

X-Ray Crystallography and Free Energy Calculations Reveal the Binding Mechanism of A_{2A} Adenosine Receptor Antagonists.

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Abstract: We present a robust protocol based on iterations of free energy perturbation (FEP) calculations, chemical synthesis, biophysical mapping and X-ray crystallography to reveal the binding mode of an antagonist series to the adenosine A_{2A} receptor (AR). Eight A_{2A}AR binding site mutations from biophysical mapping experiments were initially analysed with sidechain FEP simulations, performed on alternate binding modes. The results distinctively supported one binding mode, which was subsequently used to design new chromone derivatives. Their affinities for the A_{2A}AR were experimentally determined and investigated through a cycle of ligand-FEP calculations, validating the binding orientation of the different chemical substituents proposed. Subsequent X-ray crystallography of the A_{2A}AR with a low and a high affinity chromone derivative confirmed the predicted binding orientation. The new molecules and structures here reported were driven by free energy calculations, and provide new insights on antagonist binding to the A_{2A}AR, an emerging target in immuno-oncology.

Introduction

Computational estimation of shifts in binding free energy, associated with ligand modifications or point mutations in the receptor macromolecule, can provide the missing link between the structure of a protein-ligand complex and a panel of experimental binding affinities. Rigorous free energy perturbation (FEP) methods have been used for decades to understand the structure-affinity relationships (SAR) around a given chemical scaffold, and recent advances now allow the use of this technique routinely in ligand design projects.^[1] The same methodology can be used to analyze this problem from a complementary perspective, that is to estimate the gain or loss in binding free

energy from site-directed mutagenesis (SDM) data. The idea of in silico mutagenesis, initially introduced almost three decades ago by Kollman to study the binding and catalysis of subtilisin,^[2] was recently implemented in computational pipelines that pursue a systematic characterization of the effect on point mutations on e.g. ligand-binding or protein stability.^[3-5] The combination of both ligand and residue FEP simulations can provide a full energetic landscape of the molecular interactions governing protein-ligand binding, which underlies the design of two complementary protocols in our lab, namely QligFEP^[6] and QresFEP^[7], integrated in the molecular dynamics (MD) software package Q.^[8,9]

One area where this approach is particularly promising is the design of ligands for G-protein-coupled receptors (GPCRs), a superfamily of seven-transmembrane (7TM) cellular receptors^[10] that mediate the therapeutic effects of about 30% of all marketed drugs.^[11] There is a large amount of SAR and SDM data available for these receptors, which can be combined with the increasing growth of structural knowledge of many GPCR targets. The first integrated approach of ligand and residue FEP simulations was published by Boukharta et al. to characterize antagonist binding to the Y₁ neuropeptide receptor,^[12] which we later expanded to other GPCR families, including the related neuropeptide receptor Y₂,^[13] the orphan receptor GPR139^[14] and several members of the family of adenosine receptors.^[15-17] Among the last, the adenosine A_{2A} receptor (A_{2A}AR) was one of the first GPCRs to be crystallized^[18] and today stands out as one of the better characterized GPCRs from a structural perspective. Several structures of the inactive and active forms of the receptor have been solved within the last decade, and the integration of the available experimental data has strongly aided ligand design programs for this receptor.^[19,20]

Many A_{2A}AR antagonists have been developed targeting a number of pathologies,^[21-23] including recent clinical candidates

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in immuno-oncology. A number of these antagonists have been co-crystallized with the $A_{2A}AR$, providing unique structural information which, in combination with the extensive SDM data available,^[24] allow envisaging ligand binding mechanisms and structure-based drug design (SBDD) programs of antagonist molecules.^[19] However, suboptimal properties of traditional scaffolds, such as poor pharmacokinetics and low selectivity profiles, motivate the search of novel chemical entities as $A_{2A}AR$ antagonists,^[25] frequently through high throughput screening (HTS) campaigns. In these cases, it is not common to obtain a crystal structure of the receptor-ligand complex, which can hamper further hit to lead optimization. Instead, approximate binding modes are often inferred from the experimental data extracted from (SAR) of ligand series and SDM data,^[24] which can be complemented by computational models of the protein-ligand complex.^[26]

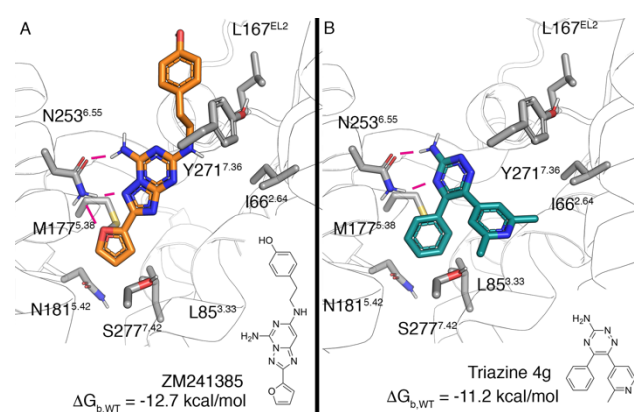


Figure 1. Binding mode and chemical structures of antagonists **ZM241385** (A) and **triazine 4b** (B), both co-crystallized with the $A_{2A}AR$ (ribbons, corresponding to the 4EIY structure)^[27] and characterized by BPM (residues labeled and depicted in gray sticks). Receptor-ligand hydrogen bonds are depicted as magenta lines.

