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Origin of asymmetry in adenylyl cyclases: structures of *Mycobacterium tuberculosis* Rv1900c

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Rv1900c, a Mycobacterium tuberculosis adenylyl cyclase, is composed of an N-terminal α/β -hydrolase domain and a C-terminal cyclase homology domain. It has an unusual 7% guanylyl cyclase side-activity. A canonical substratedefining lysine and a catalytic asparagine indispensable for mammalian adenylyl cyclase activity correspond to N342 and H402 in Rv1900c. Mutagenic analysis indicates that these residues are dispensable for activity of Rv1900c. Structures of the cyclase homology domain, solved to 2.4 Å both with and without an ATP analog, form isologous, but asymmetric homodimers. The noncanonical N342 and H402 do not interact with the substrate. Subunits of the unliganded open dimer move substantially upon binding substrate, forming a closed dimer similar to the mammalian cyclase heterodimers, in which one interfacial active site is occupied and the quasi-dyad-related active site is occluded. This asymmetry indicates that both active sites cannot simultaneously be catalytically active. Such a mechanism of half-of-sites-reactivity suggests that mammalian heterodimeric adenylyl cyclases may have evolved from gene duplication of a primitive prokaryote-type cyclase, followed by loss of function in one active site. The EMBO Journal (2005) 24, 663-673. doi:10.1038/ sj.emboj.7600573; Published online 27 January 2005 Subject Categories: structural biology; signal transduction Keywords: adenylyl cyclase; crystal structure; half-of-sitesreactivity; Mycobacterium tuberculosis; homodimer

Introduction

Worldwide, *Mycobacterium tuberculosis* is the leading cause of death from a single infectious agent. *M. tuberculosis* invades phagocytic mammalian cells such as phagocytes, monocytes and macrophages, thus evading the host's immune system. Although its ability to survive within infected cells must depend on extensive crosscommunication between host and pathogen, signal transduction pathways of *M. tuberculosis* such as the cyclic nucleotide system have not yet been extensively studied.

Cyclic nucleotides regulate diverse cellular processes (Cooper, 2003). In prokaryotes, cyclic nucleotides play key roles in diverse processes such as adaptation to nutrients, response to osmotic stress and acidification and pathogenicity (Leppla, 1982; Süsstrunk *et al*, 1998; Yahr *et al*, 1998; Ladant and Ullmann, 1999; Kimura *et al*, 2002).

Currently, the catalytic domains of nucleotidyl cyclases, enzymes that catalyze the synthesis of cAMP and cGMP, are grouped into five classes based on sequence similarity (Barzu and Danchin, 1994; Cotta et al, 1998; Sismeiro et al., 1998). Most of them, including mammalian adenylyl cyclases and guanylyl cyclases (GCs), contain class III cyclase homology domains (CHDs), which are further subdivided into four subclasses (IIIa-IIId; Linder and Schultz, 2003). Most mammalian ACs are monomeric integral membrane proteins that possess two complementary, although nonidentical, CHDs (Sunahara et al, 1996; Taussig and Zimmermann, 1998), while prokaryotes and lower eukaryotes produce both soluble and membrane-bound nucleotidyl cyclases of variant domain compositions that usually contain only one CHD (Linder and Schultz, 2003) and are catalytically active as homodimers (Guo et al, 2001; Linder et al, 2002). In eukarvotes, stimuli are usually sensed and transmitted to CHDs by a complex network of regulatory proteins, such as calmodulin, protein kinase C and those involved in the G-proteincoupled receptor (GPCR)-G-protein signal transduction pathways (Cooper, 2003). The GPCR-G-protein pathways are conspicuously absent in prokaryotes. Instead, the prokaryotic CHD isoforms, mostly identified from recently available genome sequences, appear to be directly regulated by a variety of potential regulatory domains attached directly to the CHDs. M. tuberculosis has 15 class III AC genes. Yet, their biological role is unknown. Therefore, characterization of these enzymes is essential for elucidating the role of cyclic nucleotides in *M. tuberculosis* as well as other prokaryotes.

The structures of the G protein- and forskolin-bound complexes of the N-terminal CHD from type V mammalian AC (VC₁) and the C-terminal CHD from type II mammalian AC (IIC₂) demonstrated that the catalytic site is located at the nearly symmetric interface between the CHDs (Tesmer et al, 1997). Because the catalytic domains are not identical, mammalian ACs possess only one functional active site (Dessauer and Gilman, 1996; Whisnant et al, 1996; Yan et al, 1996; Tesmer et al, 1997, 1999, 2000). In contrast, homodimeric class III nucleotidyl cyclases were expected to contain two identical active sites. From the structure of the mammalian AC $VC_1 \cdot IIC_2$ heterodimer, extensive mutational studies and the sequences of CHDs, it was inferred that six residues of the CHD fold are essential for substrate recognition and catalysis (Tesmer et al, 1997, 1999; Linder and Schultz, 2003). In mammalian ACs, these correspond to two Mg^{2+} (or Mn²⁺)-coordinating aspartates (Tang *et al*, 1995; Tesmer et al, 1997, 1999; Tesmer and Sprang, 1998) contributed by

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 C_1 subunits. The C_2 subunits contribute an asparagine (Tesmer *et al*, 1997; Yan *et al*, 1997) involved in substrate orientation, an arginine residue to stabilize the transition state (Tang *et al*, 1995; Tesmer *et al*, 1997; Yan *et al*, 1997) and an adenine-specifying lysine–aspartate couple (Tang *et al*, 1995; Tesmer *et al*, 1997; Sunahara *et al*, 1998). While catalysis in the mycobacterial ACs Rv1625c (Guo *et al*, 2001) and Rv1264 (Linder *et al*, 2002) relies on all of the six residues, several other CHDs from *M. tuberculosis* and other prokaryotes do not have a complete complement of these residues, hence cyclase activity cannot be definitively assigned to these CHDs.

