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Structural Basis for the High-Affinity Inhibition of Mammalian Membranous Adenylyl Cyclase by 2',3'
O-(N-Methylanthraniloyl)-Inosine 5'-Triphosphate[†]

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MOL #71894

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Crystal structure of adenylyl cyclase with MANT-ITP

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Number of text pages: 33

Number of Tables: 2

Number of Figures: 5

Number of References: 40

Number of words in Abstract: 244

Number of words in Introduction: 648

Number of words in Discussion: 1194

Abbreviations

mAC, mammalian membranous adenylyl cyclase; FRET, fluorescence resonance energy transfer; FS, forskolin; GTP γ S, guanosine 5'-[γ -thio]triphosphate; MANT-GTP, 2',3'-O-(N-methylanthraniloyl)-guanosine 5'-triphosphate; ITP γ S, inosine 5'-[γ -thio]triphosphate; MANT-ITP, 2',3'-O-(N-methylanthraniloyl)-inosine 5'-triphosphate; MANT-XTP, 2',3'-O-(N-methylanthraniloyl)-xanthosine 5'-triphosphate; VCI and IIC2, the N- and C-terminal catalytic domains from canine AC5 and rat AC2 expressed as soluble proteins

Abstract

2',3'-O-(N-Methylanthraniloyl)-inosine 5'-triphosphate (MANT-ITP) is the most potent inhibitor of mammalian membranous adenylyl cyclase (mAC) 5 (AC5, Ki, 1 nM) yet discovered and surpasses the potency of 2',3'-O-(N-methylanthraniloyl)-quanosine 5'-triphosphate (MANT-GTP) 55-fold (Göttle et al., J Pharmacol Exp Ther 329:1156-1165 (2009)). AC5 inhibitors may be valuable drugs for treatment of heart failure. The aim of this study was to elucidate the structural basis for the high-affinity inhibition of mAC by MANT-ITP. MANT-ITP was a considerably more potent inhibitor of the purified catalytic domains VC1 and IIC2 of mAC than MANT-GTP (Ki, 0.7 nM versus 18 nM). Moreover, there was considerably more efficient fluorescence resonance energy transfer between W1020 of IIC2 and the MANT-group of MANT-ITP compared to MANT-GTP, indicating optimal interaction of the MANT-group of MANT-ITP with the hydrophobic pocket. The crystal structure of MANT-ITP in complex with the $G_{s\alpha}$ and forskolin-activated catalytic domains VC1:IIC2 in comparison to the existing MANT-GTP crystal structure revealed only subtle differences in binding mode. The higher affinity of MANT-ITP to mAC compared to MANT-GTP is probably due to fewer stereochemical constraints upon the nucleotide base in the purine binding pocket, allowing a stronger interaction with the hydrophobic regions of IIC2 domain, as assessed by fluorescence spectroscopy. Stronger interaction is also achieved in the phosphate-binding site. The triphosphate group of MANT-ITP exhibits better metal coordination than the triphosphate group of MANT-GTP, as confirmed by molecular dynamics simulations. Collectively, the subtle differences in ligand structure have profound effects on affinity for mAC.

Introduction

Mammals express nine membranous AC isoforms (ACs 1-9) that play an important role in transmembrane signal transduction (Sunahara et al., 1996; Tang and Hurley, 1998). ACs are activated by the G-protein G_s *via* numerous receptors for hormones and neurotransmitters and catalyze the production of the second messenger cAMP. Studies with AC5^{-/-} mice indicate that potent and selective AC5 inhibitors may be valuable drugs for the treatment of heart failure, ageing, bone loss, anxiety as well as acute and chronic pain (Rottländer et al., 2007; Kim et al., 2008; Chester and Watts, 2007; Okumura et al., 2009).

Previous studies from our group showed that 2',3'-O-(N-methylanthraniloyl) (MANT)-substituted purine nucleotides are competitive AC inhibitors and inhibit AC isoforms differentially (Gille and Seifert, 2003; Gille et al., 2004). Moreover, MANTnucleotides are fluorescence probes of conformational changes in the catalytic site of AC. Crystal structures of the purified catalytic subunits of G_{sa} -bound mammalian AC (C1 subunit of AC5 (VC1) and C2 subunit of AC2 (IIC2) in complex with MANT-GTP and MANT-ATP) were resolved (Mou et al., 2005, 2006). Those studies revealed a tripartite binding pocket for MANT-nucleotides in the catalytic core consisting of sites for the base, the MANT-substituted ribosyl group and the polyphosphate chain. The combined analysis of enzymatic, fluorescence spectroscopy, crystallographic and molecular modelling data revealed that the MANT-group contributes most to the AC binding energy, whereas the base makes the smallest contribution. This is reflected by the fact that MANT-nucleotides exhibit up to 17,000-fold higher affinity for AC than their non-substituted parent nucleotides and that purine and pyrimidine nucleotides exhibit similar affinity for AC (Gille et al., 2004, 2005). Based on those data, we proposed that ACs exhibit a high degree of conformational flexibility, allowing the

catalytic site to accommodate structurally diverse bases (Mou et al., 2006; Wang et al., 2007).

During the course of our subsequent systematic studies on 2',3'-O-ribosylsubstituted nucleotides as mAC inhibitors (Göttle et al., 2009; Suryanarayana et al., 2009), we serendipitously identified MANT-ITP as the most potent mAC inhibitor known so far. Specifically, MANT-ITP inhibits AC5 with a K_i value of 1 nM and mouse heart AC (predominantly representing AC5) with a K_i value of 4 nM. Compared to MANT-GTP, MANT-ITP possesses a 55-fold higher affinity for recombinant AC5. Chemically, the base hypoxanthine differs from the base quanine by the absence of an NH₂ group at C2 of the purine ring (Fig. 1). Accordingly, hypoxanthine cannot to form a hydrogen bond with the backbone oxygen of I1019 (Mou et al., 2005). Based on analogous studies with GTP, ITP and XTP on GTP-binding proteins (Seifert et al., 1999; Gille et al., 2003; Gille and Seifert, 2004), one would also expect that the missing hydrogen bond should actually reduce affinity of mAC for MANT-ITP. This notion is further supported by the fact that MANT-XTP, bearing a keto group at C2 of the purine ring (Fig. 1), binds to mAC with much lower affinity than MANT-GTP because of electrostatic repulsion of the keto group of the xanthine group by the backbone oxygen of I1019 (Mou et al., 2005).

Therefore, the aim of our present study was to elucidate the structural basis for the high-affinity interaction of mAC with MANT-ITP, using the VC1:IIC2 heterodimer as an established model of the mAC catalytic domain. Firstly, we determined the affinities of VC1:IIC2 for MANT-GTP, MANT-ITP and MANT-XTP in enzyme activity assays (Gille et al., 2004). Secondly, we performed fluorescence studies with the three MANT-nucleotides (Mou et al., 2006). Thirdly, we analyzed the crystal structure of VC1:IIC2 in complex with MANT-ITP and compared this structure with the known VC1:IIC2 structure in complex with MANT-GTP (Mou et al., 2005).

Lastly, we conducted molecular dynamics simulations to further evaluate the structural requirements for the high-affinity interaction of MANT-ITP with AC5 in comparison to MANT-GTP (Wang et al., 2007). Here, we show that even subtle differences in ligand structure have profound effects for interaction with mAC.

Materials and Methods

Materials. MANT-ITP was synthesized according to Hiratsuka (1983) with the modifications described by Taha et al. (2009). MANT-GTP and MANT-XTP were obtained from Jena Bioscience, Jena, Germany. [α - 32 P]ATP (800 Ci/mmol) was purchased from PerkinElmer Life and Analytical Sciences (Boston, MA). Aluminum oxide 90 active, neutral (activity 1, particle size 0.06-0.2 mm) was purchased from Merck, Darmstadt, Germany. Bovine serum albumin, fraction V, highest quality, was from Sigma-Aldrich. MnCl₂ tetrahydrate (highest quality) was from Merck. FS was from LC Laboratories (Woburn, MA).

