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The gene product of a *Trypanosoma equiperdum* ortholog of the cAMP-dependent protein kinase regulatory subunit is a monomeric protein that is not capable of binding cyclic nucleotides

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Abstract

The full gene sequence encoding for the *Trypanosoma equiperdum* ortholog of the cAMPdependent protein kinase (PKA) regulatory (R) subunits was cloned. A poly-His tagged construct was generated [TeqR-like(His)₈], and the protein was expressed in bacteria and purified to homogeneity. The size of the purified TeqR-like(His)₈ was determined to be ~57,000 Da by molecular exclusion chromatography indicating that the parasite protein is a monomer. Limited proteolysis with various proteases showed that the *T. equiperdum* R-like protein possesses a hinge region very susceptible to proteolysis. The recombinant TeqR-like(His)₈ did not bind either [³H] cAMP or [³H] cGMP up to concentrations of 0.40 and 0.65 μ M, respectively, and neither the parasite protein nor its proteolytically generated carboxy-terminal large fragments were capable of binding to a cAMP-Sepharose affinity column. Bioinformatics analyses predicted that the

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Authors' contributions

Conflicts of interest

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J.B. designed research and wrote the paper. J.B., J.C.M., M.C., J.F., C.E.S.-R., V.N., J.L.E., and Y.G. performed research. J.B., J.C.M., C.E.S.-R., and S.S.T. analyzed data. All authors read and approved the final manuscript.

The authors declare that there is no conflict of interests regarding the publication of this article.

Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.biochi.2017.12.010.

carboxy-terminal region of the trypanosomal R-like protein appears to fold similarly to the analogous region of all known PKA R subunits. However, the protein amino-terminal portion seems to be unrelated and shows homology with proteins that contained Leu-rich repeats, a folding motif that is particularly appropriate for protein-protein interactions. In addition, the three-dimensional structure of the *T. equiperdum* protein was modeled using the crystal structure of the bovine PKA R^Ia subunit as template. Molecular docking experiments predicted critical changes in the environment of the two putative nucleotide binding clefts of the parasite protein, and the resulting binding energy differences support the lack of cyclic nucleotide binding in the trypanosomal R-like protein.

Keywords

Trypanosoma equiperdum; Regulatory subunits of the cAMP-dependent protein kinase; Gene cloning; Protein expression in bacteria; Protein purification; Biochemical characterization; Dourine; Trypanosomatid parasites

1. Introduction

Salivarian trypanosomes are intravascular, extracellular kinetoplastid haemoprotozoan parasites of a wide range of mammals including humans. Three species, *Trypanosoma brucei, Trypanosoma evansi* and *Trypanosoma equiperdum*, conform the subgenus Trypanozoon. *T. equiperdum* infects horses and other equids under natural conditions, and causes a venereal disease called dourine [1,2].

Cyclic AMP appears to be crucial in *Kinetoplastida*. For instance, *T. brucei* possess more than 80 different genes for adenylyl cyclases (ACs), the enzyme responsible for synthesizing cAMP from ATP [3–5]. In higher eukaryotes, the majority of the cAMP signals are mediated via the activation of the cAMP-dependent protein kinase (*aka* protein kinase A or PKA) and subsequent phosphorylation of various downstream factors, such as transcription factors, signal transduction components, cytoskeletal proteins or metabolic enzymes [6,7]. In most organisms, PKA is an inactive holoenzyme consisting of two catalytic (C) and two regulatory (R) subunits. PKA activation is achieved by the release of the C subunits from the holoenzyme complex caused by cAMP-induced conformational changes in the R subunits.

Complete or draft genomes of various trypanosomatids have been reported, providing both a huge achievement for trypanosome biology and an opportunity to consider a large amount of questions at the genome level [8–11]. Trypanosomatids possess a large set of protein kinases, comprising approximately 2% of each genome, and suggesting a key role for phosphorylation in parasite biology [12]. The AGC group of kinases is named after the PK<u>A</u>, the cGMP-dependent protein kinase or PK<u>G</u>, and protein kinase <u>C</u> (PKC) families, which are cytoplasmic serine/threonine ATP:-phosphotransferase enzymes that are regulated by cAMP (PKA), cGMP (PKG), or Ca²⁺ and diacylglycerol (PKC). The parasite genomes encode three AGC kinases that are related to the C subunits of PKA (PKAC1, PKAC2, and PKAC3), and a gene sharing high homology to the R subunits of PKA. We have identified a ~57-kDa polypeptide band corresponding to the PKA R subunit-like protein from the *T*. *equiperdum* TeAp-N/D1 isolate [13]. Antibody cross-reactivity clearly demonstrated the

high level of conservation exhibited between the parasite protein and its mammalian counterparts [13]. Here, the *T. equiperdum* PKA R-like protein was cloned, expressed in bacteria, purified to homogeneity and biochemically characterized. Similar to higher eukaryotic PKA R subunits [14–16], this novel parasite protein possessed a hinge region very susceptible to proteolysis with a variety of proteases. However, the *T. equiperdum* PKA R-like protein differed from other PKA R subunits since it was not dimeric and was not capable of binding cyclic nucleotides.

2. Materials and methods

2.1. Materials

Reagents were purchased from the following sources: [2,8-³H] cAMP (33.0 Ci/mmol), [8-3H] cGMP (13.5 Ci/mmol), Perkin-Elmer NEN® Radiochemicals; anti-mouse IgG horseradish peroxidase conjugate, SuperSignalTM West Pico chemiluminescent substrate, Thermo Scientific; mouse monoclonal anti-His tag antibody, antimouse IgG alkaline phosphatase conjugate, benzamidine, iodoacetamide, phenyl methyl sulfonyl fluoride (PMSF), 4-(2-aminoethyl) benzenesulfonyl fluoride hydrochloride (AEBSF), pepstatin, leupeptin, N-a-tosyl-L-lysinyl-chloromethyl ketone hydrochloride (TLCK), N-p-tosyl-Lphenylalaninyl chloromethyl ketone (TPCK), ampicillin, kanamycin, fibrous DEAEcellulose, DEAE-Sepharose, isopropyl-1-thio-β-D-galactopyranoside (IPTG), imidazole, complete and incomplete Freund's adjuvant, lipid A, acetylmuramyl-alanyl-isoglutamine (muramyl dipeptide), bovine serum albumin (BSA), cGMP, TPCK-treated trypsin, TLCKtreated chymotrypsin, thermolysin, Staphylococcus aureus V8 protease, papain, cysteine, Sigma; 8-(2-aminoethylamino) adenosine 3',5'-cyclic monophosphate (8-AEA cAMP), Biolog; N-hydroxysuccinimide (NHS)-activated Sepharose 4 fast flow, Superdex S75 gel filtration column, Superdex 200 gel filtration column, Mono Q HR 16/10 anionic exchange column, GE Healthcare; nitrocellulose (0.45 mm pore size), Pierce; 5-bromo-4-chloro-3indolyl phosphate (BCIP), nitro blue tetrazolium (NBT), Wizard® DNA extraction kit, Promega; OptiPhase Hisafe II (scintillation liquid), LKB; pQETrySystem vector, Qiagen; Yeast extract and tryptone (YT) medium, Luria-Bertani (LB) medium, MP Biomedicals; ProbondTM nickel-chelating resin, Mark12TM unstained standard (molecular weight protein markers), BenchMarkTM pre-stained protein ladder, precast NuPAGE[®] Novex 4e12% Bis-Tris sodium dodecyl sulfate (SDS) gels, Life Technologies; 96-well equilibrium dialyzer with a molecular weight cut off of 10,000 Da, Harvard Apparatus. All other chemicals were of the highest quality grade available.

