

Bicarbonate activation of adenylyl cyclase via promotion of catalytic active site closure and metal recruitment

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In an evolutionarily conserved signaling pathway, 'soluble' adenylyl cyclases (sACs) synthesize the ubiquitous second messenger cyclic adenosine 3',5'-monophosphate (cAMP) in response to bicarbonate and calcium signals. Here, we present crystal structures of a cyanobacterial sAC enzyme in complex with ATP analogs, calcium and bicarbonate, which represent distinct catalytic states of the enzyme. The structures reveal that calcium occupies the first ion-binding site and directly mediates nucleotide binding. The single ion-occupied, nucleotide-bound state defines a novel, open adenylyl cyclase state. In contrast, bicarbonate increases the catalytic rate by inducing marked active site closure and recruiting a second, catalytic ion. The phosphates of the bound substrate analogs are rearranged, which would facilitate product formation and release. The mechanisms of calcium and bicarbonate sensing define a reaction pathway involving active site closure and metal recruitment that may be universal for class III cyclases.

The ubiquitous second messenger cAMP regulates a large variety of essential physiological processes such as gene expression, chromosome segregation and cellular metabolism. In mammalian cells, cAMP is synthesized by a family of nine transmembrane adenylyl cyclases (tmACs) and one sAC¹. Unlike tmACs, which localize to the cellular membrane and respond to extracellular stimuli via heterotrimeric G proteins¹, sAC is found in various intracellular compartments such as the mitochondria and the nucleus². Its localization near intracellular cAMP targets is the impetus for current models of second messenger signal transduction, in which cAMP functions as a locally acting signaling molecule²⁻⁴.

sAC is insensitive to the tmAC regulators calmodulin and heterotrimeric G proteins as well as the nonphysiological activator forskolin; instead, sAC senses physiological levels of bicarbonate⁵. Aside from its role as a universal physiological buffer maintaining cellular and extracellular pH, bicarbonate functions as a signaling molecule³, regulating many biological processes in mammals such as fertility⁶, acid-base homeostasis, breathing rate, metabolism and fluid transport (reviewed in ref. 7). As the only known signaling enzyme sensitive to physiological fluctuations of bicarbonate⁵, sAC probably mediates each of these processes. Bicarbonate activation of sAC is essential for sperm motility⁸ as well as for pH-dependent acid secretion in the epididymis and possibly the kidney⁹. In addition to its bicarbonate sensitivity, sAC is synergistically activated by calcium¹⁰, and this potentiation seems to be important for sperm maturation¹¹.

Previous work has revealed the overall structure of tmAC enzymes and suggested a two-metal ion mechanism for catalysis^{12,13}. Despite their different regulation, mammalian sAC and tmACs are grouped into the nucleotidyl cyclase class III based on sequence similarities within their catalytic domains¹⁴. We set out to study the molecular

basis for the unique regulation of sAC enzymes through bicarbonate and calcium. Here, we describe a series of high-resolution crystal structures of the sAC homolog CyaC from the cyanobacterium *Spirulina platensis* in complex with ATP analogs, magnesium, calcium or calcium analogs, and with or without bicarbonate. The structures show that calcium activates sAC enzymes by replacing an active site magnesium ion that coordinates the substrate ATP. In contrast, bicarbonate stimulates sAC activity by inducing a large conformational change that leads to a remodeling of the bound nucleotide. These structures of various enzyme states further suggest general mechanisms for the catalytic pathway of class III nucleotidyl cyclases and their activation.

RESULTS

CyaC as model system for mammalian sAC enzymes

The nucleotidyl cyclase class III includes many bacterial and all known eukaryotic adenylyl and guanylyl cyclases¹⁴. However, the catalytic domains of mammalian sAC are more closely related to bacterial class III ACs than to any other known mammalian cyclases¹⁵. We therefore studied the regulation of the catalytic domain of the sAC homolog CyaC from the cyanobacterium *S. platensis* (26% sequence identity), which carries all sequence properties specific for sAC enzymes¹⁴. Like the mammalian enzyme¹⁰, the catalytic domain of CyaC is stimulated by bicarbonate through an increase in its V_{max} , potentiated by calcium through an increase in the low basal affinity for the substrate ATP typical for sAC enzymes, and is synergistically activated when both compounds are present (Supplementary Fig. 1 online). Therefore, coincident bicarbonate and calcium sensing via cAMP production seems to be an ancient and conserved biological mechanism, and CyaC can serve as a model system for studying sAC activation.