AC activity assay. AC activity was determined essentially as described in the literature (Mou et al., 2005). Briefly, reaction mixtures contained 100 μ M ATP/Mn²⁺, 10 mM MnCl₂ and MANT-nucleotides at concentrations from 0.1 nM to 1 mM as appropriate to obtain saturated inhibition curves. Additionally, assay tubes contained VC1 (8 nM) and IIC2 (40 nM). For experiments with $G_{s\alpha\text{-GTP}\gamma S}$, tubes contained VCI (3 nM), IIC2 (15 nM) and $G_{s\alpha\text{-GTP}\gamma S}$ (51 nM). Reactions were conducted in the presence of 100 μ M FS. Following a 2 min pre-incubation at 30°C, reactions were initiated by adding 20 μ l of reaction mixture containing (final) 1.0 μ Ci/tube [α -³²P]ATP, 0.1 mM cAMP and 100 mM KCl in 25 mM HEPES/NaOH, pH 7.4. Reactions were conducted for 10-20 min at 30°C and were terminated by the addition of 20 μ L of 2.2 N HCl. Denatured protein was precipitated by a 1 min centrifugation at 25°C and 15,000 x g.

Sixty-five μ I of the supernatant fluid were applied onto disposable columns filled with 1.3 g neutral alumina. [32 P]cAMP was separated from [α - 32 P]ATP by elution of [32 P]cAMP with 4 ml of 0.1 M ammonium acetate, pH 7.0. Recovery of [32 P]cAMP was ~80% as assessed with [3 H]cAMP as standard. [32 P]cAMP was determined by liquid scintillation counting using Ecolume scintillation cocktail (Fisher, Pittsburgh, PA). Competition isotherms were analyzed by non-linear regression using the Prism 4.0 software (GraphPad, San Diego, CA).

Fluorescence spectroscopy. All experiments were conducted using a Cary Eclipse fluorescence spectrophotometer equipped with a Peltier-thermostated multicell holder at 25°C (Varian, Palo Alto, CA). Measurements were performed in a quartz fluorescence microcuvette (Hellma, Plainview, NY). The final assay volume was 150 μl. Reaction mixtures contained a buffer consisting of 100 mM KCl, 10 mM MnCl₂ and 25 mM HEPES/NaOH, pH 7.4. Steady-state emission spectra were recorded at low speed with λ_{ex} = 350 nm (λ_{em} = 370-500 nm) and λ_{ex} = 280 nm (λ_{em} = 300-500 nm) with various MANT-nucleotides (1 μM each) in the absence and presence of 5 μM VC1 plus 25 μM IIC2 without and with 100 μM FS. Fluorescence recordings were analyzed with the spectrum package of the Cary Eclipse software (Varian, Walnut Creek, CA). Baseline fluorescence (buffer alone) was subtracted. Fig. 2 shows superimposed original fluorescence recordings representative for two to three independent experiments with at least two different batches of VC1:IIC2.

Preparation of Proteins. The catalytic subunits VC1, IIC2 and GTPγS activated $G_{s\alpha}$ were expressed in *Escherichia coli* BL21(DE3) cells, containing pREP4 plasmid. The plasmids encode the wild type C1a domain of canine AC 5 (residue 364-580), the C2a domain of rat AC 2 (residue 874-1018) and bovine $G_{s\alpha}$ (residue 1-396). VC1 and $G_{s\alpha}$ constructs were expressed with a hexahistidine tag at their N and

C-terminus, respectively. They were purified and stored as described previously (Tesmer et al., 2002). $G_{s\alpha}$ was further activated by incubation with 500 μ M GTP γ S and 2 mM MgCl $_2$ at 30°C for 2 hours, and digestion of the complex with trypsin was necessary for crystallization work. The smaller fragment, containing residues 39-387 was further purified using Ni-NTA column followed by MonoQ anion exchange chromatography.

Complex Formation and Crystallization with MANT-ITP. To form a stable heterotrimeric complex the recombinant proteins were mixed in the following order VC1-IIC2- $G_{s\alpha}$ -GTP γ S in a molar ratio 1.5:1:1. Thereafter, forskolin (200 μ M) was added to further stabilize the complex. The protein mixture was incubated on ice for at least 30 min and then applied to tandemly arranged Superdex 75 and 200 gel filtration columns (Amersham Biosciences). Only the fractions containing the complex were collected and concentrated to ~9.5 mg/ml in a buffer of 20 mM HEPES, pH 8.0, 1 mM EDTA, 2 mM MgCl₂, 5 mM dithiothreitol, 100 mM NaCl, 200 μM 7-acetyl-7-[O-(N-methylpiperazino)-γ-butyryl)]-forskolin and 500 μM GTPγS. The concentrated protein solution was used to grow crystals via the sitting drop method with a reservoir solution of 7.2 % (m/v) polyethylene glycol 8000, 0.5 M NaCl and 0.1 M MES, pH 5.4, at 16 °C for 3-4 weeks. Large crystals were soaked with 2 mM MANT-ITP and 3 mM MnCl₂ for at least 2 h at room temperature and then harvested in a cryoprotectant consisting of 9% (m/v) polyethylene glycol 8000, 30% (m/v) polyethylene glycol 400, 0.1 M MES pH 5.4, 0.5 NaCl, 20 mM HEPES, pH 8.0, 1 mM EDTA, 2 mM dithiothreitol, 200 μM 7-acetyl-7-[O-(N-methyl-piperazino)-γ-butyryl)]-forskolin, 100 μM GTPγS, 2 mM MANT-ITP and 3 mM MnCl₂. The cryoprotected crystals were mounted in 0.1-0.2 mm loops and stored in liquid nitrogen.

Structure Determination and Model Refinement. Diffraction data sets were collected at the Stanford Synchotron Radiation Lightsource SSRL-SMB-MC 9-1 beamline (Stanford, CA) by the oscillation method (1°/frame, 60s/frame). The incident beam wavelength was 0.9795 Å. The images were processed using HKL2000 package (Otwinowski and Minor, 1997). Due to anisotropy of the plate-type crystal, data with / index > 21 were excluded from the data set. Structures were determined by molecular replacement using the structure of the $G_{s\alpha}$ -GTP γ S:VC1:IIC2 complex (PDB code: 1AZS) as the initial phasing model (Tesmer et al., 1997). Atomic positions and thermal parameters of the mAC structure were refined by Refmac5.5 using the CCP4 program suite (Collaborative Computational Project, 1994). MANT-ITP and metal ions in the structure were located in the weighted |F_o|-|F_c| omit map computed with phases from the refined model. The model was iteratively improved by manual refitting into weighted 2|F_o|-|F_c|map using the computer graphics program Coot (Emsley and Cowtan, 2004) and subsequent refinement cycles with CCP4. The refined crystal structure was visualized with PyMOL (DeLano, 2002). Coordinates for the MANT-ITP:Mn²⁺ structure were deposited in the Protein Data Bank with the code PDB 3G82.

Molecular Dynamics Simulations. Atomic coordinates of protein-ligand complexes for MANT-ITP and MANT-GTP interacting with VC1:IIC2 were extracted from structures PDB 3G82 and PDB 1TL7. Ligand parametrization for FS, MANT-GTP and MANT-ITP was performed by extracting the relevant ligands from the crystal structures and editing them via SYBYL to ensure proper representation of valence and bond types. Thereafter the antechamber (Wang et al., 2001) module of AMBER10 was used to assign AM1-BCC charges to the ligands and calculate force field parameters for them. In order to avoid excessively large computational expense, only the VC1:IIC2 portion of each crystal structure (plus ligand and co-factor) was

retained while removing the bound $G_{s\alpha\text{-}GTP\gamma S}$. The tleap module of AMBER10 was used for the preparation of topology and coordinate files for the protein-ligand complexes using the ff99SB force field parameters for protein and the antechamber-calculated parameters for ligands. The parameters for Mn²⁺ were obtained from Bradbrook et al. (1998). The structures 3G82 and 1TL7 were separately solvated in water boxes with buffering distance of 10 Å. Assuming normal charge states of ionizable groups corresponding to pH 7.0, Na⁺ ions were added to achieve charge neutrality and to mimic the biological environment more closely.