Rv1900c is one of these unconventional cyclase genes of *M. tuberculosis*. The *Rv1900c* gene product encodes a twodomain protein, of which the 285-residue N-terminal domain has sequence similarity to α/β -hydrolases (Cole *et al*, 1998) and the C-terminal residues 291–462 (Figure 1) to class IIIc CHDs (McCue *et al*, 2000; Linder and Schultz, 2003). The presence of the hydrolase homology domain suggested that Rv1900c might possess lipase activity and the enzyme was named 'LipJ' (Cole *et al*, 1998), although consensus residues that confer esterase activity are absent (Shenoy *et al*, 2004a). The Rv1900c CHD shares less than 15 and 28% sequence identity, respectively, with structurally characterized mammalian AC and trypanosomal CHDs, and contains only four of the six residues required for mammalian AC activity. The substrate-orienting asparagine and adenine-specifying lysine are replaced by H402 and N342, respectively (Figure 1). Thus, characterization of the CHD of Rv1900c was expected to provide key insights into variations in substrate binding and catalytic mechanism among nucleotidyl cyclases.

Here we demonstrate Rv1900c is an active AC with an unusually high GC side-activity. We describe the structures of the Rv1900c CHD, the first of a homodimeric AC, in both an open, unliganded conformation and a closed, substratebound conformation. Unlike the mammalian structures (Tesmer *et al*, 1997, 1999, 2000; Zhang *et al*, 1997), which were in a forskolin-activated state, the open Rv1900c homodimer structure appears to depict the ground state. By mutagenesis, we confirmed general determinants of substrate specificity in Rv1900c and marked differences of substrate



Figure 1 Sequence alignment of selected CHDs. Sequences are as follows: *M. tuberculosis* Rv1900c (homodimer), *M. tuberculosis* AC Rv1264 (homodimer), canine AC VC₁ (heterodimer), rat AC IIC₂ (heterodimer) and human guanylyl cyclase GC-A (homodimer). Amino-acid numbering is left of each block. Secondary structures (cylinders for helices, arrows for β -strands, lines for coils) above the alignment correspond to *M. tuberculosis* Rv1900c, and those below to rat IIC₂ (Tesmer *et al*, 1997). Eight positions canonically involved in catalysis are shaded. Red: metal-binding aspartates (Rv1900c: D302 and D346); green: substrate specifying (Rv1900c: the unconventional N342 and D395, both dispensable); light blue: transition-state-stabilizing asparagine (Rv1900c: H402, dispensable); dark blue: phosphate coordinating (Rv1900c: R377, R406, K442). Note the distribution of canonical catalytic residues in the VC₁ · IIC₂ heterodimer. The Lys/Glu/polar-X-X-Gly-Asp motif of the β 2– β 3 loop is boxed.

binding by mammalian ACs and noncanonical CHDs. Unexpectedly, although Rv1900c CHDs form homodimers, they are asymmetric in both unliganded and ligand-bound states. These similarities between the homodimeric Rv1900c and heterodimeric mammalian ACs provide a structural and mechanistic basis for the evolution of asymmetry in adenylyl as well as GCs.

Results

Enzymatic activity of Rv1900c

When expressing the Rv1900c holoenzyme in Escherichia coli, only a fraction was produced as a soluble enzyme and purified to homogeneity (see Supplementary data). The specific AC activity was 0.5 µmol/mg/min (Table I). Esterase activity, which was originally anticipated from the sequence of the N-terminal domain, was not detectable using chromogenic substrates (data not shown). The Rv1900c CHD (residues 291-461) and its mutants were expressed as soluble proteins in E. coli and purified to homogeneity (see Supplementary data and Figure 2A). Purified Rv1900c CHD had a specific activity of 1.3 µmol/mg/min using $Mn^{2+} \cdot ATP$ as a substrate (Table I). The enzyme did not accept Mg²⁺ as a metal cofactor (Table I). The activation energy derived from a linear Arrhenius plot was 73 kJ/mol, with maximal activity at 45°C. Rv1900c CHDs dimerized with a $K_{\rm diss}$ of 100 nM, and dimers were detectible as a 46 kDa band even after SDS-PAGE (Figure 2A). Positive cooperativity for cAMP synthesis with respect to ATP (Table I) was consistent with two catalytic sites per functional unit. Like other mycobacterial ACs (Guo et al, 2001; Linder et al, 2002), the specific activity of the Rv1900c CHD depended on protein concentration (Figure 2B), whereas that of a CHD-CHD fusion construct did not (data not shown), confirming that cyclase activity required dimerization. Surprising for an AC, the Rv1900c CHD had considerable GC side-activity with Mn²⁺ · GTP as a substrate, corresponding to 7% of the AC activity (Table I).

 Table I
 Kinetic properties of wild-type and mutant Rv1900c CHD

Tertiary structures of Rv1900c CHDs

The structures of the Rv1900c CHD (Figure 3) and that of the complex with Mn^{2+} and the nonreactive ATP analog α,β methyleneadenosine 5'-triphosphate (AMPCPP) were determined to 2.3 and 2.4 Å resolution, respectively (Table II). The refined model (Table III) of each structure consisted of four Rv1900c monomers per asymmetric unit, arranged as two dimers, AB and CD. The unliganded Rv1900c CHD model (Figures 3 and 4A) included residues 291-457 for all four monomers in the asymmetric unit, and additionally residues 459-462 for monomers B and C. The AMPCPP-bound structure included residues 291-456 for all monomers, except residues 391-393, which were disordered in monomers B and C. A single AMPCPP molecule and one Mn²⁺ ion were bound per dimer. For both unliganded and AMPCPP-bound structures, subunits within dimers are similar, with root mean square deviations (r.m.s.d.) between corresponding $C\alpha$ atoms < 0.9 Å, whereas corresponding subunits in the two dimers of the asymmetric unit are nearly identical, with r.m.s.d. < 0.14 Å (see Supplementary data). Structural



Figure 2 (**A**) Western blot of Rv1900c CHD probed with an affinitypurified polyclonal antibody against the CHD. (**B**) CHD specific activity increases four-fold with protein concentration.