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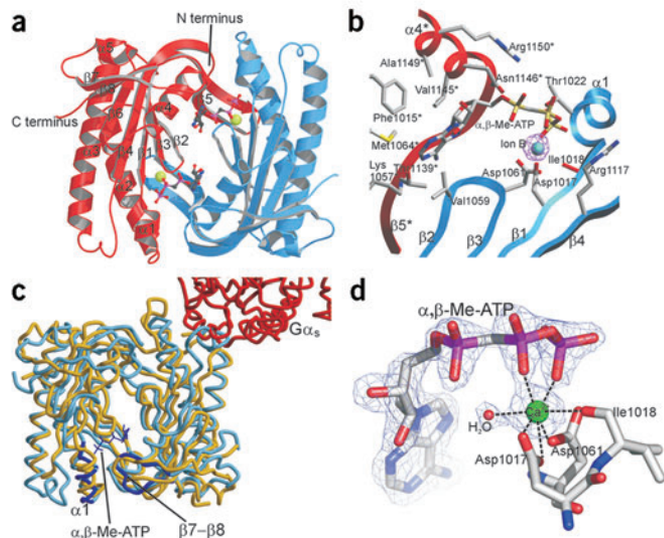


Figure 1 Open conformation of sAC in complex with α,β -Me-ATP and calcium. **(a)** Ribbon diagram of the sAC homodimer with the substrate analog α,β -Me-ATP and one metal ion bound to each active site. The two monomers are red and blue, respectively. **(b)** Active site of the sAC- α,β -Me-ATP complex with the two monomers colored red and blue, respectively. Positive $F_o - F_c$ omit electron density after soaking with the calcium analog europium is overlaid (contoured at 8σ), showing the single heavy atom bound at the ion B site. **(c)** Overlay of sAC- α,β -Me-ATP (blue) with the structure of $G\alpha_s$ -bound tmAC (yellow and red) without substrate analog. sAC helix $\alpha 1$ is shifted to a more open position to avoid clashes with the substrate analog. **(d)** Close-up view of the calcium-binding pocket in the high-resolution structure of the Sr^{2+} -soaked sAC- α,β -Me-ATP complex showing the seven-fold coordination to the metal. The $2F_o - F_c$ omit electron density defining the ligands was contoured at 1.3σ .

α,β -Me-ATP binds to an open enzyme state

To explain the molecular basis of sAC regulation, we determined a series of structures of the CyaC catalytic domain cocrystallized with the ATP analogs α,β -methylene-adenosine-5'-triphosphate (α,β -Me-ATP; $K_i = 0.3$ mM) or adenosine-5'-Rp- α -thio-triphosphate (Rp-ATP α S; $K_i \approx 0.1$ mM), in the presence of magnesium as well as calcium or calcium analogs, and with or without bicarbonate, at a resolution of up to 1.9 \AA (Table 1). Although sACs have unique regulatory properties, the overall structure and active site of CyaC (Fig. 1a,b) closely resemble the known structure of mammalian tmAC¹². The only difference is that CyaC is a symmetrical homodimer of a single catalytic domain with two complete active sites, whereas tmACs are pseudo symmetrical heterodimers of structurally similar catalytic domains (C_1 and C_2), resulting in one active site and one degenerate pseudo active site.

The cocrystal structures of the cyanobacterial sAC (CyaC; from now on referred to as sAC) differed depending on which ATP analog was used. Previously, a $G\alpha_s$ -bound tmAC structure in the absence of nucleotide revealed a more open state than in the presence of Rp-ATP α S, suggesting that substrate binding induces closure of the active site^{12,13,16}. We observed a similar closed conformation for the sAC-Rp-ATP α S complex; however, the conformation of Rp-ATP α S was not suitable for the subsequent in-line reaction (see below), and this state therefore might not resemble the substrate-bound state. The closed enzyme conformation is similar to a tmAC- $G\alpha_s$ complex soaked with product analogs, and it has been speculated that it resembles a product complex and that the substrate complex adopts a different conformation¹². Indeed, we found that in sAC in complex with α,β -Me-ATP (Fig. 1a), the ATP analog assumes a conformation in which all functional groups are arranged for the subsequent in-line reaction¹⁷ (Fig. 1b) and therefore likely resembles the conformation of the bound substrate ATP. The α -phosphate ($P\alpha$) is in minus synclinal position and oriented so that the 3' hydroxyl, as the

subsequent attacking group, and the bond to be broken between the α -phosphorus atom and the bridging function to the β -phosphate ($P\beta$), are arranged in a straight line. The nucleotide conformation also primes the enzyme for catalysis by arranging $P\alpha$ in eclipsed position with $P\beta$, which repels the pyrophosphate to facilitate its release.