Our primary simulation engine for probing the differences in the dynamic nature of the two complexes was NAMD (Phillips et al., 2005), which was chosen because of its excellent parallel scalability which enabled us to perform the simulations in an expeditious manner on 32 nodes of a large Linux cluster. In order to relieve severe steric contacts and instances of higher energy conformations that might destabilize the molecular dynamics integrator at later stages, the system was subjected to initial minimization of 20,000 steps wherein the protein backbone was held fixed (in order to relax surrounding water molecules and computationally specified hydrogen positions), followed by 20,000 steps of minimization without positional constraints (i.e., to allow the system to relax freely). The resulting lowenergy (~0K) model was then equilibrated to room temperature (~300K) by gradually increasing the system temperature in increments of 20K up to a target temperature of 300K. A each of the 15 temperature increments 15,000 dynamics steps (30 ps) were run while employing a restraint of 10 kcal mol⁻¹ Å⁻² on protein α carbons (C_{α}) to avoid any prospect of unrealistic denaturation behavior. Thereafter, the system was equilibrated for 150,000 steps (300 ps) at 310 K (approximate physiological temperature) under conditions of constant volume and temperature (NVT), and then for a further 150,000 steps (300 ps) at 310K using a Langevin piston (constant

pressure and temperature = NPT) to achieve uniform pressure of 1 bar. Finally the restraints were removed and the system was equilibrated for 500,000 steps (1 ns) to prepare the system for final analysis. For the latter, a NPT simulation was run on the equilibrated structure for 9.6 ns keeping the temperature at 310 K and pressure at 1 bar using Langevin piston coupling algorithm. The integration time step of the simulations was set to 2.0 fs, the SHAKE algorithm was used to constrain the lengths of all chemical bonds involving hydrogen atoms at their equilibrium values and the geometry was restrained rigidly via the SETTLE algorithm. Non-bonded van der Waals interactions were treated via a switching function at 10 Å and reaching zero influence at a distance of 12 Å, and the Mn²⁺ cofactors were subjected to a soft harmonic constraint (1 kcal/mol) to encourage adherence to equilibrium approach distances relative to the ligand phosphate oxygens as determined from an earlier simulation. The particle-mesh Ewald algorithm (PME) as implied in NAMD was used to handle long range electrostatic forces.

In order to probe relative contributions to the total free energy of the complexes, we employed the GBSA (Generalized Born, augmented by solvent accessible surface) method (Weiser et al., 1999; Onufriev et al., 2000; Bashford and Case, 2000). For this, the trajectory obtained from the molecular dynamics run was converted into individual coordinate files at 100 ps intervals excluding the initial 1.0 ns of the simulation. Thus, 86 frames were used in GBSA calculation for each complex. AMBER parameters (using bonding radii for GBSA calculation) were then generated from the resulting structures using tleap and were subsequently applied in GBSA calculation. The radii for Mn²⁺ was specified as being the same as Mg²⁺, since GBSA parameters for Mn are unavailable in AMBER. The change of conformational entropy was not considered. Apart from a number of settings chosen specifically for this analysis default GBSA parameters were employed (IGB = 2 to specify the OBC

model, GBSA = 1 to choose the LCPO method for solvent accessible surface area calculation, EXTDIEL = 78.50 to set the solvent dielectric for room temperature water, INTDIEL = 1.0 to specify a solute dielectric for the system, and SURFTEN = 0.0072 to describe the water surface tension).

Results

Enzymatic studies. Table 1 lists the K_i values of MANT-GTP, MANT-ITP and MANT-XTP for inhibition of VC1:IIC2 under maximally stimulatory conditions, i.e. in the presence of $G_{S\alpha\text{-GTP}\gamma S}$, as well as under submaximally stimulatory conditions, i.e. in the absence of $G_{S\alpha\text{-GTP}\gamma S}$. The latter reflect the assay conditions for the fluorescence spectroscopy studies. In accordance with the data for AC5 and mouse heart AC (Göttle et al., 2009), under maximally stimulatory conditions, MANT-ITP was a considerably more potent inhibitor of VC1:IIC2 than MANT-GTP which, in turn, was much more potent than MANT-XTP. The omission of $G_{S\alpha\text{-GTP}\gamma S}$ reduced the overall MANT-nucleotide potencies by 4-10-fold, but the rank order of affinity of nucleotides was preserved. Collectively, the enzyme inhibition studies with purified catalytic subunits of mAC confirmed the exceptionally high affinity of MANT-ITP for the catalytic site previously reported for AC5 and mouse heart AC (Göttle et al., 2009).

Fluorescence spectroscopy studies. In order to elucidate further differences in the interaction of MANT-nucleotides with VC1:IIC2, we exploited the fluorescence properties of these nucleotides (Jameson and Eccleston, 1997) and the ability of the MANT-group to bind to a hydrophobic pocket in the interface of VC1:IIC2 (Mou et al., 2005, 2006). The emission spectra of nucleotides at λ_{ex} = 350 nm for direct excitation

of the MANT-group (Jameson and Eccleston, 1997), and at λ_{ex} = 280 nm for analysis of FRET between W1020 in IIC2 and the MANT-group were determined (Mou et al., 2005). Fluorescence studies were performed in the presence of a five-fold excess of VC1 relative to MANT-nucleotides in order to ensure quantitative ligand binding to the catalytic site.

At λ_{ex} = 280 nm, MANT-nucleotides were only minimally excited, whereas at λ_{ex} = 350 nm they showed substantial intrinsic fluorescence signals with an emission peak at ~ 450 nm (Fig. 2, blue tracings). The dashed black lines indicate the endogenous tryptophan fluorescence of VC1:IIC2 at λ_{ex} = 280 nm, i.e. the fluorescence in the absence of MANT-nucleotide. Following the addition of VC1:IIC2, at λ_{ex} = 280 nm, MANT-ITP exhibited a much higher basal FRET signal, as revealed by a second emission peak at λ_{em} = 420 nm, than MANT-GTP (green tracings, panels 2A and C). At λ_{ex} = 350 nm, the interaction of MANT-ITP with VC1:IIC2 resulted in considerably higher increases in fluorescence compared with MANT-GTP (green tracings, Figs. 2B and D). The blue-shift of the fluorescence emission (Mou et al., 2005) was similar for MANT-ITP and MANT-GTP. FS (100 μM) increased basal FRET and direct fluorescence with MANT-GTP more effectively than with MANT-ITP, but the absolute FRET with MANT-ITP was still considerably larger than with MANT-GTP. These data suggest that the MANT-group of MANT-ITP binds to the hydrophobic pocket in mAC more effectively than MANT-GTP. Compared to MANT-GTP and MANT-ITP, MANT-XTP exhibited only minimal FRET and direct fluorescence (Figs. 2E and F), reflecting suboptimal binding of the xanthine ring to mAC and suboptimal insertion of the MANT-group into the hydrophobic pocket (Mou et al., 2005).

Crystallographic studies. To better understand the high inhibitory potency of MANT-ITP at ACs 1, 2 and 5 (Göttle et al., 2009) as well as at the catalytic domains VC1:IIC2 (Table 1), crystallographic studies were conducted. Crystals of FS-VC1:IIC2-G_{sα-GTPγS} were soaked with 2 mM MANT-ITP and 3 mM MnCl₂. The structure of the MANT-ITP:Mn²⁺ complex was determined at a resolution of 3.1 Å by molecular replacement using the structure of the G_{sα-GTPγS}:VC1:IIC2 complex as the initial phasing model (Tesmer et al., 1997) (PDB code: 1AZS). Crystallographic data collection and refinement statistics are summarized in Table 2. Interactions between G_{sα-GTPγS} and the pseudosymmetric VC1 and IIC2 catalytic domains center largely on IIC2 as described (Tesmer et al., 1997). The two domains form a very large interface, facilitating the binding of MANT-ITP at the catalytic site, and FS at the pseudo-dyadrelated site. The structure was very similar to the corresponding complex with MANT-GTP (Fig. 1) (Mou et al., 2005) (PDB code 1TL7). Superimposing the two structures, the overall placement of VC1 and IIC2 did not differ greatly from each other with the RMSD less than 0.5 Å (Fig. 3C).