Biophysical Mapping (BPM), is an integrated approach that has been used with success in antagonist design programs on the $A_{2A}AR$ and other GPCRs.^[28–31] Here, the binding affinity of a ligand series is evaluated via surface plasmon resonance (SPR) on a panel of mutant receptors, each bearing a single-point mutation within the putative binding site.^[28] The resulting matrix of binding affinity shifts from wild type (WT) affinities combined with SAR data and mapped to a receptor-ligand model provide further insights in the determinants of binding of the scaffold. The first application of this technique was based around the co-crystallized $A_{2A}AR$ antagonist **ZM241385** in combination with 8 receptor mutants (see Figure 1A).²³ These mutations involve residues in direct contact with the ligand, such as N253^{6.55}, L85^{3.33}, M177^{5.38}, N181^{5.42} and I66^{2.64} (Ballesteros Weinstein numbering^[32] in superscript), as well as residues not directly in contact with the ligand, namely S277^{7.42}, Y271^{7.36} and L167^{EL2} (see Figure 1). This approach was extended for 1,2,4 triazines as $A_{2A}AR$ antagonists,^[33] and the binding mode was later confirmed by X-ray crystallography (see Figure 1B).^[34] Finally, a series of chromones were identified by HTS as a novel family of $A_{2A}AR$ antagonists, and consecutively optimized to yield the potent and selective **Chromone 14** (see Figure 2). This optimization was guided by the interpretation of the BPM data and a computational model of the complex generated by docking. During this

optimization, two putative binding modes could explain the BPM data, although the SAR of the generated series seemed to favor one of them. However, this could not be validated by X-ray crystallography in the original study.^[31,35]

Here, we initially examine the SPR data available for these $A_{2A}AR$ antagonist families through a recently developed *in silico* mutagenesis tool based on free energy perturbation (FEP) simulations.^[12,15,16] The results led to the proposal of a binding mode of the chromone series, which is used as a basis to design, synthesize and pharmacologically evaluate of an extended series of compounds, confirming the SAR trend underlying their design. Finally, experimental structures of two chromone- $A_{2A}AR$ complexes were solved which confirmed the binding mode hypothesis from the computational studies.

Results and Discussion

Free Energy Perturbation calculations on existing BPM data.

The BPM data obtained for the three chemotypes of $A_{2A}AR$ antagonists were collected, and relative binding free energy changes between mutant and WT receptor were calculated from K_i values (see Table 1, $\Delta\Delta G_{bind}^{exp}$). Thereafter, relative binding free energies for each ligand were calculated based on 3D models of the receptor-ligand complexes ($\Delta\Delta G_{bind}^{calc}$, Table 1). In the case of **ZM241385**, the model was directly extracted from the high-resolution structure with the $A_{2A}AR$ (see Figure 1), and the curated receptor model (see SI Methods) was used throughout this work. The results show excellent agreement with the experimental data, with a mean absolute error (MAE) of 0.41 kcal/mol and a correlation coefficient of $R^2 = 0.94$ (Table 1). Next, the same set of BPM mutations was analyzed for **triazine 4g**. In this case, the starting configuration of the complex was obtained by aligning the **triazine 4g**- $A_{2A}AR$ complex (3UZA) to the curated model described before, retaining only the coordinates of the ligand (Figure 1B).^[36] The calculated free energies ($\Delta\Delta G_{bind}^{calc}$, Table 1) were in general good agreement with the experimental data (MAE = 0.94 kcal/mol; $R^2 = 0.66$). In contrast with the previous case, two mutations (Met177^{5.38} and Asn181^{5.42}) show qualitative discrepancy with the experimental data for this ligand. Subsequent analysis of the MD trajectories suggests that this is likely is due to a suboptimal representation of the (water mediated) H-bond network upon mutation, which has previously been recognized as a challenging factor for this particular scaffold.^[37,38]