2.2. Parasites

Cryopreserved samples of *T. equiperdum* from the Venezuelan TeAp-N/D1 isolate (*aka* TEVA1) were expanded in adult Sprague Dawley albino rats [17]. When the number of parasites reached 10^6 trypanosomes/ml, the blood was extracted from the rats by cardiac puncture using 0.5 M EDTA as anticoagulant. Parasites were purified by anion exchange chromatography using a fibrous DEAE-cellulose column [18].

2.3. Cloning of a T. equiperdum ortholog of the PKA R subunits

T. equiperdum genomic DNA was obtained using the Wizard[®] DNA extraction kit, according to the instructions of the manufacturer. Primers were designed to amplify the coding region of the gene for the PKA R-like protein from *T. equiperdum* genomic DNA by means of the polymerase chain reaction (PCR): a forward primer (5'-ACC ATG GCT GAA AAG GGA ACA TCG T -3') with an annealing temperature of 68 °C and a Nco I restriction site (underlined), and a reverse primer (5'-CCT CGA GCG ACT TCC TCC CCT CTG CCC TTA -3') with an annealing temperature of 74 °C and a Xho I restriction site (underlined) were utilized. The PCR reaction parameters were standard (Stage 1: 94 °C for 2 min; Stage 2: 35 cycles of 94 °C for 15 s, 68 °C for 20 s and 68 °C for 2 min; Stage 3: 1 cycle at 68 °C for 5 min). The PCR amplified PKA R-like protein coding region was ligated into the pQE-TrySystem vector, previously digested with the Nco I and Xho I restriction enzymes, upstream of an $8 \times$ His tag region from the vector. The identity of the construct was confirmed by means of diagnostic restriction analysis and DNA sequencing. The amplified coding region sequence of the T. equiperdum PKA R-like protein has been lodged in the NCBI GenBank database (https://www.ncbi.nlm.nih.gov/genbank/) with the Accession No. KJ636459 (NCBI ID: AID53025.1).

2.4. Construction of a site-directed mutant of the T. equiperdum ortholog of the PKA R subunits tagged with 8 His residues at its COOH-terminal [TeqR-like(His)₈]

A poly-His tagged construct of the *T. equiperdum* PKA complete R-like protein was built by modifying its stop codon to a tryptophan codon, and incorporating two additional amino acids followed by 8 His at its COOH-terminus. Thus, the following residues are present at the COOH-terminal end of the generated site-directed TeqR-like(His)₈ mutant: Trp-Leu-Glu-His-His-His-His-His-His-His. The gene sequence for the poly-His tagged R-like protein was then inserted in the pQE-TriSystem vector for its expression in bacterial cells. *Escherichia coli* competent cells (strain M15[pREP4]) were transformed with the TeqR-like(His)₈ DNA construct. Expression of the TeqR-like(His)₈ protein in bacteria was evaluated by western blotting using commercial monoclonal anti-His tag antibodies.

2.5. eterologous overproduction of the TeqR-like(His)₈ protein

YT broth cultures (50 ml) prepared in the presence of 100 mg/ml ampicillin and 50 mg/ml kanamycin were inoculated with a single colony of *E. coli* M15[pREP4] containing the gene sequence for TeqR-like(His)₈ inserted in the pQE-TriSystem vector, and bacteria cells were grown overnight at 37 °C with shaking. The overnight cultures were diluted into 12 flasks of fresh YT broth (1 L each) containing 100 mg/ml ampicillin and 50 mg/ml kanamycin and grown at 37 °C with shaking until cultures reached mid-log phase (OD_{600 nm} 0.6–0.7). The expression of TeqR-like(His)₈ was initiated by the addition of IPTG to a final concentration of 0.5 mM and grown at 16e17 °C with shaking. The broth cultures were allowed to grow for 16–24 h before harvesting the cells by centrifugation (6000×*g*) for 30 min at 4 °C. Bacteria pellets were stored at -20 °C until further use.

2.6. Purification of the TeqR-like(His)₈ protein

The TeqR-like(His)₈ protein was purified by combining the following chromatography steps: 1) a ProBond nickel-chelating affinity resin, 2) two sequential separations throughout a Super-dex 200 gel filtration column, and 3) a Mono Q anionic exchange column. During the purification steps, the purity and integrity of the TeqR-like(His)₈ protein was analyzed using SDS-polyacrylamide gel electrophoresis (SDS-PAGE) and western blots revealed with antipolyHis antibodies.

2.6.1. Affinity chromatography—Bacteria containing the overexpressed TeqRlike(His)₈ protein were resuspended in 300 ml of native lysis buffer [50 mM sodium phosphate (pH 8.0), 500 mM NaCl, 5 mM dithiothreitol (DTT), 10 mM benzamidine, 0.4 mM AEBSF, 1 mM pepstatin, 1 mM leupeptin, 28 μ M TPCK and 28 μ M TLCK] and were disrupted by passing the solution twice through a Microfluidizer high shear fluid processor (model M-110P, Microfluidics, Westwood, USA). The cellular debris was harvested by means of centrifugation (12,000×*g*) for 30 min at 4°C and the supernatant was added to a ProbondTM nickel-chelating resin (3 ml). Binding was allowed to occur overnight with gentle agitation at 4 °C. Following pelleting of the beads at 500×*g* for 5 min, the supernatant (flow through fraction) was removed and the sedimented beads were washed four times with three column volumes of native lysis buffer. The resin was poured into a column, and washed with native lysis buffer containing 20 mM imidazole. Finally, the TeqR-like(His)₈ protein was eluted with native lysis buffer containing 250 mM imidazole.