Enzyme	V _{max} (nmol cAMP/mg/min)	GC (%)	SC ₅₀ (µM)	Hill coeff.	$K_{\rm diss}$ (nM)
Holoenzyme	500	NM	590	1.4	190
CHD	1300	7	300	1.2	100
(CHD) ₂	2000	6	350	1.0	NA
N342A	700	8	300	1.1	80
N342K	900	1	420	1.4	130
N342E	500	30	200	1.3	140
D395A	1100	11	80	1.3	< 10
N342A/D395A	900	7	130	1.0	< 10
H402A	900	5	240	1.3	50
H402N	1200	2	400	1.3	50
R406A	110	2	250	1.7	ND
R377A	500	1	530	1.5	160
K442A	1000	1	600	1.4	40
E340A	2000	5	180	1.3	< 10
V341A	800	5	200	1.6	40
T343A	1400	5	180	1.6	30
F348A	1400	7	250	1.5	50

GC: relative guanylyl cyclase side-activity; SC_{50} : substrate concentration at half-maximal velocity; K_{diss} : dissociation constant of the dimer; $(CHD)_2$: two CHDs linked by a tetradecapeptide; NA: not applicable; ND: not detectable; NM: not measured. Kinetic parameters were measured using 100 nM CHD. All experiments to determine kinetic constants were conducted at least twice with duplicate samples. Variation among individual determinations is typically within 10%. Mean values are reported.

differences are confined to flexible loops and terminal residues. The refined models agreed well with the diffraction data and stereochemical criteria (Table III).

The Rv1900c CHD is compact with approximate dimensions of $40 \times 40 \times 30$ Å (Figure 3). Each domain consists of a central seven-stranded β -sheet with strand order $\beta 2$ - $\beta 3$ - $\beta 1$ - $\beta 4$ - $\beta 6$ - $\beta 8$ - $\beta 7$, with four helices, $\alpha 1$ - $\alpha 3$ and $\alpha 5$, on one face of this sheet, and a single helix, $\alpha 4$, and a β -hairpin, formed by COOH-terminal $\beta 4$ residues and $\beta 5$, on the opposite face. The Rv1900c CHD is similar in structure to other CHDs, but as a class IIIc CHD it has a shorter dimerization arm than that of mammalian ACs (Tesmer *et al*, 1997; Linder *et al*, 2002) and lacks the δ -subdomain of the trypanosomal GRESAG proteins



Figure 3 Cartoon of the Rv1900c CHD color-ramped from blue at the N-terminus to red at the C-terminus. Secondary structures are labeled. This figure and other molecular illustrations were made using PYMOL (http://www.pymol.org).

Table II Summary of data collection and phasing statistics

(Bieger and Essen, 2001). Along with palm domains of DNA polymerases, CHDs utilize $\beta\alpha\beta\beta\alpha\beta$ structural motifs to catalyze the formation of 3'–5' phosphodiester bonds (Artymiuk *et al*, 1997). In the Rv1900c CHD, this motif corresponds to $\beta2-\alpha2-\beta1-\beta3-\alpha3-\beta4$.

Table III	Refinement	statistics
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	Аро	+ AMPCPP
Residue range		
Monomer A	291-457	291-456
Monomer B	291-462	291-390, 394-456
Monomer C	291-462	291-390, 394-456
Monomer D	291-457	291-456
AMPCPP molecules	0	2
Metal ions	0	2
Water molecules	153	223
Side chains with two conformers	6	0
Data range (Å)	50.0-2.3	50.0-2.4
R_{work} (%)	19.41	23.3
$R_{\rm free}$ (%)	23.11	26.2
Average B-values (A^2)		
Main chain	49.4	39.7
Side chain	60.0	46.6
AMPCPP	—	80.4
Metal	—	110.4
Water	48.8	48.3
All atoms	54.1	38.5
R.m.s. deviations from target va- lues		
Bond lengths	0.011 Å	0.02 Å
Bond angles	1.55°	2.50°
Dihedral angles	23.9°	25.1°
B-factors between bonded	6.9 Å^2	3.4 Å^2
atoms		
Coordinate error (Luzzati, 1952)	0.26 Å	0.31 Å
Ramachandran outliers (Laskowski <i>et al</i> , 1993)	0	0

R (%): *R*-factor = $\sum_{h} |F_{obs}| - |F_{calc}|| / \sum_{h} |F_{obs}|$; test set for R_{free} consisted of 4.8% of data.

	Apo SeMet MAD data collection			+ AMPCPP
	λ_1 (peak)	λ_2 (inflection)	λ_3 (remote)	λ (1.0372 Å)
Data				
Data range (Å)	50-2.7	50-2.3	50-2.5	50-2.2
Mosaicity	0.34-0.70	0.36-0.73	0.35-0.79	0.58-0.88
Unique reflections	33 735	53 752	43131	29 564
Average multiplicity	2.58 (2.28)	2.55 (2.19)	2.57 (2.54)	3.8 (3.0)
Completeness (%)	99.8 (99.6)	99.1 (97.3)	99.7 (99.7)	95.3 (75.9)
$R_{\rm sym}$ (%)	8.4 (32.1)	8.1 (36.9)	7.3 (31.4)	7.1 (29.1)
$\langle I/\sigma_I \rangle$	9.2 (1.5)	12.1 (1.1)	11.7 (1.1)	10.4 (2.3)
Phasing statistics				
Phasing power				
Dispersive—acentric	0.554	0.0	1.463	_
Dispersive—centric	0.497	0.0	1.161	_
Anomalous—acentric	1.043	1.104	0.096	—
R_{cullis} (%)				
Dispersive—acentric	0.820	0.0	0.689	_
Dispersive—centric	0.858	0.0	0.692	_
Anomalous—acentric	0.814	0.807	0.978	_

Values in parentheses pertain to the outermost shell of data; $R_{sym} = \sum_{h,i} |I_{h,i} - \langle I_h \rangle| / \sum_{h,i} I_{h,i}$; phasing power = $\langle |F_H| \rangle / r.m.s.$ lack of closure; $R_{cullis} = r.m.s.$ lack of closure/r.m.s. isomorphous difference.