Once the QresFEP protocol was validated to reproduce the BPM data on experimentally known structures, we moved on to the chromone scaffold, for which there is analogous BPM data, but no crystal structure available. Here, two reasonable binding modes were generated by molecular docking of **chromone 14**. The two poses, denoted as A and B (see Figure 2), form at least one hydrogen bond with Asn253^{6.55} (a highly conserved interaction in AR ligand recognition)^[24] and are related by a symmetry axis along the bicyclic core of the chromone scaffold. According to the docking score function used, both poses were energetically equivalent (-9.30 vs -9.17 for pose A and B respectively) making it difficult at this point to discern the correct one, in line with the previous binding hypothesis from Andrews et al.^[31,35]

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Table 1. Comparison between experimental and calculated relative binding free energies ($\Delta\Delta G_{bind}$, in kcal/mol) for $A_{2A}AR$ mutants

Mutant	ZM241385		Triazine 4g	
	$\Delta\Delta G_{bind}^{exp}$	$\Delta\Delta G_{bind}^{calc}$	$\Delta\Delta G_{bind}^{exp}$	$\Delta\Delta G_{bind}^{calc}$
I66A ^{2,64}	0.14	0.83 ± 0.34	0.41	1.94 ± 0.34
L85A ^{3,33}	2.45	3.30 ± 0.41	1.09	1.65 ± 0.37
L167A ^{5,28}	0.00	0.60 ± 0.31	-0.14	-0.39 ± 0.36
M177A ^{5,38}	0.14	-0.09 ± 0.44	-0.27	1.66 ± 0.49
N181A ^{5,42}	1.23	1.47 ± 0.57	0.82	-0.63 ± 0.55
N253A ^{6,55}	≥ 5.86 ^c	5.81 ± 0.57	≥ 4.36 ^c	5.64 ± 0.56
Y271A ^{7,36}	1.09	0.84 ± 0.74	0.41	-0.1 ± 0.68

[a] Experimental relative binding free energies were calculated from K_i values as $\Delta\Delta G_{bind}^{exp} = RT \ln \left(\frac{K_i^{mut}}{K_i^{WT}} \right)$ with experimental errors in all cases reported as approximately 0.1 pKD unit, i.e. less than 0.1 kcal/mol.^[39] [b] Data for the mutant receptor constructs reported in^[39]. [c] Binding affinity of the ligand to the (mutant) receptor was lower than the experimental threshold (pKD < 5 in all cases). Errors are standard error of the mean (s.e.m.) over a total of 10 replicates

In pose A (Figure 2A), Asn253^{6,55} forms a H-bond with the nitrogen in the 4-methylthiazole group, leaving the carbonyl group potentially exposed to the internal water network stabilized by residues in TM7, as observed in the $A_{2A}AR$ crystal structure with **ZM241385**^[27] (see Supporting Information Figure S1). Conversely, in pose B (Figure 2B) this carbonyl forms an additional hydrogen bond with N253^{6,55}. We hypothesized that in principle, the symmetry axis would allow exchanging substituents at R₆ and R₇ between the binding poses, something that we would explore later in this work (see below).

Our strategy to select the most reliable binding mode was to compute the effect on ligand binding of the 8 mutations from the BPM panel^[39] for each pose, and compare the results with the experimental values. The results (Figure 2C, Table 1 and Supporting Information Table S1) highlight pose A as the binding mode with the best correlation to experimental data. A MAE of 0.50 kcal/mol was observed for this pose, which is comparable to the results obtained for the co-crystallized antagonist **ZM241385**. In addition, the correlation coefficient of $R^2 = 0.74$ falls between those observed for **ZM241385** and **triazine 4g**. Conversely, the corresponding values calculated on pose B are much higher (MAE = 1.53 kcal/mol) and the correlation is completely lost ($R^2 = 0.03$). Additionally, the computed loss of binding affinity upon the N253^{6,55}A mutation is closer to experiment in pose A than in pose B, indicating that the additional H-bond between Asn253^{6,55} and the carbonyl in pose B would not contribute to more favorable binding free energies. From the MD simulations, we further observed that the binding modes are not related by a symmetry axis as initially hypothesized, and the substituents at positions R₆ and R₇ are exploring different positions in the binding site, making them no longer readily interchangeable.