2.6.2. Gel filtration chromatography—Fractions from the nickel-charged affinity resin containing the TeqR-like(His)₈ protein were pooled, concentrated down to 2 ml, and loaded onto a Superdex 200 gel filtration column using a BioLogic DuoFlowTM chromatography system (Bio-Rad, Hercules, USA) with a 5 ml injection loop. The column was equilibrated and eluted with gel filtration buffer [50 mM 2-(*N*-morpholino)ethanesulfonic acid (MES, pH 5.8), 200 mM NaCl, 2 mM EDTA, 2 mM EGTA and 5 mM DTT]. The peak containing the separated TeqR-like(His)₈ protein was pooled and re-chromatographed on the same Superdex 200 gel filtration column under identical conditions.

2.6.3. Ion exchange chromatography—Fractions from the Superdex 200 gel filtration column were pooled, concentrated down to 2 ml, and loaded on a Mono Q HR 16/10 anionic exchange column using a BioLogic DuoFlowTM low pressure liquid chromatography instrument with a 5 ml injection loop. The column was equilibrated with a low salt buffer [20 mM Tris-HCl (pH 8.0), 2 mM EDTA and 5 mM b-mercaptoethanol]. After injecting the sample containing the TeqR-like(His)8 protein, the column was initially thoroughly washed with the low salt buffer, and then the bound protein was eluted using a 200-ml salt gradient from the low salt buffer to a high salt buffer [20 mM Tris-HCl (pH 8.0), 500 mM NaCl, 2 mM EDTA and 5 mM b-mercaptoethanol].

2.7. Tandem mass spectrometry (MS/MS) analysis of TeqR-like(His)₈ tryptic peptides

The purified TeqR-like(His)₈ protein (5 μ g) was digested with trypsin at a protein:enzyme ratio of 100:1. The resulting peptides were separated using Magic 2002 high-performance

liquid chromatography (HPLC) system (Michrom BioResources, Inc.) and eluted into a LTQ-Orbitrap XL mass spectrometer (Thermo Fisher Scientific, San Jose, CA) using electrospray ionization.

2.8. Circular dichroism (CD) spectroscopy

CD measurements were performed with an Aviv 202 spectropolarimeter (Aviv Biomedical, Lakewood, USA). All the spectra were collected at a constant temperature of 25.1 °C.

2.9. Proteolysis of TeqR-like(His)₈

Proteolysis was carried out at 4 °C or 37 °C for 30 min in 0.05 M ammonium bicarbonate (pH 8.3) using various ratios of protease to substrate (0, 1:500, 1:200, 1:100 and 1:50 w/w). The following proteases were used: trypsin, chymotrypsin, thermolysin, *S. aureus* V8 protease, and papain. Papain was preactivated by incubation for 30 min at 4 °C with 1.1 mM EDTA, 0.067 mM b-mercaptoethanol and 5.5 mM Cys. If the protein was run immediately on polyacrylamide gels, an appropriate aliquot of the sample buffer was added, and the tubes were placed immediately in a boiling water bath for 5 min. The samples were then loaded onto the polyacrylamide gels. For large scale proteolysis and subsequent isolation of the proteolytic fragments, the reaction was terminated by injecting the sample on a Superdex 200 or a Superdex 75 gel filtration column using a BioLogic low pressure liquid chromatography equipment and the same conditions described above.

2.10. Cyclic nucleotide binding assays

Cyclic nucleotide binding was measured by equilibrium dialysis. Equilibrium dialyses were carried out on a single plate Rotator (Harvard Apparatus, Holliston, USA) using a 96-well equilibrium dialyzer with a molecular weight cut off of 10,000 Da. Binding assays were performed in 50 mM Tris-HCl (pH 7.5), 2 mM EDTA, 5 mM b-mercaptoethanol, 1 mg/ml BSA, 30 nM of the purified TeqR-Like(His)₈ protein and increasing concentrations of [³H] cAMP (0e400 nM) or [³H] cGMP (0–650 nM). The binding reaction mixture (200 μ l) containing protein and cyclic nucleotide was placed in one side of the dialysis membrane, and 200 ml of just buffer [50 mM Tris-HCl (pH 7.5), 2 mM EDTA, 5 mM 2-mercaptoethanol and 1 mg/ml BSA] was placed on the opposite side. After an overnight incubation (16 h) at 4 °C, a 20-ml aliquot of each side was transferred into scintillation vials and counted in 6 ml of OptiPhase HiSafe II. Cyclic nucleotide binding to the protein was determined by subtraction of the free radioactive reagent. All reactions were performed in duplicates.

2.11. Preparation of polyclonal antibodies against the purified recombinant TeqR-like(His)₈ protein

The purified recombinant TeqR-like(His)₈ protein was used to produce polyclonal antibodies in ascitic fluid from 10 female BalbC mice. We pre-immunized the mice twice (at 7 and 9 weeks of age) with lipid A and muramyl dipeptide as adjuvants, under conditions that did not involve ascites fluid formation. Then, when the mice were 10 weeks old, we followed the standard protocol described by Tung et al. [19]. The animal protocol employed for the preparation of the antibodies was designed to minimize pain or discomfort to the mice, and was performed following the policies stated in the "National Institutes of Health guide for the care and use of laboratory animals (NIH Publications No. 8023, revised 1978)".

2.12. Production and purification of recombinant bovine PKA R^{I}_{α} subunit, human PKA R^{I}_{β} subunit, and mouse PKA Ca subunit

Escherichia coli BL21 (DE3) cells were transformed with the expression vectors pRSETB-RIa (containing the gene for the wildtype full-length *Bos taurus* PKA R^I*a* subunit); pRSETB-RI β (containing the gene for the wild-type full-length *Homo sapiens* PKA R^{I β} subunit); or pRSETB-Cat (containing the gene for the wild-type fulllength *Mus musculus* Ca subunit). R^{I α}, R^{I β} and C α subunits were expressed in bacteria using LB medium in the presence of ampicillin (100 mg/ml). Bacteria cells were grown at 37 °C with shaking until cultures reached mid-log phase (OD_{600 nm} 0.6–0.7). The expression of the proteins was initiated by the addition of IPTG (0.5 mM) and cell growth was maintained with shaking. Broth cultures were allowed to grow overnight before harvesting the cells by centrifugation (6000×*g*) for 30 min at 4 °C.

R subunit isotypes were purified as previously described [20–22] using 8-AEA cAMP-Sepharose resin and cGMP elution. The Ca subunit was expressed and purified according to Gangal et al. [23]. Either peak I, which contains four phosphorylated residues (Ser¹⁰, Ser¹³⁹, Thr¹⁹⁷, and Ser³³⁸) or peak II, which contains three phosphorylation sites (Ser¹⁰, Thr¹⁹⁷, and Ser³³⁸), was used to form the R^I β_2 C₂ holoenzyme. Briefly, for holoenzyme regeneration, the human R^I β subunit was dialyzed with a 5% molar excess of mouse C subunit in 10 mM MES (pH 6.5), 50 mM NaCl, 5 mM MgCl₂, 0.5 mM ATP, and 5 mM DTT. Following R^I β_2 C₂ reconstitution, the remaining C subunit was removed by gel filtration chromatography on a Superdex 200 column.