MANT-ITP was modeled into the continuous |F_o|-|F_c| map in the binding pocket. The electron density was more consistent with 3'-O-MANT-ITP than the 2'-O-MANT isomer (Fig. 3A). Our previous crystallographic studies with MANT-GTP and MANT-ATP gave similar results (Mou et al., 2005, 200). Difference electron density peaks corresponding to the two Mn²⁺ ions are observed; wherein the A-site exhibits lower occupancy than the B-site suggesting the Mn²⁺ ion is bound more tightly at the B-site as observed in other mAC crystal structures (Mou et al., 2005, 2006).

The overall conformation of MANT-ITP bound to VC1:IIC2 is similar to that observed for MANT-GTP. The ligands are similar with respect to interactions with protein residues and metal ion coordination (Figs. 3B and D). The $|F_o|$ - $|F_c|$ electron density for MANT-ITP is well defined and similar to that observed for MANT-GTP,

indicating no obvious difference in the conformation of the two ligands. One noticeable difference is that the hypoxanthine ring of MANT-ITP lacks an amino group at the C2 position (Fig. 1) that could form a hydrogen bond with the side chain of I1019 of IIC2 domain (MANT-GTP 2.6 Å; Fig. 3D). Despite the absence of this interaction, MANT-ITP binds with higher affinity to VC1:IIC2 than MANT-GTP, as indicated above (Table 1). The absence of the C2-amino group allows a higher degree of mobility of the purine moiety than in the case of MANT-GTP (Fig. 3D). The purine ring of MANT-ITP is somewhat more deeply inserted in the mAC binding site than that of MANT-GTP.

In the crystal structure the MANT-group of MANT-ITP and MANT-GTP form similar interactions at the $\alpha 4$ '- $\alpha 1$ domain interface. The aryl function of MANT engages in hydrophobic interactions with A409, L412, V413, V1024, V1026 and W1020 (Fig. 4). The increase in the fluorescence signal for MANT-ITP may e due to changes of the relative positions of W1020 and the MANT-group (Fig. 2). However, electron density is weak for the MANT moiety, suggesting that it is poorly ordered within the binding site (Fig. 3A). The oxygen of the carbonyl group of MANT-ITP is in closer proximity to N1025 compared to that of MANT-GTP (Fig. 4). However, in this orientation, the side chain of N1025 does not form an H-bond with the carbonyl group of MANT-ITP.

A stronger interaction of MANT-ITP with the phosphate binding site is supported by two observations. The side chain K1065 of IIC2 interacts with the β - and γ -phosphates of MANT-ITP, whereas the amino group of K1065 in the MANT-GTP structure is only oriented towards the γ -phosphate (Fig. 3C). The amino group of K1065 is also closer to the oxygen of the β -phosphate of MANT-ITP (2.9 Å) in comparison to the oxygen of the γ -phosphate of MANT-GTP (3.1 Å).

Of particular interest are differences between the MANT-ITP and MANT-GTP complexes with respect to coordination of the Mn^{2+} ions by the nucleotide β and γ phosphates and the side chain of D396. The carboxylate group of D396 coordinates the two metal ions, together with the nucleotide phosphates and I397 and D440 of VC1 (Fig. 3B). Diffuse electron density in the region of the β -phosphate is consistent with conformational heterogeneity of the ligand, and consequently, its ligation of the Mn^{2+} ions at the A and B sites (Fig. 3A). This may account in part for differences in metal coordination for MANT-ITP and MANT-GTP. The Mn^{2+} ions in the MANT-ITP structure form close contacts with the α - and γ -phosphates whereas in the MANT-GTP structure the Mn^{2+} ions interact predominantly with the β -phosphate. It also appears that the β -phosphate group of MANT-ITP is more tightly coordinated, due to the shorter (2.8 Å) contact with the carbonyl group of I397 than is the case for MANT-GTP (3.4 Å). The phosphate site plays a crucial role for binding affinity since phosphate group removal dramatically reduces inhibitor potency (Gille et al., 2004).

Molecular dynamics simulations. Our molecular dynamics simulations corroborate the finding that MANT-ITP has higher affinity for VC1:IIC2 than with MANT-GTP. Specifically, the GBSA free energy analysis determined a total free energy for the MANT-ITP/VC1:IIC2 system of -99.71 \pm 10.29 kcal/mol (mean \pm SD, as derived from sampling 86 time steps), whereas the MANT-GTP/VC1:IIC2 system complex had a free energy of only -69.60 \pm 8.94 kcal/mol. Nearly all of this difference can be accounted for in terms of two factors: NT-ITP complex was predicted to derive a substantial advantage from electrostatics (-951.66 \pm 36.75 kcal/mol vs. -847.41 \pm 38.66 kcal/mol for MANT-GTP) but to incur a penalty in terms of a less favorable solvation profile (890.24 \pm 30.79 kcal/mol vs. 819.68 \pm 34.53 kcal/mol). To ascertain

the source of these interaction differences, we performed distance analysis over a set of ~9,600 sample conformers (taken from each 1,000 time steps in the 9.6 ns simulation) to identify any specific receptor-ligand interatomic distances that differed significantly from one ligand to the other. The most important distinction was found not in comparing the hypoxanthine and guanine rings (the only true chemical difference between the two ligands), but rather in the interactions between the Mn²⁺ ions of the receptor and an oxygen on the β-phosphate group of the ligand triphosphate moiety. Specifically, while both ligands orient one α -, one β - and one γ phosphate oxygen in close association to one or more of the metal ions, the dynamic conformation of MANT-ITP enables a second β oxygen to remain significantly closer to a Mn²⁺ ion (2.36 \pm 0.08 Å) than is the case for MANT-GTP (3.29 \pm 0.65 Å). The substantially shorter mean distance between the MANT-ITP β oxygen and the nearest Mn²⁺ suggests a significantly stronger electrostatic interaction (Fig. 5). Specifically, a twist in the orientation of the nucleobase propagates through the ribosyl moiety (in ways that do not substantially affect the interactions of these groups) to the triphosphate chain. The triphosphate group is pulled about 0.7 Å closer to the nucleobase, which has the important effect of positioning both β oxygens of MANT-ITP (rather than just one for MANT-GTP) in an orientation that permits interaction with Mn²⁺ ions. Interestingly, the much greater standard deviation observed for the the β oxygen to Mn^{2+} interaction in MANT-GTP complex suggests that the manganese ion can interact more readily with solvent in the latter case, which is a plausible cause for the more favorable solvation energy for the MANT-GTP complex. Supplemental Figure 1 and Supplemental Tables 1-3 provide further details on the differences in interactions of MANT-GTP and MANT-IP with VC1:IIC2.

Discussion

We developed a tripartite pharmacophore model for mAC with a binding site for the base, the (substituted) ribosyl group and the polyphosphate chain (Mou et al., 2005, 2006). In a previous study, we reported that MANT-GTP γ S and MANT-ITP γ S are similarly potent inhibitors of various mACs (Gille et al., 2004). Omission of the MANT-group reduces inhibitor-affinity by several orders of magnitude, highlighting the importance of the MANT-binding site for high inhibitor-affinity (Gille et al., 2004). Interestingly, unsubstituted ITP γ S is a more potent mAC inhibitor than GTP γ S, whereas GTP and ITP exhibit similar affinity (Gille et al., 2005). In contrast, MANT-ITP is a considerably more potent mAC inhibitor than MANT-GTP (Göttle et al., 2009). These data indicate that subtle structural changes in nucleotide inhibitors (exchange guanine – hypoxanthine; exchange γ -phosphate – γ -thiophosphate) substantially change the relative contributions of the three binding sub-sites in mAC for inhibitor affinity. The main goal of the present study was to elucidate the structural basis for the exceptionally high affinity of mAC for MANT-ITP (Göttle et al., 2009).