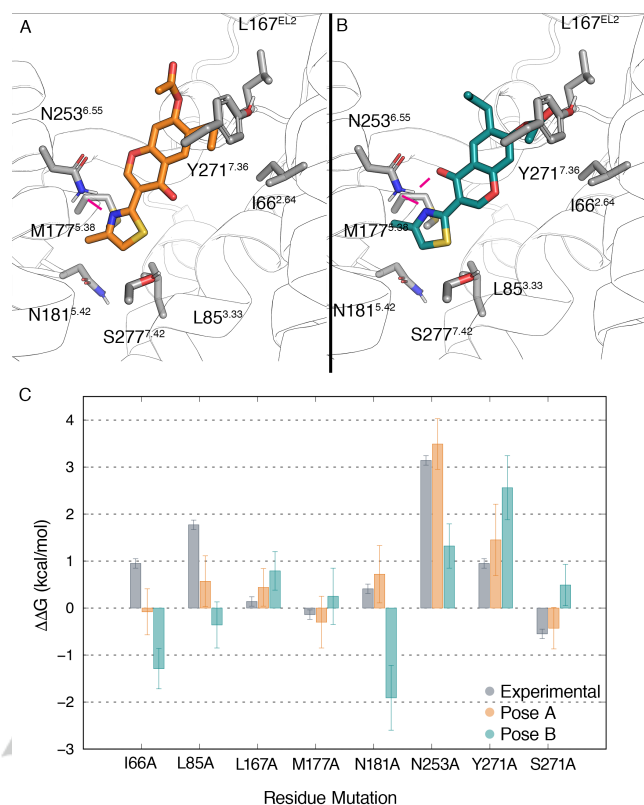


Figure 2. Putative binding modes A (orange) and B (cyan) of **Chromone 14** to the $A_{2A}AR$ (H-bonds in magenta). (C) Experimental and calculated changes in binding free energies for each mutation in the BP. The error bars correspond to the s.e.m. of the replica calculations for the calculated values, or are adjusted to the reported value of 0.1 pKD unit in the case of experimental data.^[39]

Design, synthesis and pharmacological evaluation of chromone derivatives.

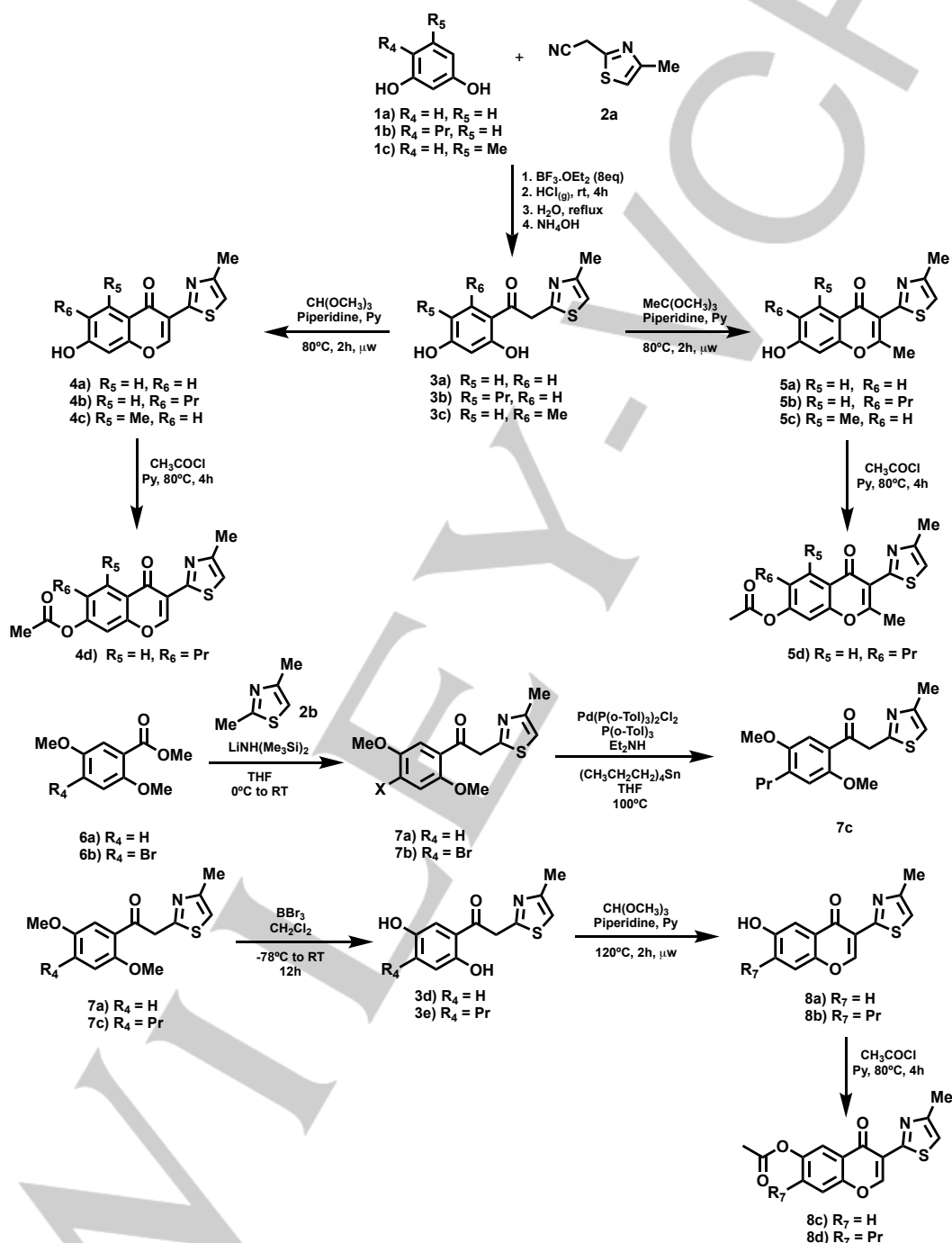
On the basis of the binding mode hypothesis assessed by the FEP calculations, we designed a small collection of chromone derivatives (**Scheme 1**), aiming to explore two different variables on $A_{2A}AR$ affinity: methylation at position 2 (series **5**) and the aforementioned effect of switching the oxygenated function (-OH or -OCOMe) from the original position 7 (series **4**) to position 6 (series **8**), swapping the alkylic substitution (Pro/H) accordingly. A collection of 11 molecules was prepared as depicted in synthetic Scheme 1. Briefly, the key 2-hydroxyphenyl ketones **3** were obtained by either the Hoesch method^[31] or Claisen condensation,^[40] employing phenols **1** or esters **6** and thiazole derivatives **2a** and **2b** as reactive precursors respectively. Treatment of ketones **3** with orthoesters (formate or acetate) enabled the efficient chromone core formation,^[41] thus defining the substituent pattern at position 2 (H or Me). Finally, the required acetates were prepared by reaction of the corresponding phenols with acetyl chloride. The full synthetic methodology is provided in the SI.

