active and the inactive state. The analysis showed that there are five conserved "inactive" interactions, which in OX1 would be Ile148-Leu306, Val75-Tyr364, Tyr364-Phe371, and Asn365-Arg372. Similarly, there are two conserved "active" interactions, which would be Leu310-Val240 and Tyr364-Lle148 in the OX2. In addition to these interactions, which locate at the intracellular side of the receptor, a triad of amino acids at the center of the receptor has been suggested to reorganize upon receptor activation.

Peptide Binding Modes. One of the aims of our study was to assess the stability of the two alternative orexin-A binding modes we had previously suggested, called the TM5-mode and the TM7-mode based on the orientation of the (arbitrarily selected) H26. In the previous study, neither binding mode was promoted by the assessment of docking scores and interactions toward residues that had been shown important by SDM.

While the 19-amino-acid fragment from the orexin-A C-terminus has been shown to suffice for the biological activity, we elected to use a full-length peptide, which we docked in the straight α-helical conformation to replicate our previous docking poses. The simulations, however, showed that this straight conformation was not stable unless hydrogen bonding between the N-terminal hook and the receptor ECL2 took place. These hydrogen bonds often involved the ECL2 main chain, and were not consistent across the simulations. In absence of hydrogen bonding between the orexin-A N-terminus and the receptor ECL2, the peptide hinge was allowed to bend, moving the peptide N-terminus to rest horizontally atop the receptor. This bending took place in the simulations 2 and 5, which feature the TM5- and TM7-modes, respectively, while other simulations with orexin-A showed the N-terminus in contact with the ECL2, following its fluctuations (Figure 3). Our simulations did not include the receptor N-terminus, as its structure and location had not been resolved at the time. The possible impact of the receptor N-terminal helix on the peptide conformation and binding is discussed below, after the description of the peptide C-terminus binding.
As the C-terminus of the peptide is the key for biological activity, we focused in its stability and interactions. To assess the stability of the predicted binding modes, we analyzed the hinge region and the conformational peptide C-terminus (N25−L33) separately, since the bending of the hinge region and the conformational flexibility of the N-terminal hook dominate the RMSD of the full-length peptide. Overall, the receptor-bound C-terminus of orexin-A maintained its conformation and location better in the TM5-mode (i.e., H26 side chain points toward TM5) than in the TM7-mode, and for both binding modes, the conformation was more stable in the pure POPC membrane than in the cholesterol-containing membrane (Figure 4).

As the binding interaction of the peptide N-terminus are heavily influenced by the (possible) bending of the peptide, we assessed the binding interactions only for the peptide C-terminus (N25−L33). These interactions, averaged across the simulations, are shown in Figure 5, panels A and B for the TM5- and TM7-mode, respectively. Simulationwise heatmaps of the interaction frequencies, along with mapped interaction energies are available in the Supporting Information as Figures S5−S12. The stable location and conformation of the peptide C-terminus in the TM5-mode is mirrored in well-defined, stable interactions across simulations 1−4, whereas the higher mobility and instability of the TM7-mode is reflected by more variable interactions in the simulations 5−8.

As described, the orexin-A C-terminus remained quite stable in the cholesterol-containing membrane (POPC membrane), or via a water-mediated hydrogen bond due to the increased distance in the simulation 3 and 4 (cholesterol membrane). H26 interacted with the side chains of the acidic residues in the ECL2 (Asp21145x51 and Glu21245x52) in the simulations 1−4, either directly as seen in the simulation 4, and (B) 3−4, (C) 5−6, and (D) 7−8. See Methods, Analysis, for definitions for the external and internal RMSD.