Quaternary structures of Rv1900c CHDs

In crystals of the unliganded as well as the AMPCPP-bound CHDs, the asymmetric unit consists of two nearly identical, pseudosymmetric, isologous homodimers (Figure 4A and B). In addition, each Rv1900c CHD homodimer is asymmetric in that the conformations of quasi-two-fold-related loops and residue side chains at the AB (or DC) subunit interface are



different (see Supplementary data). Further discussion is focused on the dimer formed by subunits A and B, and residues or secondary structural elements are distinguished by the suffix 'A' or 'B' to denote subunit-specific features. The unliganded, 'open', CHD dimer interface involves 22 residues from each subunit in polar or nonpolar interactions that bury approximately 3150 Å² of accessible surface area. Binding of Mn²⁺-AMPCPP induces dramatic, rigid-body movements of the subunits (Figure 4A) accompanied by conformational changes and repacking of the $\beta 2-\beta 3$, $\beta 4-\beta 5$ and $\beta 7-\beta 8$ loops. Thus, although monomers from the unliganded and AMPCPP-bound structures superimpose well, the dimers do not (r.m.s.d. of about 3.4 Å over 329 superimposable Cα atoms). Relative to the subunits of the open dimer, subunits of the AMPCPP-bound dimer are rotated by 16.6° and translated by 11.4 Å along the rotation axis. In this 'closed' conformation, the buried surface area increases from 3150 to 5480 Å and the number of residues from each subunit involved in interfacial contacts increase from 22 to 36-38 residues. AMPCPP buries another 230 Å². The concentration of positive charges at the subunit interface might destabilize the closed conformation of the CHD dimer in the absence of ligand. The closed homodimer (Figure 4B) resembles the $VC_1 \cdot IIC_2$ heterodimer (Tesmer *et al*, 1999) except that the shorter dimerization arms (β4-β5 hairpins) tilt away from the opposing subunit.

Rv1900c CHD active site

Two quasi-symmetrical active sites are located in a groove at the CHD dimer interface (Figure 4C). In the open homodimer, residues implicated in substrate binding and catalysis from each of the two subunits are too far apart for interactions with AMPCPP (Figure 5A). The extensive rigid-body rearrangement of subunits to the closed homodimer conformation is required to enable them to simultaneously interact with substrate (Figure 5B). Upon binding of Mn^{2+} · AMPCPP, the asymmetry of the homodimer increases as the conformations of the $\beta 2-\beta 3$, β 4- β 5 and β 7- β 8 loops of opposing subunits diverge further to accommodate substrate at one catalytic site (compare Figures 4C and 5B), but exclude the adenine moiety of ATP from the other, quasi-dyad-related site (compare Figures 4C and 5C). At the AMPCPP-containing active site, monomer A provides the two invariant aspartates that coordinate Mn²⁺, while monomer B contributes residues that potentially specify substrate and stabilize the transition state (Figure 5B).

Figure 4 Rv1900c CHD dimers. (A) Superimposed open and closed Rv1900c CHD homodimers viewed down the pseudo-dyad dimer axis of the open form. Monomers A and B of the open dimer are rendered in gray and violet, respectively, while those of the closed dimer are rendered sage and salmon, respectively. (B) Closed Rv1900c CHD dimer viewed down the pseudo-dyad dimer axis. Subunits are colored as in (A). The location of the occupied active site is indicated by the bound Mn²⁺ · AMPCPP rendered in atomic detail, and colored by atom type (O red, N blue, P orange and Mn²⁺ black, and C gray) while the unoccupied, dyad-related site is indicated by an AMPCPP molecule represented as a black stick model. (C) View along the dimer dyad axis showing the dimer active sites. Subunits are colored as in (A). Residues that may play a role in substrate binding or catalysis are rendered in atomic detail and colored by atom type (O red, N blue, P orange and Mn²⁺ black, and C according to protein subunit). Metal coordination is depicted by solid, black lines while dashed, black lines indicate hydrogen bonds.



Figure 5 The CHD active site. All three figures are viewed in similar orientations and depict the three crystallographically captured states of the Rv1900c CHD active site. Atomic details of residues that may play a role in substrate binding or catalysis are shown. Subunits, atoms and bonds are represented as in Figure 3. (A) Detailed view of the ligand-unoccupied active site of the open dimer. Active site residues are loosely and asymmetrically packed. (B) Detailed view of the Mn²⁺-AMPCPP-occupied active site of the closed dimer. Potential hydrogen bonds to the oxygen bridging the ATP α - and β -phosphates (leaving group), which is a nonscissile carbon in AMPCPP, are indicated by dashed, magenta lines. (C) Detailed view of the ligand-unoccupied active site of the closed dimer. Residues that participate in binding substrate are held in inactive conformations by a network of hydrogen bonds.