In contrast to the sAC-Rp-ATP α S complex, the state of the active site of the sAC- α,β -Me-ATP complex is even more open than in the tmAC- $G\alpha_s$ structure without nucleotide, manifested in a $3\text{--}4\text{ \AA}$ outward shift of helix $\alpha 1$ (Fig. 1c). Modeling an ATP complex based on the sAC- α,β -Me-ATP structure revealed that ATP could adopt a conformation similar to the analog, which fits well into the open enzyme state (Supplementary Fig. 2 online). Therefore, we conclude that nucleotide binding to sAC does not necessitate active site closure; in fact, it seems to precipitate an open state poised for catalysis. The open substrate analog complex observed here for sAC might also apply to tmACs and would explain why soaking or cocrystallization of activated ($G\alpha_s$ -bound) tmAC with α,β -Me-ATP was not successful¹²; in contrast to the basally active sAC enzyme used here, binding of the activator $G\alpha_s$ to tmAC may have induced partial active site closure in the absence of an ATP analog.

Calcium replaces an active site magnesium

The open state has only a single bound metal ion, rather than the two metal ions required for catalysis¹⁶. This ion (positioned analogously to the ion B magnesium in tmAC structures) serves as an anchoring point for ATP by coordinating and stabilizing the $P\beta$ and γ -phosphate ($P\gamma$) of the ATP analog. Soaking with $SrCl_2$ or $EuCl_3$, which are both

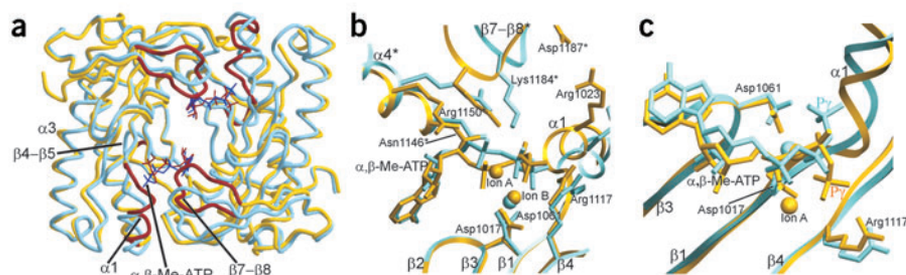


Figure 2 Bicarbonate induces active site closure. **(a)** Overlay of sAC structures in the open state (blue) and after bicarbonate addition (yellow and red). Major conformational changes include shifts of the $\alpha 1$ helix and the $\beta 7\text{--}\beta 8$ strands, a flip of the loop between $\beta 4$ and $\beta 5$ and a kinking of helix $\alpha 3$ (best seen in Supplementary Video 1 online). **(b)** Active site before (blue) and after (yellow) the bicarbonate-induced active site closure, showing the recruited second metal ion and the structural rearrangements described in the text. **(c)** Close-up view for the remodeling of $P\beta$ and $P\gamma$ of the ATP analog upon bicarbonate-induced active site closure (blue before and yellow after bicarbonate addition).