Our data suggest that a balance of binding energies among the three pharmacophores in the mAC binding site (Mou et al., 2006) considerably affects the affinity of mAC for MANT-ITP versus MANT-GTP. The MANT-ITP structure shares the common features with previously published mAC structures in complex with 2',3'—substituted purine and pyrimidine nucleotide inhibitors, where the base, triphosphate and 2',3'-ribose stubstituent reside in three distinct grooves of the substrate binding site (Mou et al., 2005, 2006). The overall conformations of MANT-ITP and MANT-GTP show only minimal differences. This result is not surprising. The crystal structures were derived with a racemic mixture of 2'-O- and 3'-O-MANT-ITP. Both structures favor the 3'-O-MANT isomer for binding to the catalytic center as shown with the MANT-GTP and MANT-ATP crystals (Mou et al., 2005, 2006).

A very intriguing finding was the subtle difference in binding mode of MANT-ITP versus MANT-GTP at the purine-binding site in comparison to the exceptionally high inhibitory potency of the nucleotide. Usually, high inhibitory potency is accomplished through strong binding of the inhibitor to the active site of the enzyme. In this case, there were actually fewer protein-ligand interactions due to a missing hydrogen bond in hypoxanthine compared to guanine (Fig. 1 and 3D). However, a gain in affinity is not only related to the number of hydrogen bonds, but also to hydrophobic interactions, residual mobility of the ligand and partial solvation of the binding pocket (Gohlke and Klebe, 2002). The loss of hydrogen bonds does not necessarily lead to a decrease in binding affinity of a ligand to a protein as assessed by molecular thermodynamic and crystallographic studies of thermolysin inhibitors (Morgan et al., 1991). Binding of those inhibitors is not only dependent on hydrogen bonding but also on metal coordination and higher ligand basicity (Grobelny et al., 1989). For binding of MANT-ITP, the absence of the C(2) amino group eliminates a potential hydrogen bond, but at the same time reduces spatial constraints at the purine binding site, thereby allowing other protein-nucleotide interactions to be optimized. Our crystallographic studies and molecular dynamics simulations clearly show that positioning of the β-oxygen of the triphosphate chain of MANT-ITP allows for more favorable interactions with Mn²⁺ than the triphosphate chain of MANT-GTP. providing a straightforward explanation for the observed difference in affinity (Figs. 3 and 5).

Stronger hydrophobic interaction of the MANT-group of MANT-ITP with the hydrophobic pocket compared to MANT-GTP was detected in direct fluorescence experiments and in FRET studies (Fig. 2). In contrast, bindig of MANT-XTP does not result in an increase in direct fluorescence or FRET. Compared to MANT-ITP and MANT-GTP, MANT-XTP is at least 60-fold less potent at VC1:IIC2 (Table 1). MANT-

XTP bears an oxygen at the C2-position of the purine ring (Fig. 1) that interacts unfavourably with D1018 in the base-binding pocket. This may move the MANT-group away from the hydrophobic pocket, leading to a strong decrease in inhibitory potency and fluorescence emmission. As suggested for metal-phosphate interactions, alleviation of binding constraints at the purine-binding pocket may allow MANT-ITP to form more favorable non-polar interactions with the MANT binding site than is possible for MANT-GTP, resulting in larger fluorescence signals

MANT-ITP is the most potent competitive inhibitor of membranous ACs known so far (Table 1) (Göttle et al., 2009). MANT-ITP is most useful as fluorescence probe for biophysical (Fig. 2) and crystallographic studies (Figs. 3 and 4) to obtain detailed molecular information on ligand/receptor interactions. Another application of MANT-ITP is to use this ligand in screening programs for the development of AC inhibitors, avoiding the use of the classic radioactive AC assay with [α - 32 P]ATP as substrate. Specifically, upon binding to the substrate-binding site, non-fluorescent inhibitors would quench the large basal or FS-stimulated direct fluorescence or FRET signals. This assay is even feasible to obtain information on relative inhibitor affinity through the analysis of cumulative concentration/quench curves in a single cuvette (Geduhn et al., 2011). Fluorescence assays with MANT-ITP could also be useful to identify allosteric AC inhibitors.

However, the usefulness of MANT-ITP *per se* as starting point for the development of AC isoform-specific inhibitors, particularly AC5 inhibitors for the treatment of heart failure, ageing, bone loss, anxiety as well as acute and chronic pain (Rottländer et al., 2007; Kim et al., 2008; Chester and Watts, 2007; Okumura et al., 2009) is limited for several reasons. First, MANT-ITP *per se* is not membrane-permeable and would have to be delivered as a prodrug (Laux et al., 2004; Hübner et al., 2011). Second, the selectivity of MANT-ITP for AC5 relative to other AC isoforms

is only very moderate (Göttle et al., 2009). This is due to the fact that the catalytic site of the membranous AC isoforms is highly conserved (Mou et al., 2005). Third, one should also keep in mind that hypoxanthine-based nucleotides do not only bind to mACs but also to other signal-transducing proteins including soluble guanylyl cyclase and G-proteins (Seifert et al., 1999; Gille et al., 2003, 2004, 2005; Hübner et al., 2011). Accordingly, pleitropic effects unrelated to direct mAC inhibition could arise (Hübner et al., 2011). Despite these reservations regarding MANT-ITP, the long-term goal of obtaining isoform-specific AC inhibitors is not elusive. Specifically, we have recently shown that certain bis-MANT-substituted nucleotides are very potent inhibitors of the *Bordetella pertussis* AC toxin, CyaA (Geduhn et al., 2011) with high selectivity relative to mammalian ACs. The identification of Bis-MANT-nucleotides as potent and selective CyaA inhibitors resulted from a relatively small medicinal chemistry program and not from an extensive high-throughput screening effort. Unfortunately, Bis-MANT-nucleotides are not very potent inhibitors of mACs because the catalytic site of these ACs is not spacious enough.

In conclusion, our data confirm the three-site pharmacophore model already postulated in previous studies (Mou et al., 2005, 2006). Our data also show that small differences in ligand structure can have a profound impact on interactions with mAC. The one missing opportunity for hydrogen bonding in MANT-ITP relative to MANT-GTP enhances mobility of the ligand in the catalytic site, thereby facilitating hydrophobic interactions of the MANT-group with surrounding amino acids and optimal positioning of the triphosphate chain to divalent cations. Together, these factors result in exceptionally high-affinity binding of MANT-ITP for mAC.

Acknowledgements

We thank the staff at the Stanford Synchotron Radiation Lightsource SMB-MC 9-1 beamline (Stanford, CA) for their assistance with data collection. Thanks are also due to the reviewers for their helpful critique that stimulated us to resolve the mAC:MANT-ITP crystal structure and to conduct molecular dynamics simulations.

Author Contributions

Participated in research design: Hübner, Mou, Dixit, Lushington, Pinto, Gille, Geduhn,

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Conducted experiments: Hübner, Mou, Pinto, Gille, Seifert

Contributed new reagents or analytic tools: Geduhn, König

Performed data analysis: Hübner, Mou, Dixit, Lushington, Pinto, Gille, Geduhn,

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Wrote or contributed to the writing of the manuscript: Hübner, Mou, Dixit, Lushington,

Pinto, Gille, Geduhn, König, Sprang, Seifert

Conducted molecular dynamics simulations: Dixit, Lushington

Aguired funding for the research: Sprang, Seifert

Directed study and coordinated collaboration of scientists: Seifert

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Footnotes

[†]Supported by the Deutsche Forschungsgemeinschaft [Se 529/5-2] (R.S.) and National Institutes of Health [DK46371] (S.R.S). M. H. was supported by a graduate scholarship from the Elite Network of Bavaria (graduate research scholarship of the Free State of Bavaria).