Metal coordination and binding of the ATP ribose and phosphate

The conformation of the occupied ribose/phosphate-binding site of the closed Rv1900c CHD homodimer is similar to that in the mammalian VC₁ · IIC₂ heterodimer with appropriately



Figure 6 Titration of 100 nM Rv1900c CHD R406A with inactive Rv1900c CHD D302A reconstitutes wild-type AC activity.

located structural equivalents of all residues involved in binding the ATP phosphate (Figures 1 and 5B; Tesmer et al, 1997, 1999, 2000). The AMPCPP γ -phosphate is positioned over the N-terminus of $\alpha 1^A$ and is hydrogen bonded to amides of the P-loop ($\beta 1^A - \alpha 1^A$ loop) and to two basic residues, R377^A in $\beta 4^{A}$ (equivalent to VC₁ R484) and K442^B in the $\beta 7^{B} - \beta 8^{B}$ loop (IIC₂ K1065). The AMPCPP α -phosphate ion pairs with $R406^{B}$ in $\alpha 4^{B}$ (IIC₂ R1029). Mutation of either R377 or K442 increases SC₅₀ for ATP, but barely affects $V_{\rm max}$, whereas mutation of R406 substantially reduces V_{max} without affecting SC50 (Table I). Evidently, these residues have roles with respect to substrate binding and catalysis, which are similar to their counterparts in mammalian ACs. Histidine 402^{B} ($\alpha 4^{B}$) is the topological equivalent of mammalian AC IIC₂ N1025, which is implicated in binding the ATP ribose oxygen and orienting the substrate during catalysis (Tang et al, 1995; Tesmer et al, 1997; Yan et al, 1997). However, H402^B is too distant from the AMPCPP ribose oxygen to form a hydrogen bond and does not participate in catalysis as demonstrated by mutagenesis (Table I). Two invariant aspartates in Rv1900c, D302 A in $\beta2^{A}$ and D346 A in the $\beta2^{A}-\beta3^{A}$ loop, coordinate Mn²⁺ at positions equivalent to the 'B' metal site of mammalian ACs where this function is filled by D396 and D440 of VC1 (Tesmer and Sprang, 1998; Tesmer et al, 1999). In addition, D346^A is also hydrogen bonded to a non-ester oxygen from the ATP α -phosphate. Mutation of either metal-binding aspartate in Rv1900c abolishes catalytic activity (Table I). Inactive D302A mutants are rescued by complementary R406A mutants (Figure 6), confirming that D302 and R406 from different subunits participate in the same interfacial catalytic site. In addition to D302^A and D346^A, the octahedral coordination shell of the Mn²⁺ ion consists of S306^A, the backbone carbonyl of I303^A, two nonester oxygens from the AMPCPP α -phosphate, the oxygen bridging the AMPCPP β - and γ -phosphates and a water molecule. The Rv1900c equivalent to the mammalian AC 'A'

Base specificity in Rv1900c

Rv1900c uses ATP and to a much lesser extent GTP as substrates because its purine-binding site does not provide determinants for specificity. The adenine moiety of AMPCPP is inserted into a pocket formed by $F300^{B}$ $(\beta1^{B}),$ $V549^{B}$ $(\beta3^{B})$ and V601^B ($\alpha 4^{B}$) and is stacked against the peptide plane of G345^A–D346^A ($\beta 2^{A} - \beta 3^{A}$ loop). Rv1900c residues D395^B ($\beta 4^{B} - \beta 3^{A}$ $\beta 5^B$ hairpin) and $N342^B$ ($\beta 2^B \text{--}\beta 3^B$ loop) are topological equivalents of D1018 and K938 in mammalian AC IIC2. The latter residues directly hydrogen bond to adenine N1 and N6 (Tesmer et al, 1997). However, in Rv1900c, neither D395^B nor N342^B hydrogen bonds to the AMPCPP adenine. Individual alanine replacements of N342 or the neighboring residues E340, V341, T343 and F348 of the $\beta 2\text{--}\beta 3$ loop do not substantially affect cyclase activity, whereas replacement of N432 by a charged lysine or glutamate decreases or increases relative GC activity, respectively (Table I). Similarly, mutation of D395A did not affect cyclase activity (Table I). Even the N342A/D395A double mutant had nearly normal cyclase activity (Table I). Thus, the enzymatic data for the mutant enzymes, together with the crystal structure of Rv1900c, indicate that neither of the latter two canonical residues nor their neighbors in the $\beta 2-\beta 3$ loop are crucial for activity. However, residues in this loop do appear to have a role in the half-of-sites reactivity of Rv1900c (see below). The limited preference of Rv1900c for ATP over GTP is probably dictated by general size and electrostatic complementarity of the purine-binding pocket rather than by base-specific hydrogen bonds. The absence of base-specific interactions may explain why Rv1900c is not sensitive to P-site inhibitors (data not shown). In this respect, Rv1900c is quite distinct from the mammalian ACs.

Substrate binding in dyad-related active sites of Rv1900c CHD homodimers

The $\beta_2-\beta_3$ loop at the subunit interface of Rv1900c is the central feature of an ATP-driven structural switch that promotes substrate binding in one catalytic site while occluding the dyad-related site (Figures 4C, 5B and C). A network of hydrogen bonds among D346^B, S306^B, T307^B and H402^A pull the $\beta 2^{B} - \beta 3^{B}$ loop toward the unoccupied and away from the occupied ATP-binding sites (Figure 5B and C). Further, hydrogen bonds among N395^A, T343^B and N342^A position N342^A within 2.0 Å of the potential purine-binding pocket of the unoccupied site and thereby exclude the purine moiety of ATP (Figure 5C). The same interactions withdraw $N342^{B}$ from the quasi-dyad-related adenine-binding pocket and allow substrate to bind (Figure 5B). Binding of the AMPCPP ribose phosphate and Mn²⁺ disrupts hydrogen bonds from D346^A to H402^B and S306^A, induces reorientation of D346^A and stabilizes the alternative conformation of $\beta 2^{A} - \beta 3^{A}$.