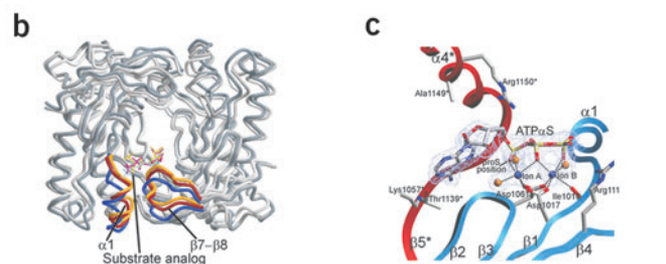
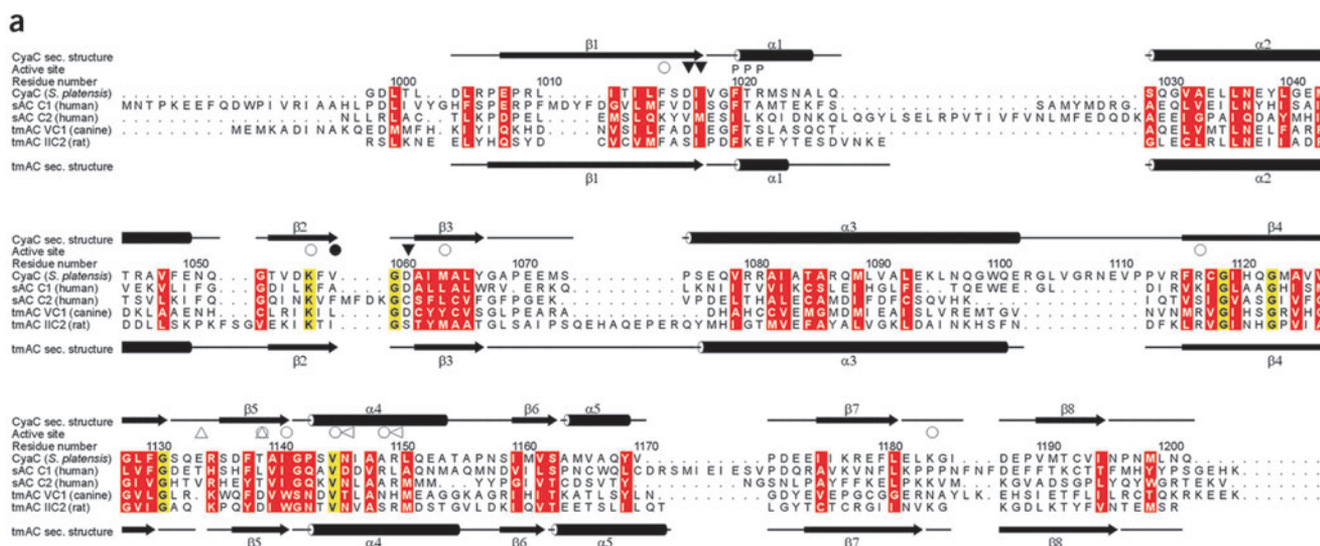


Figure 3 Conformational states and comparison of AC enzymes.

(a) Structure-based sequence alignment of bicarbonate responsive sAC enzymes and the G protein-regulated tmAC domains VC1 and IIC₂ (PDB entry 1A2S). Secondary structure elements of sAC and IIC₂ are indicated on top and bottom, respectively. Ion-binding residues (V) and residues binding the substrate (○) or the transition state (◁) are labeled (filled and empty symbols label C₁ and C₂ residues, respectively). Thr1139* and the insertion characteristic for sAC enzymes are indicated (Δ). Conserved amino acids are shaded yellow, and residues with conserved physicochemical properties are shaded red. (b) Overlay of the sAC-α,β-Me-ATP structure (open state, darkest gray, with α1 helix and β7-β8 loop in blue), the sAC-Rp-ATPαS complex (partially closed, middle gray and red), and the bicarbonate-soaked Rp-ATPαS structure (closed, lightest gray and yellow). Structures were superimposed on sAC-α,β-Me-ATP by optimizing positional agreement for residues 1014-1018, 1056-1065, 1117-1126 and 1143-1167 in both subunits. (c) sAC active site in complex with Rp-ATPαS and two magnesium ions, with the two monomers colored red and blue, respectively. The dashed lines indicate the octahedral coordination of the ions through the ATP analog, protein residues and one and two water molecules (gold spheres), respectively. The 2F_o - F_c omit electron density for the ligands was contoured at 1.1 σ. In its tmAC complex, Pα of Rp-ATPαS was modeled differently but with limited electron density for the ribose and its link to the Pα¹⁶, and we speculate that this density might also be interpretable with the inhibitor conformation observed here for its sAC complex.

known to occupy calcium sites^{18,19}, unambiguously identified this ion B site as a calcium-binding pocket (Fig. 1b). We found no allosteric binding sites for calcium, and soaking in the calcium analogs did not lead to any substantial conformational changes. Although calcium is not often found in active sites, the ion B site is a typical calcium-binding site formed by less flexible ligands and few solvent molecules^{19,20}. The main chain carbonyl oxygen of Ile1018, the side chains of Asp1017 and Asp1061, the phosphate oxygen atoms of Pβ and Pγ of the substrate analog, and a single water molecule coordinate the ion B site calcium (Fig. 1d). The bidentate interaction with Asp1017, which results in a seven-fold coordination that is more favorable for calcium than for magnesium, probably explains the higher affinity of calcium for this site. As calcium has also been reported to have a higher affinity for ATP in solution²¹, this dual preference explains how calcium lowers the K_m for ATP in both mammalian and bacterial sACs, an effect necessary for high sAC activity at physiological ATP concentrations¹⁰ because of the enzyme's low basal substrate affinity (K_m = 12.3 mM; Supplementary Fig. 1 online).