The affinity of these compounds for the WT $A_{2A}AR$ was then evaluated with the same SPR assay used to generate the BPM data (see SI methods).^[39] The data confirmed that methylation at position 2 (compounds **5a-d**) is generally unfavorable for binding, compared to the parent series (compounds **4a-d**). This is in agreement with previous work by Andrews et al.,^[31] where

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methylation of the low affinity compound **4a** (corresponding to chromone **8** as reported by Andrews et al.^[31]; $pK_i = 5.7$) resulted in a relative decrease in affinity of 0.6 log units (Chromone **12**, $pK_i = 5.1$), an effect that we observed to be amplified in the case of the high affinity precursor (see **4d** as compared to **5d**). More surprising was the observation of a similar effect when the

substituents between R_6 and R_7 are swapped (compounds **8a-d**), which again was most pronounced for the high affinity compound **4d** (see Table 2), with a drop of affinity of 2.7 log units for the corresponding methylated derivative **8d**.




Scheme 1. Structure and synthetic pathways employed to assemble chromones **4**, **5** and **8**

Computational evaluation of the proposed binding modes.

From the pharmacological data on the novel series of chromones and the initial MD simulations, it appears that the two binding poses differ more than just purely on the rotation axis of the chromone scaffold.

Table 2. SPR affinity data for the series of Chromone derivatives synthesized in this work.



Compound	Substituents				pK _D ^[a]
	R ₂	R ₅	R ₆	R ₇	
4a	H	H	H	OH	5.95
4b	H	H	C ₃ H ₇	OH	6.20
4c	H	CH ₃	H	OH	5.80
4d	H	H	C ₃ H ₇	OCOCH ₃	8.60
5a	CH ₃	H	H	OH	5.10
5b	CH ₃	H	C ₃ H ₇	OH	5.42
5c	CH ₃	CH ₃	H	OH	5.36
5d	CH ₃	H	C ₃ H ₇	OCOCH ₃	5.90
8a	H	H	OH	H	5.50
8b	H	H	OH	C ₃ H ₇	5.70
8d	H	H	OCOCH ₃	C ₃ H ₇	5.90

[a] Experimental errors in all cases reported as approximately 0.1 pK_D unit, i.e. less than 0.1 kcal/mol

The following step was then to investigate whether all compounds in the series would adopt the preferred binding mode A, or if binding mode B is accessible by some of the compounds, depending on the pattern of substitutions. Such a hypothesis was experimentally observed for caffeine, which in contrast to other xanthine derivatives presents a dual binding mode to the A_{2A}AR (see **Figure 3A**). Indeed, an isoenergetic dual binding mode for caffeine was previously hypothesized on the basis of free energy calculations,^[42] before experimental observation of dual-occupancy crystal structures in complex with the A_{2A}AR.^[43,44] Conversely, the X-ray structure of the N₇-demethylated analogue theophylline shows that this molecule adopts only one of these binding modes, where the acidic hydrogen at position N₇ makes an additional H-bond contact with N253^{6.55} (**Figure 3**). We thus evaluated whether our recently developed dual-topology protocol QligFEP was suitable to capture this different behavior, by a direct estimation of relative binding free energies between the two poses, in analogy to the efficiency of this protocol to compare topologically unrelated ligands (e.g. scaffold hopping).^[6] The results for the A_{2A}-xantines system shown in Table 3 show that this is the case: the negligible calculated free energy difference