![Figure 4. RMSD of the peptide C-terminus. In green and orange, the “external” RMSD, and in blue and red, the “internal” RMSD. (A) Simulations 1−2, (B) 3−4, (C) 5−6, and (D) 7−8.](image-url)
Phe227\textsuperscript{3443}. The NH\textsubscript{2} of the amidated C-terminus mainly bonded to the G29 or the I30 carbonyl. An interesting network of interactions took place at the bottom of the binding cavity, between the exposed carbonyls of L31 and T32, along with Gln134\textsuperscript{3432} and Tyr317\textsuperscript{6448}. In the beginning of the simulations, Gln134\textsuperscript{3432} side-chain nitrogen usually formed a hydrogen bond to the carbonyl of T32, while Tyr317\textsuperscript{6448} hydroxyl pointed down. This was most likely imposed by the starting configuration. During the simulations, the Gln134\textsuperscript{3432} hydrogen bond shifted to the carbonyl of L31, Tyr317\textsuperscript{6448} hydroxyl flipped upward, and a water molecule moved in to bridge interactions between the Gln134\textsuperscript{3432} side-chain carbonyl, 

![Figure 5. Heatmap of binding interactions of the peptide C-terminus, averaged across all simulations sharing the same binding mode (1–4 for TM5-mode, 5–8 for TM7-mode). The heatmaps display the fraction of simulation time that any interatomic distance was below 4 Å. H:xx denotes the presence of a hydrogen bond and W:xx the presence of a water-mediated hydrogen bond. These are shown only when the frequency was over 20%.](image)

![Figure 6. Orexin-A binding interactions in the TM5-mode. (A) Binding site with close receptor residues shown as sticks. (B) Hydrogen bonds between the peptide C-terminus and the receptor. The atomic coordinates for the TM5-mode receptor complex presented here are available as Supporting Information.](image)
Tyr317-648 hydroxyl and the T32 carbonyl. The Gln332-L31 interaction was also prominent in terms of interaction energy (Figures S5–S8 in Supporting Information). In the simulation 2, this reorganization did not happen, as Tyr317-648 hydroxyl immediately bound to the Gln134-332 side-chain carbonyl, leaving no space for the bridging water molecule. While the hydroxyl of the Tyr354-742 was consistently close to the carbonyl of L31, it participated in the network only transiently, preferring to bind with Thr111-2466; this interaction was almost exclusively seen only in the TM5-mode (simulations 1–4). The carbonyl of L33 also took part only rarely.

The instability of the orexin-A helix in the TM7-mode rendered it difficult to sum up the interactions across the simulations 5–8. The simulation 7 showed an almost complete “melting” of the peptide C-terminus, and in simulation 6 and 8 the hinge region unfolded and fell into the cavity between the TMs 2 and 7. The common observations between the simulations were that N25 was close to Asp211-451 and Cys210-450, I30 inhabited a pocket lined by Ile320-651, Asn324-655, Lys327-658, and Phe346-734, and L33 was close to His350-738, often hydrogen bonded to the C-terminal amide. In terms of interaction energy, there was no apparent consistency between the simulations (Supporting Information, Figures S9–S12). The simulation 7 with the most prominent peptide unfolding aside, the three other simulations (5, 6, and 8) showed the side chain of A27 facing a salt bridge formed by Glu212-452 and Arg328-659, A28 residing under the ECL2 and the side chain of L31 facing the pocket of Pro131-329, Thr135-333, Gln187-461, and Met191-465. The C-terminal carbonyls did not form uniform interactions such as those described above for the TM5-mode.

Given the difference in stability, in terms of both peptide conformation and binding interactions, between the TM5- and the TM7-mode of binding, the TM7-mode seems less likely to be the correct binding mode. Therefore, if the C-terminus of the orexin-A indeed binds in an α-helical conformation, we suggest the TM5-mode as the binding mode for orexin-A at the OX2. The binding mode should also be valid for the OX1 receptor and for orexin-B, since the orexin peptide C-terminus is conserved and the receptor structures are similar. However, with current data, we cannot rule out the possibility of nonhelical bioactive conformation. Endothelin, for example, adopts a conformation that is mostly helical but the C-terminus is extended into the receptor cavity.75 While it remains unclear whether these different locations represent different functional states or if either one is a crystallization artifact, the N-terminal helix was shown, through mutagenesis, to be important in the peptide-induced receptor activation.73 This suggests direct interaction with the peptide, which in turn would promote the location next to the ECL2. If the N-terminal helix would indeed lie atop the receptor in interaction with ECL2, as seen in the OX1 structure,73 it would prevent the binding of orexin-A in the straight α-helical conformation such as the one we used for docking and as the starting conformation for the simulations. However, the peptide bent toward the TMS would fit snugly underneath the N-terminal helix, presenting R15 and Y17 among others toward the downward-facing polar residues of the N-terminal helix (Figure 7). Interestingly though, the N-terminal helix in the crystal structure features lipophilic residues (Phe31, Leu32, Tyr34, Leu35, and Trp36) facing toward the solvent, which seems unfavorable. A roll of 180° would bring them in contact with L16, Y17 and L20 in the orexin-A. It is also possible that the N-terminal helix adopts a distinct peptide-bound