Bicarbonate induces active site closure

Unlike calcium's ability to increase substrate affinity, bicarbonate stimulates substrate turnover. Cocrystallization of sAC with bicarbonate did not produce any crystals and soaking bicarbonate into preformed crystals dissolved them, suggesting that bicarbonate caused

a structural change. Like sAC activation in solution⁵ (Supplementary Table 1 online), the bicarbonate effect on sAC crystals is specific and pH-independent and cannot be induced with other anions, such as nitrate or acetate (see Supplementary Fig. 3 and Supplementary Table 1 online). Flash soaking and freezing permitted us to catch a glimpse of the bicarbonate-induced conformational changes. Most markedly, bicarbonate induced closure of the active site and facilitated binding of the second metal ion (ion A), which serves as the catalytic metal (Fig. 2a,b and Supplementary Video 1 online).

Bicarbonate 'closes' the active site mainly by inducing a 4-5 Å movement of the β7-β8 loop toward the dimer center and a shift of the α1 helix in the same direction. This conformation is stabilized by a newly formed salt bridge between Arg1023 within the α1 helix and Asp1187* (asterisk indicates the partner monomer within the dimer) in the β7-β8 loop. The active site closure induced movements of a several active site residues conserved in all class III cyclases. The β7-β8 loop movement shifts Lys1184*, which in turn pushes Arg1150* of the neighboring α4 helix by 6 Å. Arg1150* is positioned by Asn1146* to be oriented toward the ATP ribose 3' hydroxyl group and Pα such that it could stabilize the additional negative charge in the transition state²². The shift of the α1 helix rearranges the phosphate chain of the ATP analog, leading to a 180° flip of Pγ (Fig. 2c). The shift pushes Pγ out of its binding site and orients it toward Arg1117. This residue was previously thought to bind the substrate¹² but now seems to

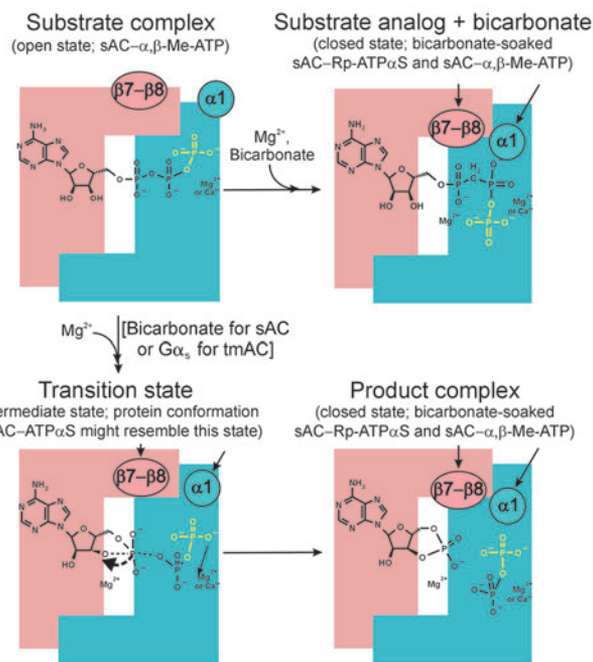


Figure 4 Model for catalysis by class III nucleotidyl cyclases. The model for catalysis (bottom pathway) is based on the conformational changes observed with the sAC-substrate analog complexes (top). The arrows at $\alpha 1$ and $\beta 7$ - $\beta 8$ indicate the movements undergone by these protein parts. The individual catalytic states (open, intermediate and closed) are extrapolated from the different sAC structures presented in the text, with the protein conformation of the sAC-Rp-ATP α S complex being a speculative approximate intermediate state.

Potential bicarbonate recognition sites

Bicarbonate induced these structural changes, but it was not detected in any of the soaked crystals. Its absence, along with the observations that bicarbonate (i) does not disrupt microcrystals of free sAC, and (ii) is very specific (**Supplementary Table 1** online), yet requires high concentrations (physiological concentrations between 5 and 25 mM) to activate sAC^{5,10,24} (corresponding to an estimated off-rate of $\sim 10^6$ s⁻¹, much higher than the turnover rate k_{cat} of ~ 1 to 4 s⁻¹; **Supplementary Methods** online), might suggest that bicarbonate interacts only very transiently with the enzyme during each catalytic cycle.