between the two poses of caffeine is in line with the equally populated dual binding mode in the crystal structure with the A_{2A}AR (PDB code 5MZP, **Figure 3A**). In contrast, the single binding mode observed in the crystal structure of theophylline (blue color in **Figure 3A**) is energetically favored by 1.6 kcal/mol. Subsequent application of the same strategy in the generated chromone series showed a similar energy gap between the two binding poses considered for the simplest chromone in our series (**4a**, **Figure 3B**), with pose A being 1.7 kcal/mol more favorable than pose B. Notably, this energy gap increases significantly for the highest affinity compound **4d**, suggestive of an optimal anchoring of the R₆ acetamide and R₇ n-propyl substituents in pose A (**Table 3**).

Table 3. Calculated free energy difference between two alternative poses for A_{2A}AR antagonists

Ligand A	A → B		H → CH ₃	
	ΔΔG _{calc} (kcal/mol)	Ligand B	ΔΔG _{exp} (kcal/mol)	ΔΔG _{calc} (kcal/mol) ^[a]
Caffeine	0.47 ± 0.49	-	-	-
Theophylline	1.59 ± 0.87	Caffeine	0.6 ^[b]	0.31 ± 0.32
4a	1.65 ± 0.93	5b	1.16 ± 0.1 ^[c]	2.06 ± 0.72
4d	8.76 ± 0.81	5d	3.68 ± 0.1 ^[d]	1.66 ± 1.09

[a] Calculations performed in the selected pose (see text). [b] ΔG_{bind} (caffeine – theophylline), extracted from ChEMBL^[45]. [c] ΔG_{bind} (5a – 4a) and [d] ΔG_{bind} (5d – 4d), data from **Table 2**.

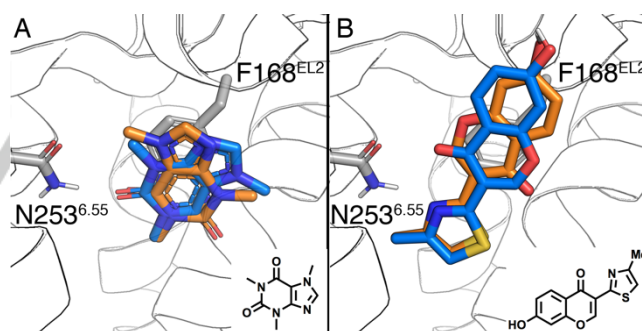


Figure 3. (A) Dual binding mode of caffeine, as extracted from the A_{2A}AR crystal structure with the A_{2A}AR (PDB code 5MZP). Colour code is orange (binding mode A) and blue (binding mode B). (B) modelled binding modes of chromone **4a**, following the same colouring scheme as in panel A.

We consequently retained the selected binding pose for each scaffold (i.e. xanthine and chromone) and computed the affinity shifts observed upon methylation. The calculations reproduced the experimental decrease in affinity observed for caffeine, as compared to the N₇ demethylated analogue theophylline (**Table 3**). Similarly, the corresponding FEP calculations capture the experimentally observed negative effect on A_{2A} affinity due to methylation on R₂ in the original chromones series **4** (see series **5**, see **Table 2**).^[31] The two pair comparisons performed (**4a** → **5a** and **4d** → **5d**, see **Table 3**) qualitatively capture this effect, which is experimentally more pronounced in the last case. Notably, a comparison of the endpoint

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configurations of the **4d** → **5d** FEP transformation provide the structural reason of this effect: to avoid the induced steric hindrance of the new methyl group with Asn253^{6.55}, the core scaffold of **5d** moved approximately 1.5 Å away from the original

pose A of **4d** while maintaining the interaction between the nitrogen in the 4-methylthiazole group and the key asparagine residue.