Discussion

Variant substrate recognition in Rv1900c

This study presents one of several possible variations to the paradigm of catalysis by CHDs that was originally inferred from analyses of mammalian ACs. Rv1900c is the first example of a variant class III adenylyl cyclase catalytic site to be characterized. Coordination of the metal cofactor and binding of ATP phosphate is similar to that of the mammalian ACs. A remarkable feature of Rv1900c activity is its strict requirement for Mn²⁺. Many bacterial CHDs prefer Mn²⁺ as a cofactor, possibly because the radius of the metal coordination sphere is too large to ligate the smaller Mg^{2+} ion (Coudart-Cavalli et al, 1997; Kasahara et al, 2001). However, mammalian ACs and a few bacterial ACs can also use Mg²⁺, which probably is the physiologically relevant cofactor (Dessauer et al, 1997). Rv1900c and other ACs from M. tuberculosis are among those that require millimolar concentrations of Mn²⁺ for activity (Guo *et al*, 2001; Linder et al, 2002, 2004; Shenoy et al, 2005). It is well possible that this is physiologically relevant as the M. tuberculosis genome contains a putative Mn²⁺ transporter that may be capable of sustaining a sufficiently high concentration of intracellular Mn²⁺ (Cole *et al*, 1998; Reddy *et al*, 2001; Linder *et al*, 2004).

Our mutational and structural analysis demonstrates that the ribose- and purine-binding subsites of Rv1900c differ from those in mammalian ACs. H402, the Rv1900c equivalent of the mammalian AC asparagine that appears to orient the substrate during catalysis, does not contact the substrate analog (Tesmer *et al*, 1997; Yan *et al*, 1997). However, this does not appear to have a major impact on the affinity of Rv1900c for ATP or its rate of catalysis, which are comparable to those for unactivated mammalian ACs (Dessauer *et al*, 1997).

Analyses of nucleotidyl cyclases from different organisms have established that specificity for ATP in ACs is very stringent compared to that for GTP in GCs (Beuve et al, 1993; Coudart-Cavalli et al, 1997; Sunahara et al, 1998; Linder et al, 2000; Guo et al, 2001; Kasahara et al, 2001; Weber et al, 2004). Indeed, Rv1900c is the first example of an AC with significant GC side-activity. In mammalian AC heterodimers, a lysine and an aspartate specify ATP by hydrogen bonds to the adenine N1 and N6, and also sterically prevent GTP binding (Tesmer et al, 1997). Structure-based sequence comparisons with ACs indicated that in GCs these residues are substituted by a glutamate and a cysteine, respectively, which were proposed to interact with the guanine N2 and O6 (Sunahara et al, 1998; Tucker et al, 1998). Rv1900c may be able to use GTP as substrate simply because N342 and D395 do not specify adenine as do their lysine and aspartate counterparts in mammalian AC. Accordingly, mutations of these residues did not dramatically alter activity. In contrast, similar mutations in other adenylyl cyclases completely abolish enzyme activity (Kasahara et al, 2001; Shenoy et al, 2003) and the implementation of the lysine-aspartate couple in GCs switches specificity from GTP to ATP (Sunahara et al, 1998; Tucker et al, 1998; Linder et al, 2000). Therefore, mutational analysis of Rv1900c and other ACs indicates that the canonical lysine-aspartate couple specific to most class III ACs are not the sole determinants of nucleotide specificity, and general determinants such as electrostatic gradients, shape complementarity and size

constraints of the purine-binding pocket also play important roles in dictating nucleotide specificity. It is likely therefore that several of the recently identified CHDs that show deviations in the canonical residues required for catalysis by the mammalian ACs (Linder and Schultz, 2003) are also functional cyclases like Rv1900c, but will display differences in substrate specificity, affinity, rate of catalysis and perhaps in mechanism.

Cooperativity in the catalytic cycle

Although CHD homodimers contain two chemically identical and catalytically competent active sites, the structural snapshots of unliganded and AMPCPP-bound Rv1900c CHD dimers show that, at any instance, only one active site binds. Catalytic activity of the D302A-R406A complementation mutant, which has only one functional active site, is comparable to that of the wild-type enzyme (Figure 6), supporting the structural observation that only one site of the homodimer is catalytically competent at a time. Mechanisms of half-of-sites reactivity have been previously observed in enzymes such as adenylyl transferases (Izard and Geerlof, 1999), tyrosyl-tRNA synthetase (First and Fersht, 1993) and dehydrogenases (Eby and Kirtly, 1976; Nichols et al, 2004), as well as ligandbinding proteins like the aspartate receptors (Biemann and Koshland, 1994). However, unlike these proteins, where halfof-sites-reactivity results in negative cooperativity, Rv1900c shows slight positive cooperativity for product formation with respect to ATP. This cooperativity may arise from productive ATP binding in one site, and perhaps weak and nonproductive binding of ATP as an allosteric effector in the dyad-related site.

Progress through the Rv1900c catalytic cycle appears to require a flexible dimer interface, with the open homodimer conformation predominating in the absence of ligand. Entry of ATP into one of the active sites of the homodimer triggers a cooperative movement of the subunits relative to each other, forming the closed homodimer. The productive site, and the ATP bound within, must undergo additional conformational changes relative to the complex to permit full occupancy of Mn^{2+} at the A-site, as required for catalytic activation of the ribosyl 3'-hydroxyl group for in-line nucleophilic attack on the ATP α -phosphate (Tesmer *et al*, 2000). The pentavalent phosphate transition state or intermediate may be stabilized by R406^B and the catalytic Mn²⁺ ion. As in mammalian ACs, steric conflict with the cyclized nucleotide phosphate may trigger ejection of cAMP and the A-site Mn²⁺ (Tesmer *et al*, 2000). This would allow the $\beta 2^{B} - \beta 3^{B}$ loop and Asn^{342B} to revert to the conformation that is nonproductive for substrate binding at the evacuated site. The release of Mn^{2+} · pyrophosphate then may be the rate-limiting step as in the mammalian ACs, prompted by return of the $\beta 2^{A} - \beta 3^{A}$ loop and D346^A to ligand-unoccupied conformations (Dessauer et al, 1997). We have no proof that release of Mn^{2+} · pyrophosphate from one catalytic site is a prerequisite for productive substrate binding in the dyad-related site. However, the comparable rates of cAMP synthesis in wild-type Rv1900c and the D302A-R406A complementation couple indicate that initiation of a new catalytic cycle within the same active site is not rate limiting. Thus, it is plausible to assume that immediately after product release, the two sites will be indistinguishable and therefore equally likely to bind substrate.