2.13. Molecular docking

Modeling of cAMP and cGMP binding to both putative CNB domains of the TeqR-like protein was performed using the Auto-Dock Vina program [24]. Docking to the CNB B was chosen first given that this domain corresponds to the high affinity/low dissociation cAMP binding site in mammalian PKA R subunits. A homology model of the *T. equiperdum* protein was generated using the crystal structure of the PKA R^Ia subunit homodimer from *B. taurus* (PDB ID: 4MX3 [25]) as a framework template, and the SWISSMODEL server [26]. As a control, a model was also built for the PKA R^Ia subunit from *B. taurus* complexed with (Rp)-adenosine 3', 5'-cyclic monophosphothioate (PDB ID: 1NE4, [27]. Since all cyclic nucleotide binding assays were carried out at pH 7.5, the protonation state of the ionizable residues was calculated using the web server H [28]. Ligand binding affinity was estimated by predicting empirical binding energy differences.

2.14. Other procedures

Protein concentration was determined using BSA as protein standard [29]. SDS-PAGE was carried out as reported by Laemmli [30] on 1.5-mm thick precast NuPAGE[®] Bis-Tris slab SDS gels containing a gradient of 4e12% polyacrylamide. Coomasie Blue R-250 was used

for protein visualization on gels. For western blot analyses, proteins separated by SDS-PAGE were electrotransferred from the gels to nitrocellulose sheets as described by Towbin et al. [31]. To prepare 8-AEA cAMP-Sepharose, 100 mmol of 8-AEA cAMP was coupled to 25 ml of NHS-activated Sepharose fast flow using the protocol supplied by the vendor.

3. Results

The DNA database of the T. b. brucei genome project was used to obtain the complete gene sequence encoding for the T. b. brucei ortholog of the PKA R subunits. On the basis of predicted homologies between T. brucei and T. equiperdum genes, appropriate oligonucleotides were designed to amplify the corresponding gene from the genomic DNA of *T. equiperdum*. The resulting PCR fragment was cloned and verified by sequencing. Fig. 1 illustrates that the gene for the *T. equiperdum* PKA R-like protein only possesses one nucleotide difference with respect to the corresponding sequence of the T. brucei Tb11.02.2210 PKA R-like protein [Tb927.11.4610 from the TriTrypDB database (http:// tritrypdb.org); and accession No. AF326975.1, NCBI ID: AAG49383.1 from the NCBI GenBank database (https://www.ncbi.nlm.nih.gov/genbank/)]. Recently, a first draft of the T. equiperdum genome has been reported [11] [Accession No. CZPT02000000 (https:// www.ncbi.nlm.nih.gov/nuccore/)]; however, the last version lodged in the NCBI Nucleotide database does not include the gene encoding for the TeqR-like protein cloned here. The translated amino acid sequence of the parasite R-like protein is also shown in Fig. 1. At the protein level, the *T. equiperdum* protein was 100% identical to the *T. brucei* protein. The open reading frame of the T. equiperdum protein predicts an acidic protein (pI 5.24) of 499 amino acids, with a calculated molecular weight of 56,734.3. As also shown in Fig. 1, the trypanosomal R-like protein contains the inhibitor or pseudosubstrate site that in other eukaryotes is involved in binding and inactivation of the PKA C subunits, and two carboxyterminally located putative cyclic nucleotide binding (CNB) domains containing their presumptive phosphate binding cassette (PBC) consensus motifs.

When the amino acid sequence of the *T. equiperdum* R-like protein was compared with the four human PKA R subunit isoforms, $R^{I}\alpha$, $R^{I}\beta$, $R^{II}\alpha$, and $R^{II}\beta$, it was clear that the parasite protein shares sequence homology with mammalian R subunits (Fig. 2A), especially at the carboxy-terminal region (residues 243–499) holding the hypothetical CNB domains. Although the four human R subunits possess high level of conservation at the aminoterminal region containing the dimerization/docking (D/D) domain, the corresponding region of the *T. equiperdum* R-like protein (residues 1e201) shares very little homology with the mammalian R subunits (Fig. 2A). Additionally, the N-terminal region is unusually long in the parasite protein. The results of the comparison of the domain organization between the *T equiperdum* R-like protein and the PKA R subunits are shown diagrammatically in Fig. 2B. The figure illustrates that in contrast to its mammalian orthologs, the parasite protein lacks the D/D domain at its amino-terminal region.

A recombinant T. *equiperdum* R-like protein containing a poly-tag was generated by sitedirected mutagenesis [TeqR-like(-His)8]. Once the IPTG-dependence on the induction of the expression of the recombinant protein was checked, cells were grown in 12 L of YT culture media. Although some protein remained in the particulate fraction, most of the recombinant

protein was expressed in a soluble manner (Fig. S1A) and was purified to homogeneity by consecutively using the following chromatography steps: 1) a Probond TM nickel-chelating affinity resin, 2) two sequential separations throughout a Superdex 200 gel filtration column, and 3) a Mono Q anionic exchange column (Figs. S1 and S2). A pure polypeptide band that migrated with an apparent molecular mass of ~56 kDa was obtained by SDS-PAGE, which corresponded to the expected size for the T. equiperdum ortholog protein. We obtained a yield of 5–6mg of the recombinant purified protein per liter of media, which was soluble up to concentrations higher than 40 mg/ml.

To verify the identity of the parasite PKA R-like protein, the recombinant polyHis-tagged protein was digested with trypsin and the resulting tryptic peptides were separated by HPLC and analyzed by mass spectrometry. The sequence information that was obtained is summarized in Fig. 1 and Table SI, showing coverage of ~64% of the parasite protein by MS/MS (317/499 residues). The N-terminal region of the protein was almost completely confirmed (residues 4–237) (Fig. 1). In contrast, the C-terminal region of the parasite R-like protein, containing the putative CNB domains, appeared to be more protected to proteolysis by trypsin and, consequently, fewer sequences related to this region were acquired by mass spectrometry (Fig. 1, Table SI).

The conformation of the recombinant TeqR-like(His)₈ protein was analyzed by CD, and the protein appeared to be correctly folded (Fig. 3). A comparison with the CD measurements obtained for the PKA bovine R^I α and human R^I β subunits is also shown (Fig. 3A and B). Using the K2D3 program to estimate secondary structure from CD spectra [32], percentages of 36.78% and 4.36% of α -helix and β -strand, respectively, were predicted for the *T. equiperdum* protein.