In the absence of a defined bicarbonate-binding site, sequence differences between tmACs and sACs provide the best insight into the unique bicarbonate regulation of sAC enzymes (**Fig. 3a**). One difference corresponds to sAC active site residue Thr1139*, which is conserved in the bicarbonate-responsive sACs but replaced by a conserved aspartate in bicarbonate-insensitive bacterial ACs and tmACs²⁴. Mutating the analogous threonine to aspartate abolished bicarbonate responsiveness in CyaB1, a cyanobacterial sAC²⁴, consistent with Thr1139* contributing to the bicarbonate stimulation of sACs. A potential scenario is that a single bicarbonate molecule could replace the two water molecules that coordinate ion A (**Fig. 3c**) and facilitate recruitment of this catalytic ion. TmACs with an aspartate replacing sAC Thr1139* would be insensitive to bicarbonate because of the negative charge that decreases the bicarbonate population near the ion A site. However, the effect of bicarbonate on the sAC-Rp-ATP α S complex indicates that ion recruitment can be only part of the bicarbonate function.

A second sequence variation between sACs and tmACs localizes to the $\beta 4$ - $\beta 5$ loop¹⁴, in which bicarbonate-responsive sACs contain a single-residue insertion (**Fig. 3b**). The additional residue causes a different conformation of the $\beta 4$ - $\beta 5$ strands, opening a hole in the back of the enzyme, which may be a conduit for bicarbonate access to the active site. However, we could not confirm the role of this insertion because shortening this loop by one amino acid abolished enzyme activity (data not shown).

DISCUSSION

The structures described here reveal two novel mechanisms of stimulating production of the ubiquitous second messenger cAMP by sAC enzymes. In contrast to its more common role as an allosteric regulator, calcium binds to the active site and directly contributes to substrate binding. We also show that bicarbonate, which was previously only known to serve as a substrate for enzymes, plays an allosteric role by inducing a stimulatory conformational change.

For a model for the dynamic catalytic pathway and the specific activation of all class III adenylyl and guanylyl cyclases (**Fig. 4** and **Supplementary Video 2** online), we assembled the structures presented here. In this model, substrate binding would lead to the formation of an enzyme-substrate complex in the open conformation, with one magnesium bound at the ion B site. In sAC enzymes, this ion can be replaced by calcium to increase substrate affinity. The next step of catalysis is binding of the second metal to the ion A site and the concomitant active site closure. This is the step facilitated by the

attract the reaction product pyrophosphate to aid its exit. These same structural elements were found to be flexible in the G α_s -bound tmAC structures; therefore, we predict that G α_s facilitates similar conformational changes in tmACs as bicarbonate induces in sAC.

Rp-ATP α S binds nonproductively to sAC

As stated above, Rp-ATP α S binds in a nonproductive conformation to a more closed sAC state than α,β -Me-ATP (**Fig. 3b**). This intermediately closed enzyme state resembles G α_s -bound tmAC soaked with Rp-ATP α S¹⁶ and is slightly less closed than the bicarbonate-soaked sAC- α,β -Me-ATP structure. The sAC-Rp-ATP α S complex has two magnesium ions bound, and the electron density clearly reveals that instead of the pro-R oxygen being positioned for coordination of ion A, as observed in the α,β -Me-ATP structure (**Fig. 1b,d**) and as necessary for productive ATP binding, its replacement by sulfur in Rp-ATP α S forces P α to turn such that the pro-S oxygen now coordinates ion A (**Fig. 3c** and **Supplementary Fig. 4** online). This twisted conformation prevents the in-line arrangement of attacking and leaving group, that is, the ribose 3' hydroxyl group and the bridging oxygen between the α - and β -phosphorus atoms. This observation rationalizes why Rp-ATP α S stereospecifically inhibits class III ACs¹⁷, whereas Sp-ATP α S, whose modification does not prevent binding in a productive, α,β -Me-ATP-like conformation, is a substrate for tmACs¹⁷ and likely for sAC enzymes (**Supplementary Fig. 5** online).

The sAC-Rp-ATP α S complex, in addition to being partially closed, has the second metal ion, essential for catalytic activity, bound to the ion A site. During catalysis this ion is recruited after binding of the substrate²³, and this partially closed enzyme state with two bound ions therefore might resemble some features of an intermediate conformation of the enzyme during catalysis. This conclusion is consistent with the successful soaking of Rp-ATP α S into tmAC crystals already partially closed by G α_s ¹⁶. Because of its artificial binding conformation, Rp-ATP α S induced a partially closed state of sAC in the absence of bicarbonate stimulation, and soaking bicarbonate into these crystals resulted in further closure into a structure similar to the bicarbonate-soaked structure of sAC in complex with α,β -Me-ATP (**Fig. 3b**).