![Image](https://example.com/image.png)

**Figure 7.** Simulation-derived bent conformation of orexin-A overlaid with the recently published OX1 structure, which displays the N-terminal receptor helix.
conformation, which differs from the small-molecule-bound structures.

Small Molecule Binding. In addition to orexin-A, we carried out simulations with bound small molecules, the antagonist suvorexant (simulations 9−10) and the agonist Nag26 (simulations 11−14). Suvorexant binding mode seen in the crystal structures was stable through the simulations in both membranes. The external RMSD fluctuated around 1 Å, and the internal RMSD between 0.5−1 Å (Figure 8). The chlorinated benzoxazole remained in the pocket lined by Cys107, Ala110, Thr111, Val114, Trp120, Ile130, Pro131, and Gln134, with the oxazole ring...
packing next to Pro131\textsuperscript{329} (Figure 9A). A water molecule was often seen binding with the oxazole nitrogen, but it did not form a stable water-mediated interaction. The seven-membered ring occupies a central location with the methyl facing Phe227\textsuperscript{543}. The linker carbonyl hydrogen bonds to Asn324\textsuperscript{645} and via water to His350\textsuperscript{748}. The methylated benzene ring interacts with His350\textsuperscript{748}, Val353\textsuperscript{741}, and Tyr354\textsuperscript{742} in TM7, and is flanked by Gln134\textsuperscript{332} and the benzanazole ring on the other side. Underneath, the Gln134\textsuperscript{332}–Tyr354\textsuperscript{742} hydrogen bond is maintained throughout the simulations, which is not seen consistently in any other simulations. The triazole ring is next to Ile320\textsuperscript{51} and Phe227\textsuperscript{543}, and traps 2–3 water molecules into a pocket between the TMs 5 and 6, where the water molecules form a hydrogen-bond network between Ser321\textsuperscript{642}, Thr231\textsuperscript{546}, and His224\textsuperscript{540}.

Concerning the pool of crystallized orexin receptor antagonists, the binding mode of suvorexant is near identical between the OX1 and OX2 subtypes,\textsuperscript{32,73} and closely replicated out the simulations, which is not seen consistently in any other simulations. The triazole ring is next to Ile320\textsuperscript{51} and Phe227\textsuperscript{543}, and trapping of water molecules into a pocket between the TMs 5 and 6 is where the water molecules form a hydrogen-bond network between Ser321\textsuperscript{642}, Thr231\textsuperscript{546}, and His224\textsuperscript{540}.

Figure 10. Small molecular agonist Nag26 was mobile during the simulations, rendering the binding mode identification difficult. Representatives of the three most visited conformations are shown.