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Legends for Figures

- Fig. 1. Structure of MANT nucleoside 5'-triphosphates (NTPs). Represented are the MANT nucleotides used for enzymatic studies, fluorescence spectroscopy, crystallography and structure activity evaluation, MANT-ITP, MANT-GTP and MANT-XTP. The MANT-group isomerizes between the 2' and 3'-O-ribosyl function. Note the different substitution of the C2 carbon atom of the purine ring in the various nucleotides.
- **Fig. 2. Fluorescence emission spectra of MANT-GTP, MANT-ITP and MANT-XTP.** Emission at λ_{ex} = 280 nm (λ_{em} = 300 500 nm) and at λ_{ex} = 350 nm (λ_{em} = 370 500 nm) are represented. Experiments were conducted at 25°C. Addition of MANT-nucleotides (1 μM) *blue* lines; subsequent addition of VC1 (5 μM) and IIC2 (25 μM), *green* lines; subsequent addition of FS (100 μM), *red* lines. Dashed lines in panels A-C represent endogenous tyrosine/trypophan fluorescence of VC1:IIC2. Reaction mixtures contained a buffer of 100 mM KCl, 10 mM MnCl₂ and 25 mM HEPES/NaOH, pH 7.4. Three independent experiments with at least two different batches of VC1:IIC2 were performed. A and B, MANT-GTP; C and D, MANT-ITP; E and F, MANT-XTP. Fluorescence intensities are shown in arbitrary units (a.u.). For FRET (panels A, C and E, λ_{ex} = 280 nm), the fluorescence observed with VC1:IIC2 was set to 100%. In direct fluorescence experiments (panels B, D and F, λ_{ex} = 350 nm), the fluorescence observed with MANT-nucleotides alone was set to 100%.
- Fig. 3. Binding mode of MANT-ITP and two Mn²⁺ ions in the catalytic site.

 MANT-ITP and two metal ions are bound in the cleft between the soluble C1a and C2a domains. VC1 and IIC2 are colored *wheat* and *light pink*, respectively. MANT-ITP is shown as stick model, carbon atoms are *cyan*, nitrogen atoms are *dark blue*,

oxygen atoms are red, and phosphorus atoms are green. The two Mn²⁺ ions are shown as orange spheres. A, Difference electron density for 3'-O-MANT-ITP and Mn²⁺. The *lime green* wire represents the |F_o|-|F_c| electron density for MANT-ITP contoured at 2.5 σ . The *blue* wire corresponds to the $|F_o|-|F_c|$ electron density for the two Mn²⁺ ions contoured at 5 σ. The coordinates for the ligands were omitted from the phasing model. The secondary structure elements of the complex are labeled as defined previously (Tesmer et al., 1997). B, Detailed view of substrate binding site of VC1:IIC2 with MANT-ITP:Mn²⁺. The catalytic site of VC1:IIC2 shows MANT-ITP, Aand B- site of two Mn²⁺ ions and the protein residues that are responsible for ligand interaction. The interaction among protein residues and MANT-ITP, Mn²⁺ are shown as gray dashed lines. C, Superimposed crystal structures of 3'-O-MANT-ITP and 3'-O-MANT-GTP. The derived MANT-ITP crystal structure was superimposed and compared with the crystal structure of MANT-GTP, shown as a transparent yellow stick model (Protein Data Bank code 1TL7) (Mou et al., 2005). The protein residues are in almost identical conformation and the inhibitors are situated in the substrate binding pocket in a similar fashion. D, Superimposed purine binding site of 3'-O-MANT-ITP and 3´-O-MANT-GTP. The interaction of the hypoxanthine ring and guanine ring of MANT-ITP and MANT-GTP are shown as black and olive green dashed lines, respectively. The distances of hydrogen bond between the hypoxanthine ring and surrounding protein residues of MANT-ITP are indicated in A. The hydrogen bond between I1019 and the amino group of MANT-GTP is missing in the MANT-ITP structure. K938 and the oxygen of the hypoxanthine ring are further apart. The hypoxanthine ring has less binding constraint in the purine binding pocket in comparison to the guanine ring of MANT-GTP.

Fig. 4. MANT-binding site. A detailed view of the MANT-binding site is depicted. MANT-ITP is shown as a stick model; carbon atoms are *cyan*, nitrogen atoms are *dark blue*, oxygen atoms are *red* and one phosphorus atom is displayed in *green*. VC1 and IIC2 are colored *wheat* and *light pink*, respectively. MANT-GTP is shown as a transparent *yellow* stick model. The carbonyl group of MANT-ITP is in closer contact to N1025, but does not interact with the side chain of N1025 in this orientation. Apart from this, no conformational differences between MANT-ITP and MANT-GTP are detected. However, MANT-ITP might exert stronger hydrophobic interactions due to changes of the relative positions of W1020 and the MANT-group.

Fig. 5. Comparison of the binding of MANT-ITP and MANT-GTP by molecular dynamics simulations. Overlaid graphical representations of the terminal (t = 9.6 ns) time steps for the MANT-GTP (CPK-colored sticks with *green* carbons and *orange* phosphorus atoms) and MANT-ITP (CPK-colored sticks with *cyan* carbons and *tan* phosphorus atoms) interacting with the VC1:IIC2 receptor (*pale green* ribbons) and its cofactor Mn²⁺ ions (*magenta* spheres for MANT-GTP simulation and *purple* for MANT-ITP). Additional details on differences in the interactions of MANT-GTP and MANT-ITP with VC1:IIC2 are provided in Supplemental Figure 1 and Supplemental Tables 1-3.

Table 1. Inhibitory potencies of MANT-NTPs on the catalytic activity of VC1:IIC2

	VC1:IIC2 Mn ²⁺ +	
MANT-nucleotide	FS + G _{sα} -GTPγS	VC1:IIC2 Mn ²⁺ + FS
	<i>K_i</i> (nM)	K_i (nM)
MANT-GTP	18 ± 6.0	130 ± 20
MANT-ITP	0.7 ± 0.1	7.0 ± 3.2
MANT-XTP	1,200 ± 370	4,600 ± 510

Catalytic activities of VC1:IIC2 were determined as described in "Materials and Methods". Reactions were conducted in the presence of 10 mM MnCl₂ and 100 μ M FS in the absence or presence of $G_{s\alpha}$ -GTP γ S. Data were analyzed by non-linear regression to calculate K_i values. The catalytic activity of C1/C2 in the presence of Mn²⁺+ FS+ $G_{s\alpha}$ -GTP γ S with 100 μ M ATP as substrate was 2,700 \pm 350 nmol/mg/min and in the presence of Mn²⁺+ FS, the activity was 300 \pm 110 nmol/mg/min. The K_m values for VC1:IIC2 were previously reported (Mou et al., 2005) for each experimental condition (430 and 620 μ M, respectively) and were used to calculate K_i values from IC_{50} values. Data are the mean values \pm SD of 2-4 independent experiments performed in duplicates with at least two different batches of protein.

Table 2. Summary of crystallography data collection and refinement statistics

Parameters	MANT-ITP:Mn ²⁺	
Cell constants (Å)		
а	117.6	
b	133.4	
c	70.6	
No. of crystals	1	
D _{min} (Å)	3.1	
Average redundancy	3.0 (1.8) ^a	
$R_{\text{sym}}\left(\%\right)^{b}$	17.9 (34.5)	
Completeness (%)	81.4 (55.1)	
(l)/(g)	4.5 (1.7)	
Resolution range for refinement $(\mathring{A})^c$	15-3.1	
Total reflections used	15824	
No.of protein atoms	5645	
No. water molecules	5	
No. ligand atoms	106	
rmsd bond length (Å)	0.007	
rmsd bond angle (°)	1.21	
Rwork (%) ^d	24.1	
R_{free} (%) $^{\text{e}}$	29.4	
Average B-factor (Ų)	45.2	

^a numbers in parentheses correspond to the statistic data from the highest resolution shell.

^b $R_{sym} = \sum_{h} \sum_{i} |I(h) - I(h)_{i}| / \sum_{h} \sum_{i} I(h)_{i}$, where I(h) is the mean intensity after rejections.

^c Due to anisotropy, data with an *l* index greater than 21 were omitted from refinement.