Table 1 Data collection and refinement statistics

	CyaC Rp-ATP α S	CyaC α,β -Me-ATP	CyaC α,β -Me-ATP + SrCl ₂	CyaC α,β -Me-ATP + EuCl ₃	CyaC α,β -Me-ATP + bicarbonate	CyaC Rp-ATP α S + bicarbonate
Data collection						
Space group	C222 ₁	P2 ₁ 2 ₁ 2 ₁	P2 ₁ 2 ₁ 2 ₁	P2 ₁ 2 ₁ 2 ₁	P2 ₁	C222 ₁
Unit cell dimensions (Å)						
<i>a</i>	54.2	53.7	53.5	53.6	53.3	51.1
<i>b</i>	78.9	71.5	70.5	70.2	71.2	74.0
<i>c</i>	283.6	99.6	99.4	99.6	106.7	266.7
β (°)	90.0	90.0	90.0	90.0	95.8	90.0
Resolution (Å)	20.0–1.9	20.0–2.4	20.0–1.9	20.0–3.0	20.0–2.3	20.0–2.5
Unique reflections	48,991	15,574	30,126	8,038	33,968	14,233
$\langle I/\sigma \rangle$	9.7	7.0	10.8	5.1	8.9	8.3
Completeness (%) ^a	99.6 (98.6)	99.2 (99.8)	98.8 (94.1)	99.4 (99.2)	95.0 (98.0)	80.0 (69.0)
R_{merge} (%) ^a	7.2 (36.7)	8.1 (34.4)	5.6 (21.7)	13.8 (36.2)	7.0 (42.2)	9.2 (27.7)
Refinement						
Resolution (Å)	15.0–1.93	15.0–2.4	15.0–1.9	15.0–3.0	15.0–2.3	15.0–2.5
Total reflections used	40,595	12,704	25,021	5,601	26,793	11,732
No. atoms						
Protein	4,495	3,042	3,025	3,041	4,510	4,447
Ligand	99	64	64	64	132	99
Solvent	385	141	227	2	99	86
R.m.s. deviations						
Bond lengths (Å)	0.006	0.007	0.006	0.012	0.007	0.010
Bond angles (°)	1.2	1.4	1.2	1.6	1.2	1.5
Average <i>B</i> -factor (Å ²)	30.8	33.9	28.8	—	39.0	29.2
Final $R_{\text{cryst}}/R_{\text{free}}$ (%) ^b	19.8 / 22.9	21.9 / 27.1	20.4 / 23.6	27.4 / 31.5	20.2 / 26.2	24.1 / 29.1

^aValues in parentheses are for the highest-resolution shell. ^b R_{free} was calculated from 6–7% of measured reflections omitted from refinement.

physiological stimulators of ACs, either bicarbonate in the case of sAC enzymes or G α_s proteins for tmACs. Binding of the ion A magnesium enables transition state formation by contributing to the activation of the ribose 3' hydroxyl group and by stabilizing the charges at the triphosphate. The force from the active site closure would shift P γ and P β toward the front, where the products will ultimately exit. Unlike the phosphate flip observed for the α,β -Me-ATP substrate analog, P γ and P β of ATP would move as a rigid body, thereby lengthening and weakening the bond to P α (Fig. 4). The movement of P γ and P β would force P α to swing toward the 3' hydroxyl, leading to ring formation and release of pyrophosphate. The closed conformation observed for the tmAC–G α_s –Rp-ATP α S complex and for the sAC complexes after bicarbonate addition probably corresponds to this product complex with one major exception: the positions of P γ and P β are swapped compared with the product pyrophosphate (Fig. 4). This model is reinforced by the observation that both protein conformation and the position of pyrophosphate in a tmAC–G α_s –product analog structure¹² match those observed in the Rp-ATP α S and bicarbonate-soaked structures. Dissociation of pyrophosphate finally enables the active site to open again by releasing the interactions of the phosphates with the α 1 helix and β 7– β 8 loop.

In summary, the high-resolution structures of different enzyme states presented here predict that class III cyclases might use the same reaction pathway, and that modulators, such as G proteins in tmACs and bicarbonate in sAC, might increase activity by promoting the catalytic cycle via active site closure and metal recruitment.