The simulations showed distinct preferences for water molecule locations. Within the receptor core, Asp100\textsuperscript{2a50} was often accompanied by few water molecules (Figure 11). In some simulations, a semistable chain of water molecules connected these waters to the binding cavity. Two to three water molecules usually resided between the TMs 5 and 6, bound to Thr231\textsuperscript{546} and Ser321\textsuperscript{642}, sometimes also to His224\textsuperscript{540} and Asn324\textsuperscript{645}. Often, the simulations 11 and 12, the group rose a bit and then pushed in-between the TMs 4 and 5. This rise brought the amide oxygen in water-mediated contact with Asn324\textsuperscript{645} or Arg328\textsuperscript{659}. The push also moved Thr231\textsuperscript{546} and Phe227\textsuperscript{543} toward the TM4, resulting in a marked counterclockwise rotation of the upper section of the TM5 and tightening of the helical turn at the bulge near Tyr232\textsuperscript{547}. The sulfonamide moiety, which is crucial for activity, lay next to Pro131\textsuperscript{329} in all simulations but found different binding partners. In the simulation 13, where the Nag26 conformation was the most stable, the sulfonamide formed both direct and water-bridged hydrogen bonds from the oxygens to Gln134\textsuperscript{332}. Despite close proximity, Gln189\textsuperscript{461} and Thr135\textsuperscript{333} did not hydrogen bond with the sulfonamide. The amide nitrogen formed a stable water-bridge to Glu212\textsuperscript{545}, which in turn remained bonded to Arg328\textsuperscript{659} through the simulation. The simulation 14 showed a more mobile sulfonamide moiety, which hydrogen-bonded first with Arg328\textsuperscript{659} and then moved higher and toward the center of the binding site and switched to Lys327\textsuperscript{648}. This movement also allowed the amide nitrogen to bond with Asp211\textsuperscript{451}. Arg328\textsuperscript{659} and Glu212\textsuperscript{545} remained in interaction through the simulation, but the ligand movement moved both side chains toward the TMs and appeared to constrict the entrance of the binding site. Simulations 12 and 14 showed Arg328\textsuperscript{659} hydrogen bonding with the sulfonamide and Glu212\textsuperscript{451}/Asp211\textsuperscript{451}, while the amide nitrogen interacted with Cys210\textsuperscript{450}’s main-chain oxygen via water. Water also bridged transient interactions between the sulfonamide oxygens and the side chains of Asp211\textsuperscript{451} and Glu212\textsuperscript{451}. The hydrogen bonding between the sulfonamide nitrogen and Lys327\textsuperscript{648}/Arg328\textsuperscript{659} seen in simulations 11, 12 and 14 was also a strong contributor to the interaction energy (Supporting Information, Figures S13, S14, and S16). The rest of the molecule was quite mobile; if the ligand retained or readopted the packed conformation, the methylated ring folded in-between the TM7 and the rest of the ligand, favoring interactions with Phe346\textsuperscript{734}. In the case of ligand extension, the ring moved close to the ECL1, which brought the polar atoms of the chain linker in hydrogen-bonding distance of the exposed carbonyl of Cys210\textsuperscript{450}, and also Thr111\textsuperscript{260}.

It is noteworthy that we employed only one starting conformation for the Nag26 agonist. As also our peptide simulations highlighted, the starting conformation can heavily impact the simulation trajectory, and it is possible that another starting conformation for the Nag26 would have yielded different results. Then again, our simulations showed the Nag26 sampling multiple conformations within the binding site, unable to find a markedly stable binding mode. This could mirror the weaker binding interactions in comparison to suvorexant; the Nag26 $K_i$ is 140 nM at the OX2, while the corresponding value for suvorexant is 0.35 nM.\textsuperscript{2,7} Alternatively, the initial conformation might be far from correct, but given the resemblance between our docking pose and the structurally related small molecular antagonist EMPA (not shown), that seems unlikely.

**Water Molecules**. The simulations showed distinct preferences for water molecule locations. Within the receptor core, Asp100\textsuperscript{2a50} was often accompanied by few water molecules (Figure 11). In some simulations, a semistable chain of water molecules connected these waters to the binding cavity. Two to three water molecules usually resided between the TMs 5 and 6, bound to Thr231\textsuperscript{546} and Ser321\textsuperscript{642}, sometimes also to His224\textsuperscript{540} and Asn324\textsuperscript{645}. Often, the...
simulations showed also stable water molecules that took part in the ligand–receptor interactions; these were discussed in detail above for each complex.

Many of the observed stable water molecules are also present in the high-resolution crystal structures, such as between the TMs 5 and 6 (Thr231 and Ser321), around Asp100, at the TM6 kink and atop the TM6, and in the intracellular cavity between TMs 3, 4, and 5. Water molecules were also seen around the ECL2 in matching positions to the crystal structures, and close to the TMs 5 and 6. Interestingly, the water molecule that takes part in suvorexant binding, bridging the carbonyl oxygen to His350, was also stable in simulation (orexin-A in the TM5-mode), where the water molecule bridged a three-way interaction between Asn324, His350 and the main-chain carbonyl of N25. In simulation 13, the carbonyl oxygen in the linker chain of the Nag26 occupied a similar position. The water molecules between the TMs 5 and 6 provide the most interesting case; there were two-to-three water molecules in all simulations, unless the TM5 made the counterclockwise turn, moving Thr231 further from Ser321 and thus breaking the water-binding pocket. With the Nag26 simulations, this was linked with the dimethylamide pushing between the TMs 4 and 5, but in the simulation 2 (orexin-A in the TM5-mode), this turning appeared to take place without clearly defined ligand-dependency.