A comparative analysis by gel filtration chromatography was carried out using the tetrameric $R^{I}\beta_{2}C_{2}$ holoenzyme, the dimeric $R^{I}\beta_{2}$ subunit, and the monomeric C subunit with native molecular weights of ~172,000, 96,000 and 38,000, respectively. As illustrated in Fig. 4A, these experiments showed that the recombinant parasite R-like protein possesses a size of approximately 60,000 Da. An HPLC separation by gel filtration chromatography of TeqR-like(His)₈ was also performed in the presence of various molecular weight markers (Fig. 4B). The molecular mass of the parasite protein was empirically estimated to be 57,323 Da by plotting the *K*av value of each standard *versus* the logarithm of its molecular weight (Fig. 4B). All these results demonstrated that the *T. equiperdum* R-like protein is a monomer, differing from R subunits of higher eukaryotic cells, which are dimeric.

A calibration curve of $(-\log Kav)^{1/2}$ versus the Stokes radii of each standard was also obtained. The elution volume yielded the $(-\log Kav)^{1/2}$ for TeqR-like(His)₈, and a Stokes radius of 3.45 nm was determined by interpolation (data not shown). The frictional coefficient f/f_o can also be evaluated from the molecular mass and the Stokes radius [33]. Deviations of the ratio of f/f_0 from unity reflect non-spherical molecular shape and/or hydration effects. The calculated frictional ratio for TeqR-like(His)₈ was 1.36. This f/f_o value suggested some asymmetry in the molecule and indicated that the native parasite R-like monomeric protein folds into a conformation that lies in the boundary between globular and moderately elongated, when compared with compact spheres [34]. Since the frictional

coefficient contains information not only about the molecular shape of the protein, but also about the hydration effects, an upper limit on the contributions of hydration can be calculated by assuming that all deviations of f/f_0 from unity are due to hydration. Then, the maximal value of hydration (w_{max}) in grams of H₂O bound per gram of TeqR-like(His)₈ can be calculated using the following equation [35]:

$$w_{max} = \frac{\overline{V}}{\overline{V}_{water}} \left[\left(\frac{R_s}{R_0} \right)^3 - 1 \right]$$

where \overline{V} is the partial specific volume of TeqR-like(His)₈ which is assumed to be 0.703 cm³ g⁻¹, \overline{V}_{water} is the partial specific volume of bound water (1 cm³ g⁻¹), R_s is the experimental Stokes radius, and R_0 corresponds to the minimal radius for a perfect sphere possessing the calculated molecular weight. The w_{max} for TeqR-like(-His)₈ is 1.07 g of H₂O per gram of protein, which deviates from the value of 0.3–0.5 g of H₂O per gram of a typical medium sized compact protein.

Higher eukaryotes R^I and R^{II} subunits contain a hinge region that is very susceptible to proteolysis by a variety of enzymes [14-16]. In order to assess whether the parasite R-like protein also contained this hinge region, and to define which fragments were more stable, limited proteolysis experiments were carried out with trypsin, chymotrypsin, the V8 protease from S. aureus, papaín, and thermolysin at two temperatures, 4 °C and 37 °C. As shown in Fig. 5, the *T. equiperdum* R-like protein appears to possess the characteristic hinge region that has been reported for mammalian R subunits. At 4 °C, enzymatic digestions showed that the protein that originally had a molecular mass of ~56 kDa was predominantly cleaved into two fragments; a large fragment of 31e37 kDa and a small fragment of 21–23 kDa. At 37 °C, a similar proteolytic pattern was observed, especially at low concentrations of the proteases (Fig. 5). However, at higher levels of proteases, the 21-23 kDa fragments were further cleaved into smaller fragments at both temperatures (Fig. 5). Therefore, the resulting 31-37 kDa fragments seemed to be more resistant to proteolysis than the 21-23 kDa fragments as the amount of proteases increased in the reaction mixtures. Given that the T. equiperdum protein contained a polyHis tag at its COOH-terminus, the large proteolytic fragment was identified as the corresponding carboxy-terminal region of the protein by affinity chromatography throughout a Probond TM nickel-chelating resin (data not shown). From these results, we concluded that the C-terminal fragment appears to be more stable than the small N-terminal fragment. These findings coincided with the results previously attained by sequencing using MS/MS (Fig. 1, Table SI). Western blotting showed that both fragments were recognized by polyclonal antibodies raised in mice ascites against the recombinant T. equiperdum R-like protein (data not shown).

Since amino acids 243e354 and 363e475 of the *T. equiperdum* R-like protein appeared to form the putative CNB domains A and B, respectively (Fig. 1), the cyclic nucleotide binding properties of the parasite protein were also measured here. In contrast to the PKA bovine R^Ia and human R^I\beta subunits, which showed dissociation constant (K_d) values for cAMP of 40.1 nM and 56.9 nM, respectively, the *T. equiperdum* R-like protein did not bind [³H] cAMP up to concentrations of 0.4 mM (Fig. 6A). The K_d 's (cAMP) obtained for the

mammalian R^I subunits were comparable to published values (K_d 22 nM for the PKA bovine $R^{I}\alpha$ [36]). The generation of such reproducible digestion patterns using various proteases (Fig. 5), prompted us to purify the resulting proteolytic fragments from the T. equiperdum R-like protein using gel filtration liquid chromatography (data not shown). Similar to the non-digested R-like protein, none of the COOH-terminal proteolytic large fragments containing the putative CNB sites were capable of binding [³H] cAMP (data not shown). The inability to bind cAMP was also analyzed employing chromatography throughout a cAMP-Sepharose affinity column. Neither the entire parasite R-like protein (Fig. 6B) nor the C-terminal large fragments resulting from enzymatic limited proteolysis (data not shown) were able to bind to a cAMP-Sepharose column. These experiments confirmed that the T. equiperdum R-like protein is not capable of binding cAMP, which agrees with the results reported by Shalaby et al. [37] for the *T. brucei* R-like protein that is 100% identical to the *T. equiperdum* protein. Given that Shalaby et al. [37] reported that the T. brucei R-like protein was capable of binding cGMP rather than cAMP, we also evaluated the binding of [³H] cGMP to the *T. equiperdum* R-like protein. As shown in Fig. 6A (inset), the *T. equiperdum* protein was not capable of binding [³H] cGMP up to concentrations of 0.65 mM, which curiously argued against the results reported by Shalaby et al. [37]. In addition, none of the generated C-terminal proteolytic fragments that contain the putative CNB sites were capable of binding [³H] cGMP (data not shown). Our experiments clearly showed that neither the *T. equiperdum* R-like protein nor its large proteolytic fragments were capable of binding either cAMP or cGMP.