The regulation of the CHD activity may well be exerted by the uncharacterized, N-terminal Rv1900c α/β -hydrolase domain in response to signals from hitherto unidentified ligands or protein partners. It can be imagined that a domain interaction occurs between surface-accessible elements of the CHD such as $\alpha 1$, the $\alpha 1-\alpha 2$ loop, $\alpha 2$ or the dimerization arm and the N-terminal domain. The poor expression of the Rv1900c holoenzyme precluded attempts to obtain the structure of the full AC.

Exploiting asymmetry for CHD regulation

Like the Rv1900c CHD, other homodimeric CHDs may be expected to form asymmetric homodimers as a consequence of the bifunctionality of the $\beta 2-\beta 3$ loop. This loop contains a conserved Lys/Glu/polar-X-X-Gly-Asp sequence, of which the first residue regulates nucleotide base binding or specificity in one active site, while in the quasi-dyad-related site, the peptide plane of the invariant glycine-aspartate stacks against the nucleotide base and the invariant aspartate also coordinates metal-phosphate. However, spatial constraints appear to prevent this loop from participating in both active sites simultaneously, permitting only one active site to bind substrate productively and enforcing conformational and mechanistic asymmetry upon the homodimer. This asymmetry might allow differential interaction of sensory/regulatory domains or small molecules with the two subunits of CHD homodimers, and provide mechanistic checkpoints for regulation of cAMP synthesis.

The asymmetric structural features observed in the Rv1900c CHD seem to have been amplified in the evolution of other cyclases. For example, the pseudoheterodimeric mammalian ACs appear to have evolved to enhance and optimize the inherent asymmetry of the CHD homodimer, using one binding site for catalysis while the quasi-dyadrelated site acquired a regulatory function. The plant diterpene forskolin, a fortuitous activator of all mammalian AC isoforms except type IX (Seamon and Daly, 1981; Tang and Gilman, 1995; Sunahara et al, 1996), binds to this regulatory site. However, it cannot be accommodated in the catalytic site of homodimeric CHDs. In mammalian ACs, structural asymmetry is then further exploited by regulatory proteins such as $G\alpha$ subunits, which preferentially bind at nonoverlapping sites on either C₁ or C₂ domains (Tesmer et al, 1997; Dessauer et al, 1998) and probably act by affecting the interplay between open and closed states (Tesmer et al, 1997) or by disrupting nonproductive or inhibitory interdomain contacts.

Materials and methods

Plasmid construction

The gene Rv1900c (GenBank accession number Z97193) was amplified by PCR from *M. tuberculosis* genomic DNA (gift of Dr Boettger, University of Zürich Medical School), and cloned into pET-16b between 5' *Ndel* and 3' *Xhol* sites, adding an N-terminal MGH₁₀SSGHIEGRH tag. The isolated Rv1900c CHD, residues 291– 462, was cloned into pQE30 between 5' *Bam*HI and 3' *XmaCl* sites, adding an N-terminal MRGSH₆GS tag. Point mutations were introduced by standard methods. In the linked dimmer, the connecting sequence was TRAAGGPPAAGGLE. DNA inserts were verified by double-stranded DNA sequencing.

Expression and purification of Rv1900c proteins

Rv1900c CHD and mutants were expressed in E. coli BL21(DE3)[pRep4]. Expression was induced by 60 µM IPTG (4-20 h, 22°C). Bacteria were washed (50 mM Tris-HCl, 1 mM EDTA, pH 8) and stored at -80° C. Cells from a 200–600 ml culture were suspended in 20 ml lysis buffer (50 mM Tris-HCl, 2 mM 3-thioglycerol, pH 8), sonicated for 30s, and then incubated on ice for 30 min each with 0.2 mg/ml lysozyme and 5 mM $MgCl_2 + 10 \mu g/ml$ DNAseI. After centrifugation (31 000 g, 30 min), 15 mM imidazole pH 8 and 250 mM NaCl (final concentrations) were added to the supernatant. Protein was equilibrated on ice for 60 min with 250 µl Ni²⁺-nitrilotriacetic acid (Ni-NTA)agarose, and then successively washed with 10 ml of lysis buffer containing 15 mM imidazole, 250 mM NaCl and 5 mM MgCl₂ and 5 ml of lysis buffer containing 15 mM imidazole and 5 mM MgCl₂. The protein was eluted with 0.4 ml of elution buffer (37.5 mM Tris-HCl pH 8, 250 mM imidazole, 2 mM MgCl₂ 1.5 mM 3-thioglycerol), and then stored at -20° C in elution buffer containing 20% glycerol.

Expression and purification of the Rv1900c holoenzyme was similar except that *E. coli* BL21(DE3)[pLysS] was used for expression, a French Press was used for lysis, lysozyme and DNAseI treatments were omitted, the supernatant and wash buffers contained 30 mM imidazole and 150 μ l Ni-NTA-agarose was used for purification.

Cyclase assays

AC activity was determined at 37°C for 10 min in 100 µl reactions (Salomon *et al*, 1974) containing 50 mM TES–NaOH pH 8.1, 22% glycerol, 3 mM MnCl₂, 500 µM [α -³²P]ATP and 2 mM [2,8-³H]cAMP. Mutant mixtures were incubated on ice for 10 min prior to the start of the reaction. Standard deviations of the data were usually <10%. GC activity was determined identically using GTP as substrate (Klumpp and Schultz, 1982).