**Activation.** The activation of a GPCR is a chain of events propagating through the receptor from the ligand-binding site to the G-protein-binding site at the intracellular face. In our simulations, the intracellular conformation of the receptor remained in the inactive state, as anticipated, which was mirrored by the absence of the common contacts observed in the structures of activated GPCRs. Concerning the transitional steps along the activation cascade, the core triad, which lies underneath the binding site and is thought to be relevant in relaying the activation signal, also remains in the initial, inactive conformation. It is noteworthy that orexin receptors have a valine residue at 3x40, whereas a bulkier residue such as isoleucine is often found in the receptors that display a clear “activation” of the core triad. The M2 receptor also features Val3x40, but there both active and inactive receptor structure resemble the activated core triad of e.g. β2 receptor. At the binding site, the activation determinants differ between receptors, rendering comparison of orexin receptors to the available structures of active GPCRs dubious. However, a common theme in activating interactions is a change in interhelical interactions across the binding site. With this in mind, we noted that suvorexant simulations showed a consistent hydrogen bond between Gln134 and Tyr354, which was not seen in other simulations, but is replicated by other crystallized antagonists. In contrast, the simulations with orexin-A in the TM5-mode showed this hydrogen bond only 0–8% of the simulation time. On the other hand, the TM5-mode propagated Tyr354–Thr111 hydrogen bonding. We speculate that enforcing the Gln134–Tyr354 hydrogen bond could stabilize the inactive state of the receptor, and that the alternative Tyr354–Thr111 hydrogen bond could enable receptor activation, even though Nag26 did not show as strong effect as the orexin peptide did. The comparison between suvorexant and Nag26 highlights the agonist interacting with Thr231, a location which is central to the activation of adrenoceptors, for example. The Nag26 also interacted with the ECL2, and the salt bridge Glu212–Arg328, which crosses the
binding site. One could speculate that this interaction with the salt bridge could induce an inward tilt of the extracellular end of the TM6, which, in turn, could pivot the TM6 and open the intracellular binding site. Nag26 was also seen in two simulations pushing between the TMs 4 and 5, which induced a tightening in the TM5 helix and changed the interactions between the TMs 4, 5, and 6. This could also serve in destabilizing the inactive receptor conformation, thus facilitating receptor activation.

**CONCLUSIONS**

We performed extensive molecular dynamics simulations on the OX2 receptor. On the basis of the simulations, we suggest that orexin-A binds the receptor in a configuration where N25 faces the TM7 (Tyr343[32]), H26 points toward the TM5 but interacts either with the ECL2 or the ECL3, A27 faces the ECL1, A28 and G29 pack next to His350[34,35] I30 and L31 flank the TM3 and Pro131[36,37] with the L31 side chain occupying a pocket beneath the ICL1, T32 faces the TM7 and L33 the TMs 5 and 6. The exposed carbonyls in the last helical end of the TM5 into a counterclockwise rotation, which stabilizes the inactive receptor conformation, thus facilitating receptor activation.

**ASSOCIATED CONTENT**

* Supporting Information
  The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcb.8b10220.

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  Alex Bunker: 0000-0002-1236-9513
  Henri Xhaard: 0000-0002-3000-7858
  *Author Contributions*
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**ABBREVIATIONS**

GPCR, G protein-coupled receptor; POPC, 1-palmitoyl-2-oleoylphosphatidylcholine; SDM, site-directed mutagenesis; MD, molecular dynamics; ECL1–3, extracellular loop 1–3; ICL1–3, intracellular loop 1–3; APJ, area per lipid; TM, transmembrane helix; CCM, cholesterol consensus motif.