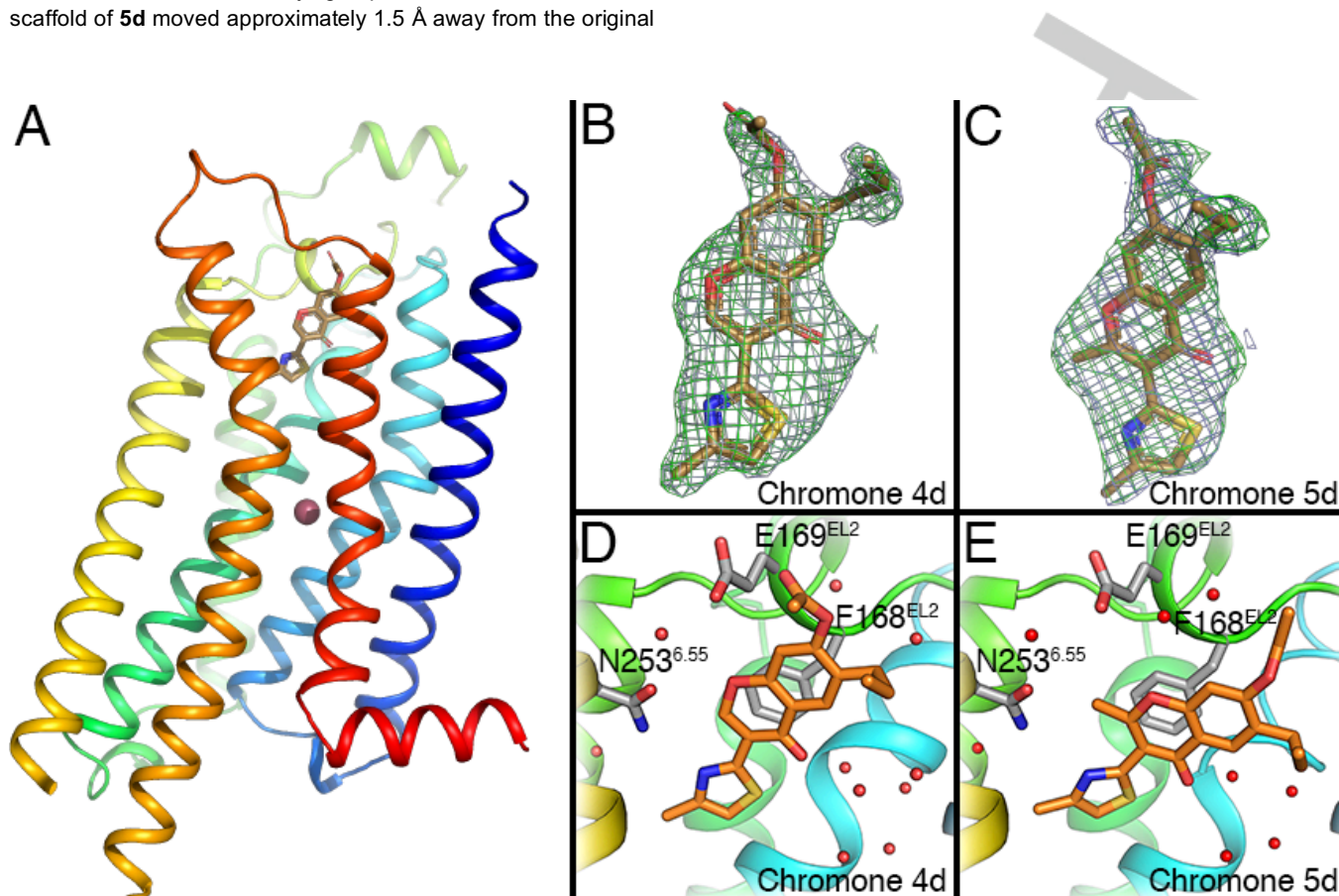


Figure 4. (A) Crystal structures of the $A_{2A}AR$ and compound **4d**, ligand shown in sticks and sodium ion shown as a sphere. Electron densities of chromones **4d** (B) and **5d** (C). Binding mode of compound **4d** (D) and **5d** (E); ligands and the conserved residue N253^{6.55} shown as sticks, water molecules in red spheres.

X-ray Crystallography.

In an effort to univocally reveal the binding mode of the chromone scaffold, the $A_{2A}AR$ -STAR2 construct was crystallized with compound **4d** and its 2-methyl analogue **5d**. The structures were obtained following the *in meso* soaking approach (see methods), and could be refined down to a resolution of 1.92 and 2.13 Å respectively. Statistics for data collection and refinement are given in Supporting Information Table S2. The overall structure of the $A_{2A}AR$ receptor is highly similar to previously solved structures with other antagonists (see Figure 4A), with an RMSD of 0.46 Å for the $C\alpha$ trace as compared to the **ZM241385** structure (PDB 4E1Y, calculated using PyMOL^[46]). The two structures show clear positive omit density at 1σ for the presence of the chromone compounds in the orthosteric binding site (Figure 4B and C), in both cases adopting binding mode A (Figure 4D and 4E). All residues in the binding site show a conserved rotameric state between the two structures and the $A_{2A}AR$ -**ZM241385** complex. The only exception is Y271^{7.36}, which slightly rotates outward served to accommodate the alkyl tail of the chromone at R₆ (see SI Figure S2A), an induced-fit effect that was previously