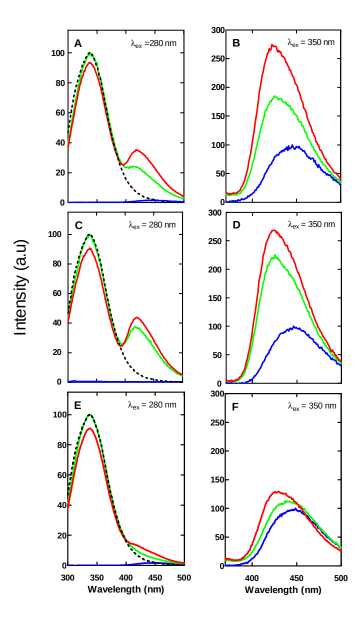
 $^{^{}d}$ $R_{\text{work}} = \sum_{h} ||F_o(h)|| - ||F_c(h)||/\sum_{h} |F_o(h)||$, $F_o(h)$ and $F_c(h)$ are the observed and calculated structure factors, respectively

 $^{^{}m e}$ 5.1% of the complete data set was excluded from refinement to calculate $R_{
m free}$

guanine

hypoxanthine

xanthine



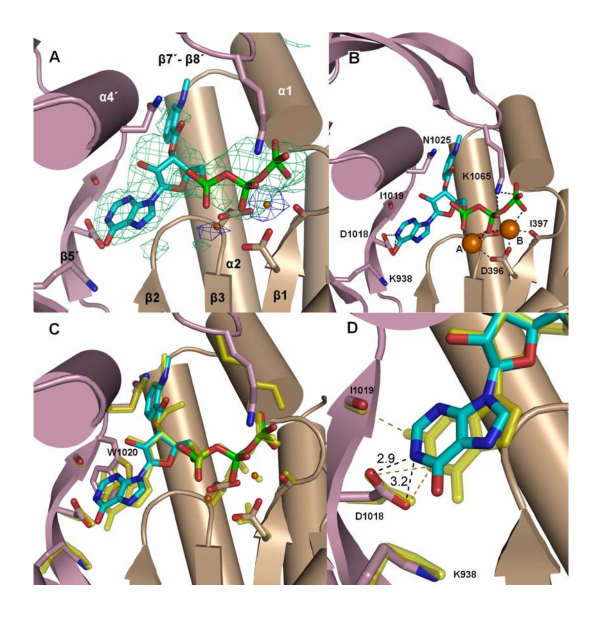
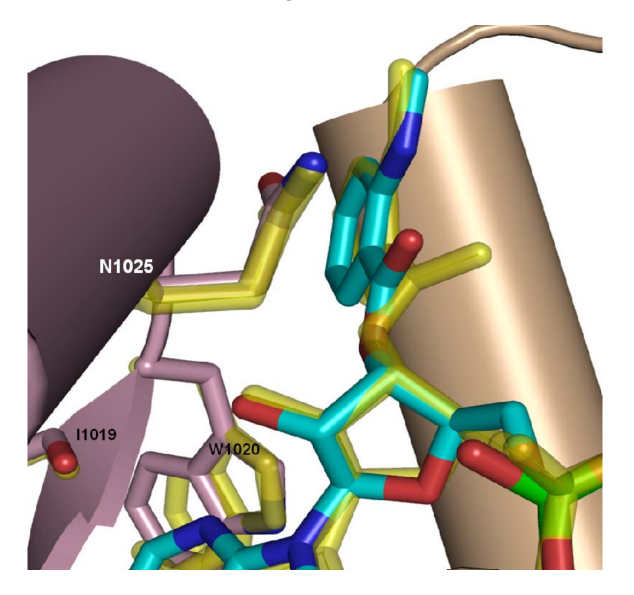
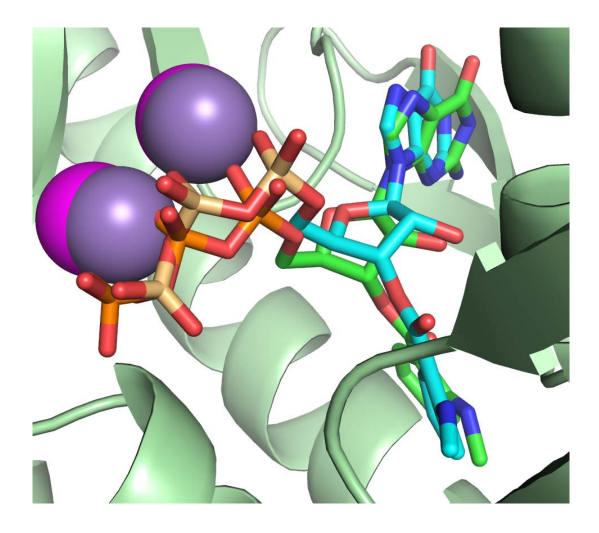


Figure 4





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Supplemental Material

Structural Basis for the High-Affinity Inhibition of Mammalian Membranous Adenylyl Cyclase by 2',3'-O-(N-Methylanthraniloyl)-Inosine 5'-Triphosphate

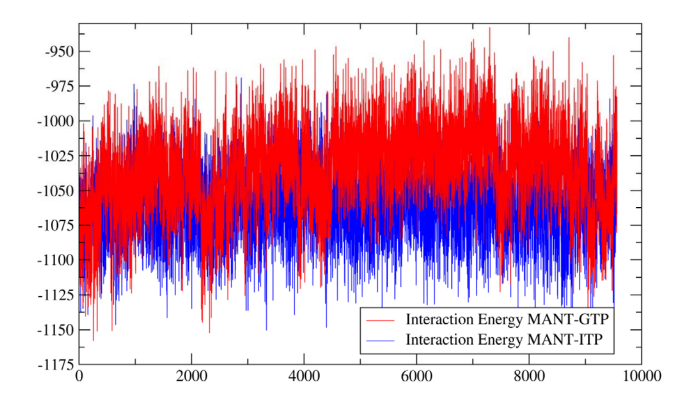
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1. Ligand / VC1:IIC2 interaction energies for MANT-GTP and MANT-ITP via molecular dynamics simulations

The interaction energy ($E_{interaction}$) between ligands (MANT-GTP and MANT-ITP) and rest of the system (protein + water + ions) was calculated for each of the snapshots (~9600) as implied in NAMD2.7. It is the difference of the sum of van der Waals and electrostatic energies of the complex ($E_{complex}$), ligand (E_{ligand}) and the rest of the system ($E_{protein+Water+ions}$).

Thus E_{interaction} = E_{complex} - E_{ligand} - E_{rest}

This was calculated on each frame of the ~9600 frame trajectories and, therefore, should give a fairly good estimation of the interaction between the ligand and the rest of the complex (protein + water + ions). The energy values were plotted against simulation time and the two curves obtained for the energies for MANT-GTP and MANT-ITP were superimposed (*Supplemental Figure 1*). The red curve shows the plotted values for simulation with MANT-GTP while the blue curve shows the same for simulation using MANT-ITP. It can be seen from the plot that MANT-ITP has better energies of interaction as compared to MANT-GTP. The average of the interaction energies for MANT-ITP is - 1061.48 kcal/mol while for MANT-GTP it is -1035.10 kcal/mol, thus a difference of 26.38 kcal/mol exists between the two ligands in favor of MANT-ITP in terms of vdW and electrostatic energy. *Supplemental Tables 1 and 2* provide additional details on the differences regarding the interactions of VC1:IIC2 with MANT-GTP and MANT-ITP in molecular dynamics simulations.



Supplemental Figure 1. Ligand-receptor interaction energies for MANT-GTP and MANT-ITP in VC1:IIC2 as computed over time via molecular dynamics simulations.