METHODS

Cloning, protein purification and activity assays. Residues 998–1202 and 1005–1202, respectively, which comprise the catalytic domain of CyaC from *S. platensis* (TrEMBL entry O32393)²⁵, were cloned into the pET28a expression vector with an N-terminal His-tag and expressed in *Escherichia coli* BL21(DE3) for 18 h at 20 °C. The protein was purified using Ni-NTA affinity and Q-Sepharose ion exchange chromatography, followed by gel filtration in 20 mM Tris, pH 7.8. Adenylyl cyclase assays were carried out with purified CyaC(998–1202) as described¹⁰. Protein used for activity assays was stored at –20 °C in 50% (v/v) glycerol.

Crystallization and X-ray data collection. For crystallization of CyaC(998–1202) in complex with Rp-ATP α S, drops were mixed from 1 μ l protein solution (8 mg ml^{−1} in 20 mM Tris, pH 7.8, 5 mM Rp-ATP α S, 7.5 mM MgCl₂, 10 mM CaCl₂) and 1 μ l reservoir solution, equilibrated against 0.4 ml reservoir (100 mM HEPES, pH 7.2, 8% (w/v) isopropanol, 10% (w/v) PEG 4000), and microseeded after 24 h. CyaC(1005–1202) in complex with α,β -Me-ATP was crystallized by mixing 1 μ l protein solution (6 mg ml^{−1} in 20 mM Tris, pH 7.8, 5 mM α,β -Me-ATP, 5 mM MgCl₂, 5 mM CaCl₂) and 1 μ l reservoir solution and equilibration against 0.4 ml reservoir (100 mM cacodylate, pH 6.7, 6% (w/v) isopropanol, 11% (w/v) PEG 4000). Both crystal forms were frozen 60 s after addition of two-drop volumes cryoprotectant solution (25% (v/v) isopropanol in reservoir solution).

For identifying the calcium-binding site, CyaC(1005–1202)– α,β -Me-ATP crystals were soaked with 5 mM EuCl₃ for 23 h and with 10 mM SrCl₂ for 13 h, respectively. Bicarbonate soaking was done by adding 150 mM potassium bicarbonate, pH 7.5, to the cryoprotectant solution and freezing of the crystals after 15 to 25 s. Diffraction data were collected at 100 K at Brookhaven National Laboratory beamline X4A (Table 1) and indexed, scaled and merged with DENZO and SCALEPACK²⁶.

Structure determination. Initial Patterson search trials with the CyaC–Rp-ATP α S data and with a polyaniline model of tmAC VC₁ (PDB entry 1AZS; 20% identity with CyaC) or with conserved side chains included did not yield a prominent solution. We therefore generated a homology model of CyaC from the tmAC VC₁ structure (HM-CyaC). Only after converting HM-CyaC to a polyaniline model were we able to obtain two of the three monomers in the asymmetric unit using MOLREP²⁷: monomer A, which forms a dimer with its symmetry mate along a crystallographic two-fold rotation axis and an isolated monomer B. The missing monomer C expected from Matthews coefficient calculation, crystal packing analysis and the tmAC structure was generated manually by applying two-fold rotation symmetry on B resulting in a BC dimer. A fine grid search for C using Como²⁸ then located the correct orientation and position of this monomer. For rigid body refinement and simulated annealing, all monomers had to be replaced with a tmAC VC₁ model trimmed to include only the conserved core of the protein. Electron density quality was sufficient for model building after being markedly improved through three-fold noncrystallographic symmetry averaging and solvent flattening with DM²⁹. Model building was done using O³⁰, and the model was refined using CNS³¹ with an overall anisotropic B-factor, a bulk solvent correction and individual isotropic Debye-Waller factors. All further complex structures were solved by molecular replacement using MOLREP²⁷. Structural figures were generated with MolScript³² and Raster3D³³ (Figs. 1a,c, 2a and 3a), SETOR³⁴ (Figs. 1b, 2b,c and 3b), and PyMOL (http://www.pymol.org) (Fig. 1d), and the alignment (Fig. 3c) with Alscript³⁵.

Coordinates. The coordinates and structure factors have been deposited in the Protein Data Bank (accession codes: CyaC in complex with Rp-ATP α S, 1WC1; with α , β -Me-ATP, 1WC0; with α , β -Me-ATP and Sr²⁺, 1WC3; with α , β -Me-ATP and Eu³⁺, 1WC4; activated α , β -Me-ATP and Rp-ATP α S complexes after soaking with bicarbonate, 1WC5 and 1WC6, respectively).

Note: Supplementary Information is available on the Nature Structural & Molecular Biology web site.

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COMPETING INTERESTS STATEMENT

The authors declare that they have no competing financial interests.

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