observed for other $A_{2A}AR$ ligands.^[36] The structure with the high affinity ligand **4d** shows outstanding agreement with the computational model (see Figure 5A) with an RMSD of 0.67 Å between the docked and experimental poses, and is indeed highly similar to previously reported binding modes of the compound.^[31,35] The core of the scaffold is anchored by a H-bond between the nitrogen in the 4-methylthiazole ring and N253^{6.55}, and complemented by a π - π stacking with F168^{EL2}, in line with the binding pattern observed for many other $A_{2A}AR$ antagonists.²¹ The ester at position R₇ is lined towards the Glu169^{5.30} – His264^{7.29} ionic pair, which closes the EL2-EL3 interface, while the propyl at R₆ makes contacts with Tyr271^{7.53} and Met270^{7.52}. Given the good resolution of the structure, we could observe a well-defined water network in the interface between the carbonyl moiety of the chromone core and the receptor (Figure 4D). While most water positions are conserved in the $A_{2A}AR$ -**ZM241385** complex (see Supporting Information Figure 2B), ligand **4d** displaces a number of water molecules observed in the first hydration shell around that ligand. Some of these waters were previously associated to a high-energy or “unhappy” state,^[31] which could partially explain the high affinity of this particular compound.

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The A_{2A}AR-5b complex (Figure 4E) further shows that methylation at position R₂ displaces the chromone core by 1.79 Å, in accordance with observations from the FEP simulations (Table 3 and Figure 5B, RMSD = 1.24 Å between docked and experimental ligand configuration). The H-bond interaction between the nitrogen in the 4-methylthiazole ring of compound 5d and Asn253^{6,55} is maintained, but the methylation does not disrupt the position of Asn253^{6,55}, which remains linked to Glu169^{5,30} via a conserved water molecule. The relative displacement of the chromone core situates the substituents at R₆ and R₇ in sub-optimal pockets (Figure 4D), displacing two of the water molecules described for compound 4d (see Figure 5D). Conversely, the water molecule deeper in the binding crevice remains, mediating a common anchoring point between the carbonyl group of the ligands and His276^{7,43}.

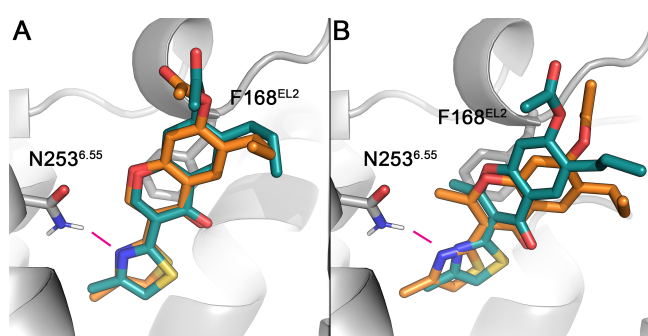


Figure 5. Crystal structure (orange) and modelled coordinates (cyan) of (A) the highest affinity compound 4d and (B) the methylated derivative 5d (panel B) with the A_{2A}AR. H-bond interactions are indicated in magenta.

Conclusion

We described a robust workflow to iteratively improve receptor-ligand binding models, based on mapping of available experimental data onto structural information via free energy calculations. We applied this protocol to provide novel insights in the binding mode of a recent series of A_{2A}AR antagonists. The optimization included the identification of an initial binding mode hypothesis, based on the exploration of BPM and SAR data of the original chromone series using FEP. Thereafter, new derivatives were designed, synthesized and evaluated, which was followed by a second iteration of FEP calculations that reinforced the binding hypothesis. Finally, X-ray crystallography experimentally confirmed this binding mode, supporting the rational design of these compounds. These structures, combined with the FEP calculations, provide structural and energetic insights in the determinants of high affinity binding of the chromone scaffold series. We expect the presented workflow to be of general applicability in structure-based drug design, particularly in the case of GPCRs where structures of receptor-ligand complexes are increasingly available.

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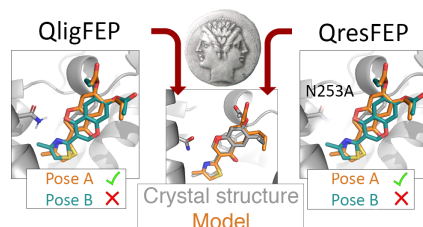
Keywords: Adenosine receptors • Biophysical mapping (BPM) • Free energy perturbation (FEP) • G protein-coupled receptor (GPCR).

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Entry for the Table of Contents

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In molecular design, structural, pharmacological and chemical information can be interconnected by computational estimations of binding free energies. Using a combined FEP approach to examine both mutagenesis and ligand SAR, we designed new analogues of the A_{2A} AR antagonist series of chromones. Subsequent crystal structures supported the rational design of these compounds, linking the structural and energetic understanding on ligand binding.

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