Hydrolase assays

Purified holoenzyme was tested for hydrolysis of *p*-nitrophenyl caproate (4.5 mM), *p*-nitrophenyl acetate (saturated) and *p*-nitrophenyl phosphate (1.35 mM). In all, 14–50 µg/ml enzyme was used in a microtiter plate setup and the product *p*-nitrophenol quantified photometrically at 410 nm for 10–20 min at ambient temperature. Buffers were varied to cover a range of conditions. Esterase assays were performed in 0.09% Triton X-100, 9 mM Tris-HCl pH 7.5–9.0 and 0.9–9 mM CaCl₂, and phosphatase assays in 50 mM buffer (pH 4.5–8.0), supplemented with 1 mM MgCl₂ and 0.1 mM EGTA. *Staphylococcus hyicus* lipase and calf intestinal phosphatase were used as controls.

Western blot analysis

Proteins samples blotted onto PVDF membranes after 15% SDS– PAGE were probed with affinity-purified, anti-Rv1900c CHD, rabbit polyclonal antibodies, followed by peroxidase-conjugated goat antirabbit IgG secondary antibodies (Dianova). Peroxidase was detected using the ECL-Plus kit (Amersham-Pharmacia).

Modified purification protocol for structural analysis

His-tagged selenomethionyl (SeMet) Rv1900c CHD was expressed in *E. coli* strain B834(DE3)[pRep4]. Cultures (1 l) grown at 37°C in minimal media containing selenomethionine were induced with 60 μ M IPTG for 12 h at 20°C. Purification was as described above. A HiPrep 26/10 desalting column (Amersham-Pharmacia) was used for fast buffer exchange to 20 mM Tris–HCl, pH 7.5, 10% w/v glycerol, 1 mM MgCl₂ and 0.05% β -mercaptoethanol. Purified protein was concentrated to 7.5 mg/ml, flash frozen and stored at -180° C.

Crystallization and crystal mounting

SeMet Rv1900c CHD was crystallized at 20°C by hanging-drop vapor diffusion from a 1:1 mixture of protein solution and well solution (100 mM HEPES pH 7.5, 8–16% PEG 1000, 450 mM NaCl). Rod-shaped crystals harvested in 50 µl of well solution were cryoprotected by five successive exchanges of 10 µl of harvesting solution with an equal volume of cryosolution (50% PEG 1000, 100 mM HEPES pH 7.5) every 5 s. The substrate analog AMPCPP was similarly soaked into crystals using cryosolution containing 2 mM AMPCPP and 2 mM Mn²⁺. The CHD could not be cocrystallized with several potential ligands under similar condi-

tions. Cryoprotected crystals were immediately flash frozen in liquid $\mathrm{N}_2.$

Data collection

Data were recorded as 5-s exposures over 0.5° crystal rotation per image on a 3×3 mosaic CCD detector, at a crystal to detector distance of 200 mm at the Structural Biology Center beamline 19ID at the Advanced Photon Source, Chicago. MAD data used to solve the structure were collected from a single SeMet Rv1900c crystal as a single 241.5° sweep at each of three wavelengths selected from X-ray fluorescence spectra, the Se K peak (0.979290 Å), inflection point (0.979399 Å) and a remote wavelength (0.956668 Å). Data at each wavelength were collected from areas separated by ~ 0.1 mm along the long crystal axis. Data from an AMPCPP-soaked crystal were recorded as a single 200° sweep at 1.0372 Å.

Data were processed using HKL2000 (Otwinowski and Minor, 1997) (Table I). Both crystals belonged to space group P2, with P2₁22₁ pseudosymmetry, but were nonisomorphous. Unit cell parameters are a = 90.7 Å, b = 44.4 Å, c = 80.4 Å, $\beta = 90.00^{\circ}$, and a = 91.1 Å, b = 48.9 Å, c = 68.2 Å, $\beta = 90.00^{\circ}$ for the unliganded and AMPCPP-soaked crystals, respectively.

MAD structure determination of the Rv1900c CHD

Eight out of 16 Se sites (figure-of-merit = 0.61) were located from unmerged MAD data using SOLVE (Terwilliger and Berendzen, 1999). Refinement of the Se partial structure and MAD phasing were performed using SHARP (Bricogne and Gilmore, 1990) (Table I). The preliminary C α backbone model for four copies of the Rv1900c polypeptide per asymmetric unit ($V_m = 2.0 \text{ Å}^3/\text{Da}$) was built in the initial 3.0-Å map. A total of 10 cycles of phase refinement and extension via four-fold averaging using noncrystallographic symmetry (NCS) operators determined by superposition of secondary structure elements from each subunit, solvent flattening and histogram matching in the program DM (Cowtan, 1994) yielded an interpretable 2.5-Å map from $|F_{obs}|$, ϕ_{best} .

Molecular replacement solution of Rv1900c- AMPCPP complex

Positions and orientations of monomers A, B and D were determined by molecular replacement in EPMR (Kissinger *et al*, 1999), using a single Rv1900c monomer as a search model (correlation coefficient = 0.474). Monomer C was placed, by applying the NCS operator relating subunits A to B, to monomer D.

Model building and refinement

Models for unliganded and AMPCPP-bound structures were built using O (Jones *et al*, 1991) and refined using CNS (Brunger *et al*, 1998) (Table II). The models were restrained by four-fold NCS during preliminary refinement cycles, but during later cycles, as it became apparent that pairs of subunits (A and D, and B and C) from different dimers were more similar than subunits within a dimer, the protein core was restrained by four-fold NCS and the remainder by two-fold NCS restraints relating AB to DC. The unliganded and AMPCPP-bound structures are available in the Protein Data Bank with accession codes 1YBT and 1YBU, respectively.

Supplementary data

Supplementary data are available at The EMBO Journal Online.

Note added in proof

While this manuscript was under review, Steegborn *et al* published a homodimeric adenylyl cyclase structure from *Spirulina platensis* (Steegborn C, Litvin TN, Levi LR, Buck J, Wu H (2005) Bicarbonate activation of adenylyl cyclase via promotion of catalytic active site closure and metal recruitment. *Nat Struct Mol Biol* **12**: 32–37).

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