The primary structures of the *T. equiperdum* R-like protein and PKA R subunits are homologous, especially at the COOH-terminal region bearing the CNB domains (Fig. 2). On the basis of this homology, the Phyre2 server that is available on the web [38] was employed to predict and analyze the structure of the trypanosome protein. The model retrieved is shown in Fig. 7A. The NH_2 -terminal portion of the trypanosomal R-like protein appears to be unrelated to PKA R subunits (Fig. 7B). Yet, bioinformatics analyses predicted that the carboxy-terminal region of the parasite protein folds in the same way that the corresponding region of all known PKA R subunits from higher eukaryotes (Fig. 7B). Similarly, we carried out a secondary structure prediction bioinformatics analysis of the first 201 residues of the protein using the Phyre2 web portal [38] in order to obtain structural information on the amino-terminal region of the parasite protein. As shown in Fig. S3, the analysis predicted that the N-terminal portion of the parasite R-like protein is enriched in α -helices, and appears to possess a structural unit consisting of a small β -strand followed by an α -helix. When this result was compared with proteins stored in the data bank, the N-terminal sequence of the parasite R subunit had homology with proteins that contained Leu-rich (LRR) repeats or LRR proteins. The LRR motif is a structural fold vastly spread in proteins, which consists of 20-30 residues with a characteristic repetitive sequence pattern that is enriched in Leu [39]. The consensus sequence defining the LRR motif is LxxLxLxxNxL (x any aminoacid), although other hydrophobic residues may substitute Leu and Asn at the consensus positions. In LRR proteins, two or more repetitions in tandem form a horseshoeshaped or curved solenoid type of structure that is particularly appropriate for proteinprotein interactions.

The AutoDock Vina docking program [24] was employed to model how the CNB domains are arrayed in the TeqR-like protein, and to predict differences in the environment that outline the putative binding pockets in the parasite protein. Docking was carried out using the crystal structure of the *B. taurus* PKA $R^{I}\alpha$ subunit as a template. In order to calculate the deviation between the template and the structure modeled by the SWISS-MODEL server [26], a model for the *B. taurus* PKA $R^{I}\alpha$ subunit complexed with the (Rp)-adenosine 3', 5'cyclic monophosphothioate analog of cAMP was also built. A root mean-square deviation of 1.863 Å was found between the two resulting structures of the mammalian proteins. In the case of the $R^{I}\alpha$ subunit, both cAMP (Fig. 8A) and cGMP (Fig. 8B) fitted comfortably and tightly inside the cyclic nucleotide binding clefts. The conserved residues, Glu²⁰² and Arg²¹¹ in CNB domain A and Glu³²⁶ and Arg³³⁵ in CNB domain B, which have been reported to be involved in cyclic nucleotide binding by crystallography studies [40], consistently appeared to participate directly in the binding of both cAMP and cGMP. Crystal structures have shown that Glu residues interact with the 2'eOH of the ribose ring while the Arg residues interact with the negative charge of the phosphate group. In both domains, an aromatic side chain, Trp²⁶² in CNB domain A and Tyr³⁷³ in CNB domain B seemed to stack with the ring of the nitrogenous base moieties, as also has been shown by X-ray diffraction [40]. Interestingly, when photoaffinity labeling with $8-N_3$ cAMP was used to identify residues that are located close to the cAMP binding sites, both Trp²⁶² in CNB site A and Tyr³⁷³ in CNB site B were labeled in $R^{I}a$ [41]. These two aromatic side chains in $R^{I}a$ are optimally aligned [40], which presumably accounts for the exceptionally high efficiency in photolabeling. Molecular predictions illustrated that cAMP (Fig. 8A) and cGMP (Fig. 8B) can also be accommodated into the two CNB sites of the TeqR-like protein. However, chemical ligation appears to be unfavorable in the para-site protein since a loose and weak fit was obtained between the cyclic nucleotide ligands and all the relevant residues that participate in binding, showing much empty space in both binding clefts. Specifically, in the modeled structure for the bovine PKA R^Ia subunit bound to either cAMP or cGMP, all distances were in the range of 1.9–3.4 Å between the side chains of the functional residues and the various moieties of the cyclic nucleotide. In contrast, the same distances were 4.6 Å in the modeled structure of the TeqR-like protein. Moreover, even though the parasite protein possesses residues Glu³⁰⁹ and Glu⁴³³ that in mammalian PKA R subunits are involved in hydrogen bonding with the ribose group, surprisingly, the strongly conserved Arg residue, which makes a salt bridge interaction with the nucleotide cyclic phosphate, is replaced by Thr at position 318 and Asn at position 442 in the CNB domains A and B, respectively. Therefore, the electrostatic/ionic balance within the nucleotide binding pockets is clearly destabilized. In the TeqR-like protein, Trp²⁶² of R^Ia is replaced with Tyr³⁷⁰ in CNB site A, maintaining the aromatic features of this side chain; however, Tyr³⁷³ of R^Ia is swapped by Pro⁴⁸⁰ in CNB site B of the TeaR-like protein, which introduced a kink that affects the structural conformation and chemical nature of this region in the nucleotide binding cleft (Fig. 8). However, Tyr⁴⁸² of the parasite protein appears to be located close to where Tyr³⁷³ of R^Ia is located in the predicted three-dimensional structures (Fig. S4), which is interesting since these residues are not at the same position in the amino acid sequences of the two proteins. Therefore, the aromatic characteristics of this particular section of the CNB domain B also seem to be retained. In addition, the COOH-terminal regions of the mammalian PKA $R^{I}\alpha$ subunit and the trypanosomal R-like protein were super-imposed to

reveal the similarities and differences between both modeled structures within the two putative cyclic nucleotide binding sites (Fig. S5).

Docking results were also used to determine the binding affinities for cAMP and cGMP of the TeqR-like protein and PKA R^Ia subunit CNB domains. While the mammalian protein showed binding energies for cAMP of -9.4 and -9.6 kcal/mol in CNB domains A and B, respectively, the parasite protein showed binding energies for cAMP of -5.9 and -5.5 kcal/mol in CNB domains A and B, respectively. Similarly, the mammalian protein showed binding energies for cGMP of -10.2 and -9.5 kcal/mol in CNB domains A and B, respectively, whilst the parasite protein showed binding energies for cGMP of -6.1 and -5.5 kcal/mol in CNB domains A and B, respectively, whilst the parasite protein showed binding energies for cGMP of -6.1 and -5.5 kcal/mol in CNB domains A and B, respectively. On the basis of the predicted binding affinities, no preference of the TeqR-like protein to bind cGMP over cAMP can be inferred. On top, a loss in binding energy of approximately 4 kcal/ mol was acquired in the two CNB domains of the parasite protein for both cAMP and cGMP, which supports and might account for the lack of cyclic nucleotide binding that was obtained experimentally (Fig. 6).