Supplemental Table 1. MANT-GTP ligand-receptor interactions from molecular dynamics simulations

Metal-Phosphate interactions (Cutoff: dist. 3.5 Å)

Interaction	Avg. distance	Occupancy
Mn380-ONM_382_O1G	4.445	0
Mn380-ONM_382_O2A	4.599	0
Mn379-ONM_382_O3G	5.874	0
Mn380-ONM_382_O3G	2.256	100
Mn379-ONM_382_O3B	3.293	50.920
Mn379-ONM_382_O2B	4.925	0.010
Mn380-ONM_382_O3A	6.161	0
Mn379-ONM_382_O2G	6.799	0
Mn379-ONM_382_O2A	2.245	100
Mn380-ONM_382_O3B	2.281	100
Mn379-ONM_382_O3A	4.435	0
Mn380-ONM_382_O2B	4.383	0.031
Mn380-ONM_382_O2G	4.071	0.125
Mn379-ONM_382_O1G	7.490	0

#LYS1065-Phosphate Interactions (Salt Bridge)

(Cutoff: dist. 3.2 Å)

Interaction	Avg. distance	Occupancy	
LYS1065-ONM_382_O1G	4.821	0.596	
LYS1065-ONM_382_O2A	6.854	0	
LYS1065-ONM_382_O2B	3.382	73.149	
LYS1065-ONM_382_O2G	3.406	57.267	
LYS1065-ONM_382_O3A	5.868	0.732	
LYS1065-ONM_382_O3B	5.427	0	
LYS1065-ONM 382 O3G	5.490	0	

#Hydrogen_bonds

(Cutoff: dist. 3.0 Å)

Phosphate	oxygens:
_	

donor	acceptor	Occupancy
THR401-Side	ONM382-Side	78.380
THR401-Main	ONM382-Side	46.690
LYS1065-Side	ONM382-Side	32.820
Guanine:		
LYS938-Side	ONM382-Side	39.910
ONM382-Side	ASP1018-Side	23.670

No HB observed with ribosyl part

Supplemental Table 2. MANT-ITP ligand-receptor interactions from molecular dynamics simulations

Metal-Phosphate interactions

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Metal I hospitate interactions		(Cutoff: dist: 5:5 ff)	
Interaction	Avg. distance	Occupancy	
Mn380-MI3_382_O1G	4.239	0	
Mn380-MI3_382_O2A	4.867	0	
Mn379-MI3_382_O3G	5.683	0	
Mn380-MI3_382_O3G	2.298	100	
Mn379-MI3_382_O3B	2.365	100	
Mn379-MI3_382_O2B	4.343	0	
Mn380-MI3_382_O3A	6.183	0	
Mn379-MI3_382_O2G	6.384	0	
Mn379-MI3_382_O2A	2.280	100	
Mn380-MI3_382_O3B	2.381	100	
Mn379-MI3_382_O3A	4.452	0	
Mn380-MI3_382_O2B	4.046	0	
Mn380-MI3_382_O2G	3.956	0	
Mn379-MI3_382_O1G	6.570	0	

#LYS1065-Phosphate Interactions (Salt Bridge) (Cuto

	dist.		

Interaction	Avg. distance	Occupancy	
LYS1065-MI3_382_O1G	5.058	0	
LYS1065-MI3_382_O2A	6.662	0	
LYS1065-MI3_382_O2B	3.218	77.993	
LYS1065-MI3_382_O2G	3.368	61.360	
LYS1065-MI3_382_O3A	5.563	1.770	
LYS1065-MI3_382_O3B	5.463	0	
LYS1065-MI3_382_O3G	5.525	0	

#Hydrogen_bonds

Phosphate oxygens:

(Cutoff: dist. 3.0 A)	3.0 A)
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donor	acceptor	Occupancy
THR401-Side	MI3_382-Side	75.26%
LYS366-Side	MI3_382-Side	33.39%
PHE400-Main	MI3_382-Side	23.48%
THR401-Main	MI3_382-Side	58.84%
Xanthine:		
MI3_382-Side	ASP1018-Side	32.37%
LYS938-Side	MI3_382-Side	36.95%

No HB observed with ribosyl part

2. GBSA calculation

The Generalized Born (GB) model is an approximation to the exact (linearized) Poisson-Boltzmann equation. It is based on modeling the protein as a sphere whose internal dielectric constant differs from the external solvent. GBSA is generalized Born model augmented with the hydrophobic solvent accessible surface area (SA) term. The use of this model with molecular mechanics is known as MM/GBSA. It is among the most frequently used implicit solvent model combinations. This formulation has been shown to successfully identify the native states of short peptides with well-defined tertiary structure. However, in some studies the results of GBSA differ significantly from those produced by explicit solvent and deviate from the protein's native state. Variants of the GB model have also been developed to approximate the electrostatic environment of membranes, which have had some success in folding the transmembrane helices of integral membrane proteins. In the current study we have used a modified generalized Born model reported by Onufrieve et al. (2000), popularly known as OBC model and is suitable for macromolecules.

The GB model approximates Δ Gel by the formula

$$\Delta G_{el} \approx \Delta G_{GB} = -\frac{1}{2} \sum_{ij} \frac{q_i q_j}{f^{GB}(r_{ij}, R_i, R_j)} \left| 1 - \frac{e^{-k f GB}}{\epsilon_w} \right|$$

where r_{ij} is the distance between atoms i and j, R_i is the so-called effective Born radius of atom i, and f^{GB} is

$$f^{GB} = \left[r_{ij}^2 + R_i R_j \exp\left(-r_{ij}^2 / 4R_i R_j\right)\right]^{1/2},$$

The effective Born radius of an atom reflects the degree of its burial inside the molecule. The efficiency of computing the effective radii is an important issue. It has been shown that a good agreement between the GB and PB models can be achieved if the effective Born radii match those computed using the PB approach (Onufriev et al., 2000). The so-called Coulomb field approximation is often used for the calculation of the effective radii and the following expression for effective radii (R_i) can be derived (Onufriev et al., 2000; Bashford and Case, 2000).

$$R_i^{-1} = \rho_i^{-1} - \frac{1}{4\pi} \int_{\text{solute}} \theta(|\dot{r}| - \rho_i) - \frac{1}{r^4} d^3 \dot{r}.$$

The integral is over solute volume excluding a sphere of radius ρ_i .

The AMBER model for the calculation of R_i is as follows:

$$R_i^{-1} = \rho_i^{-1} - \rho_i^{-1} \tan h (\alpha \Psi - \beta \Psi^2 + \gamma \Psi^3)$$

Where ψ =integral above, α , β and γ are the adjustable dimensionless parameters. Within the GB models currently available in Amber, each atom in a molecule is represented as a sphere of radius ρ i with a charge qi at its center; the interior of the atom is assumed to be filled uniformly with material of dielectric constant of 1. The molecule is surrounded by a solvent of a high dielectric ε w (78.5 for water at 300 K). Supplemental Table 3 summarizes the results of the GBSA calculations.

Supplemental Table 3. Ligand-receptor binding energy contributions from GBSA calculations performed on molecular dynamics simulation trajectory

Parameter	MANT-GTP		MANT-ITP	
	mean	stdev	mean	stdev
ELE	-847.41	38.66	-951.66	36.75
VDW	-41.87	6.46	-38.33	6.88
GAS	-889.28	37.07	-989.98	35.01
PBSUR	-7.49	0.13	-7.4	0.18
PBCAL	839.96	34.93	923.88	32.87
PBSOL	832.47	34.91	916.48	32.8
PBELE	-7.45	10.72	-27.77	12.89
PBTOT	-56.81	8.54	-73.5	10.24
GBSUR	-8.73	0.17	-8.61	0.24
GB	828.4	34.54	898.85	30.87
GBSOL	819.68	34.53	890.24	30.79
GBELE	-19	10.43	-52.8	12.71
GBTOT	-69.6	8.94	-99.74	10.29

General strategy

The total free energy of the system $G=G_{pol}+G_{nonpol}+E_{mm}\text{-}TS$

Where,

G_{pol} is estimated by generalized born model.

 G_{nonpol} is estimated by $\gamma SASA \gamma$ is surface tension taken as 0.0072.

SASA is solvent-accessible surface area calculated using LCPO method (Weiser et al., 1999).

E_{mm} is sum of electrostatic (Coulombic), van der Waals (Lennard-Jones), and internal energies (bonds, angles, and dihedrals).

Methodology

The trajectory obtained from the MD run was converted into individual crd files at 100ps interval excluding the initial 1ns of the simulation. Thus total 86 frames were used in GBSA calculation for each simulation. The AMBER parameters (using bonding radii for GBSA calculation) were then generated using tleap and were subsequently used in GBSA calculation. The radii for Mn was set as similar to Mg as there are no parameters available for Mn in GBSA module of AMBER. The change of conformational entropy was not considered.

The following settings were used for the GBSA calculation.

IGB=2 (The OBC model)

GBSA=1 (LCPO method for solvent accessible surface area calculation)

EXTDIEL=78.50 (solvent dielectric)

INTDIEL=1.0 (solute dielectric)

SURFTEN=0.0072

References

Bashford D, and Case DA (2000) Generalized Born models for macromolecular salvation effects. *Annu Rev Phys Chem* **51:**129-152.

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