4. Discussion

In this work, a PKA R-like protein from the Venezuelan TeAp-N/ D1 isolate of *T. equiperdum* was cloned, expressed in bacteria, purified to homogeneity and biochemically characterized. At the protein level, a similarity and identity of 100% was achieved when the R-like proteins from *T. equiperdum* and the TREU 927 strain of *T. b. brucei* were compared. The similarity and identity between the R-like protein from *T. equiperdum* and its counterpart from the *T. evansi* STIB 805 strain stored in TriTrypDB (http://tritrypdb.org) (TevSTIB805.11_01.4760) were 100% and 99.6%, respectively, and only two conservative substitutions were detected at positions 215 and 409 from a total of 499 residues [13].

Similar to mammalian PKA R subunits, this novel parasite protein possessed a hinge region very susceptible to proteolysis with a variety of proteases. Yet, the T. equiperdum R-like protein differed from PKA R subunits since it was not dimeric. This result is in agreement with the fact that the trypanosomal protein bears no identifiable functional D/D domain in its amino acid sequence, justifying the monomeric state of the protein. Another major difference with mammalian PKA R subunits is that the parasite TeqR-like protein did not bind cAMP. In agreement with these results, Calabokis et al. [13] have also shown that a partially purified fraction of the T. equiperdum native PKA R-like protein was not capable of binding cAMP. These findings clearly suggest that the TeqR-like protein is not a direct downstream cAMP effector in this parasite. Absence of cAMP binding is probably caused by replacements on specific amino acids that are functionally essential for the ligation of the nucleotide phosphate in both PBC motifs [13]. It has been reported that the variety observed within PBC sequences in several organisms suggests that the CNB domain has evolved as a scaffold for binding not only cAMP, but also an ample diversity of other ligands [42,43]. Since both CNB sites of the T. equiperdum R-like protein hold non-canonical PBC sequences, these structural motifs might be used to bind a different kind of ligand that has not been identified to date.

Shalaby et al. [37] reported that the *T. brucei* R-like protein was capable of binding cGMP rather than cAMP, yet the *T. equiperdum* R-like protein, which is 100% identical to the *T.* brucei protein, was not able of binding either nucleotide. Thus, a rigorous comparison of our findings with those previously published by Shalaby et al. [37] is essential. Most likely, technical variations in the methodological conditions employed in the two studies might account for the discrepancies that were found. Firstly, Shalaby et al. [37] expressed the T. brucei recombinant protein in bacteria as a glutathione S-transferase (GST)-fusion molecule, but it was all produced as an insoluble protein. Attempts to solubilize the recombinant fusion protein with urea were not successful, and the protein remained insoluble and denatured. Therefore, none of the cyclic nucleotide binding experiments that they performed was actually carried out using the full-length recombinant T. brucei protein. Instead, deletion mutants restricted to only one of the two CNB domains (either CNB A or CNB B) and that carried a V5 immunological tag and a His6-tag at their C-termini, were expressed individually in the Drosophila cell line Schneider 2 (S2) and purified by affinity chromatography using a nickel-chelating resin. These truncated CNB sites A and B constructs were capable of binding cGMP but not cAMP, and exhibited K_d values for cGMP of about 8 µM and 11 µM, respectively [37]. On the basis of these findings, the authors concluded about the cGMP binding ability of the parasite protein. In contrast, our experiments were performed using the full-length T. equiperdum R-like protein containing all its functional domains.

Another dissimilar issue is the concentration of cAMP and cGMP used by Shalaby et al. [37], which was much higher than the concentration of cyclic nucleotides employed in the present study. It is critical to state which is the estimated amount of cyclic nucleotide that is physiologically relevant. Strickler and Patton [44] measured the intracellular concentrations of cAMP in Trypanosoma lewisi, a blood protozoan of rats that undergoes differentiation from a rapidly reproducing form to a nonreproducing form in response to the host antibody ablastin. They found that the concentration of cAMP was different in the two developmental stages of T. lewisi (0.167 µM in the reproducing form versus 0.328 µM in the nonreproducing form). Similar low micromolar levels of cAMP have been found in other cellular systems [45], and multiple studies have established that the basal concentration of cAMP in several cell types is about 1 μ M [46–49], which is well above the reported concentration of cAMP required to half-maximally activate PKA that is in the 100-300 nM range [50,51]. Moreover, intracellular cGMP levels have been reported to be even lower than cAMP levels in certain cells or tissues, for example in cultures of fibroblast cells [52], and the reported concentration of cGMP required to half-maximally activate PKG is also in the nanomolar range (~90–200 nM) [53]. Therefore, we strongly believe that 20 µM cGMP, which is the concentration used by Shalaby et al. [37], is extremely high and nonphysiological, and under these conditions nonspecific binding may occur. Conversely, the cAMP and cGMP concentration range that was assayed in the present manuscript is much lower (up to 0.4 μ M for cAMP and up to 0.65 μ M for cGMP), which lie in the expected physiological range. It is known that ligand binding proteins bind their ligand molecules in a concentration dependent manner, which allows the description of two parameters: i) the maximum number of ligand molecules bound per mole of protein and ii) the K_d of the reversible binding process. So, for a putative cyclic nucleotide binding protein as the T.

equiperdum PKA R-like protein, and under the pretense that the protein is capable of ligating cyclic nucleotides, it would be expected to measure increasing number of ligand molecules bound to the protein as the concentration of the cyclic nucleotide increases. Even if we assume a threshold concentration of ~1 μ M cyclic nucleotide to elicit a response [54], the cAMP and cGMP concentrations used here should be more than enough to evaluate cyclic nucleotide binding on a purified protein that contains two putative CNB domains, and is homologous to the mammalian PKA R sub-units and PKG enzymes. Therefore, the lack of cAMP and cGMP binding to the parasite protein clearly demonstrates that the CNB domains of the TeqR-like protein are defective for cyclic nucleotide binding.

It is commonly known that cAMP and cGMP behaved as cyclic nucleotide analogs in structure-activity studies. PKAs and PKGs are defined as cAMP-dependent protein kinases and cGMP-dependent protein kinases, respectively, on the basis of their affinity for each nucleotide; explicitly, PKA exhibits a >50-fold higher affinity for cAMP than for cGMP and preferentially binds cAMP, similarly, PKG has a >50-fold higher affinity for cGMP than for cAMP and prefer-entially binds cGMP [55-57]. Accordingly, PKAs and PKGs are activated relatively specifically by cAMP and cGMP, respectively. However, either cyclic nucleotide can cross-activate the other enzyme *in vitro* and due to the lack of absolute specificity of either enzyme, PKAs and PKGs are capable of binding both nucleotides. It has even been postulated that it is possible that either cyclic nucleotide could cross-activate the other protein kinase in certain tissues [58]. Hence, it seems unusual to find a cyclic nucleotide binding protein that is capable of binding only cGMP but not cAMP as reported by Shalaby et al. [37], especially at such a high concentration as 20 µM. It appears more convincing to find that the trypanosomal protein is not capable of binding neither cAMP nor cGMP, as we are reporting here. Our results from docking experiments are consistent and strengthen these findings since they revealed binding energy differences that support the lack of both cAMP and cGMP binding to the TeqR-like protein. Moreover, no preference of the parasite protein to bind cGMP over cAMP can be presumed based on the predicted binding affinities disclosed by molecular modeling.

Besides the previous report of Shalaby et al. [37] showing that the *T. brucei* PKA is stimulated by cGMP, not any other role of cGMP has been proposed in these parasites. Likewise, the cGMP dependence of the *T. brucei* PKA R-like protein has not been confirmed by any other group. It is commonly known that guanylyl cyclases (GC) are the enzymes responsible for synthesizing cGMP from GTP, and cGMP-dependent phosphodiesterases (PDE) are the enzymes responsible for degrading cGMP to GMP. Strikingly, no GC activities have been identified in trypanosomes, nor is there any measurable cGMP-dependent PDE activity in trypanosomes [59]. These results also argue against a physiological role of cGMP in these parasites.

A plausible explanation for the inability of the recombinant trypanosomal R-like protein to bind cyclic nucleotides might be that the protein is not correctly folded. Although analysis by CD demonstrated the presence of secondary structures, it is known that the occurrence of secondary structures does not necessary mean that the protein possesses its functional tertiary conformation given that some secondary structures could be resistant to denaturation. However, the purified protein seemed to be properly folded because it

remained soluble up to concentrations higher than 40 mg/ml. It would not be thermodynamically viable for a partially unfolded protein to stay soluble at such a high concentration. In addition, formation of stable fragments by limited proteolysis with trypsin, chymotrypsin, V8 protease, papaín, and thermolysin, argues in favor of the recombinant protein possessing its native conformation. In all cases, the protein that originally had a molecular mass of ~56 kDa was primarily digested into two fragments; a large piece of 31–37 kDa and a small piece of 21–23 kDa. A partially unfolded protein would yield randomly generated proteolytic peptides following incubation with such different proteases.

Bioinformatics analysis of the N-terminal region of the TeqR-like protein established its homology with LRR proteins, a series of molecules that contain structures involved in protein-protein interactions. In all mammalian PKA R subunit isoforms, the D/D domain allows binding to the so-called A-kinase anchoring proteins (AKAPs), which function as key-regulator scaffolds of cAMP intracellular specificity synchronizing spatial and temporal aspects of cell signaling. Since no conventional AKAPs have been reported in trypanosomes, and the parasite R-like protein lacks the putative D/ D domain, it is appealing to postulate that under certain conditions this R-like protein might be capable of working in an analogous manner as an AKAP, tethering and directing trypanosome kinases that may be related to the PKA C subunits to their downstream substrate targets. As illustrated above (Figs. 1 and 2), the R-like protein possesses a site that contains a substrate recognition sequence (inhibitor site), and as some AKAPs from higher eukaryotes [60], it could be a preferred substrate for its associated protein kinase. In any case, its presence in every trypanosomatid genome elucidated to date clearly suggests that PKA R-like proteins might possess a significant function in trypanosome physiology.

5. Conclusions

The *T. equiperdum* ortholog of the PKA R subunit is a monomer, which is incapable of ligating cyclic nucleotides and contains an exposed hinge region that is vulnerable to hydrolysis by several proteolytic enzymes. Analyses using bioinformatics tools predicted that the COOH-terminal regions of the parasite R-like protein and mammalian PKA R subunits are related and possess similar 3D structures; however, the NH₂-terminal region of the trypanosomal protein is unrelated to that of its mammalian counterparts and shows homology with Leu-rich repeats-containing proteins. Predictions by molecular docking support the lack of cyclic nucleotide binding in the *T. equiperdum* R-like protein.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Abbreviations:

| РКА | cAMP-dependent protein kinase |
|--------------------------------------|--|
| R | PKA regulatory subunit |
| С | PKA catalytic subunit |
| Teq R-like (His) ₈ | poly-His tagged construct of the <i>Trypanosoma equiperdum</i> ortholog of the PKA R subunit |
| AC | adenylyl cyclase |
| PMSF | phenyl methyl sulfonyl fluoride |
| AEBSF | 4-(2-Aminoethyl) benzenesulfonyl fluoride hydrochloride |
| TLCK | N-a-tosyl-L-lysinyl-chloromethyl ketone hydrochloride |
| ТРСК | N-p-tosyl-L-phenylalaninyl chloromethyl ketone |
| IPTG | isopropyl-1-thio-β-D-galactopyranoside |
| muramyl dipeptide | acetylmuramyl-alanyl-isoglutamine |
| BSA | bovine serum albumin |
| 8-AEA | cAMP 8-(2-aminoethylamino) adenosine 3',5' -cyclic monophosphate |
| NHS | N-hydroxysuccinimide |
| BCIP | 5-bromo-4-chloro-3-indolyl phosphate |
| NBT | nitro blue tetrazolium |
| YT | Yeast extract and tryptone |
| LB | Luria-Bertani |
| SDS | sodium dodecyl sulfate |
| SDS-PAGE | SDS-polyacrylamide gel electrophoresis |
| MS/MS | tandem mass spectrometry |
| PCR | polymerase chain reaction |
| MES | 2-(N-morpholino)ethanesulfonic acid |
| DTT | dithiothreitol |
| HPLC | high-performance liquid chromatography |
| Mr | molecular weight |

| CD | circular dichroism |
|-------------------------|--|
| CNB | cyclic nucleotide binding |
| PBC | phosphate binding cassette |
| D/D | dimerization/docking |
| <i>f/f</i> ₀ | frictional coefficient |
| \overline{V} | partial specific volume |
| \overline{V}_{water} | partial specific volume of bound water |
| w _{max} | maximal hydration |
| R_s | Stokes radius |
| R _o | minimal radius |
| LRR | Leu-rich repeats |
| GST | glutathione S-transferase |
| GC | guanylyl cyclases |
| PDE | phosphodiesterase |
| AKAP | A-kinase anchoring protein |
| | |

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| 1 | atg M | tcto S | gaa E | aago K | ggaa G | icat T | cgt S | L | aac N | L | F | ctcç L | A A | A | gco C | aaa Q | aago K | gag E | ggto G | v V |
|-------------|-----------|------------------|-----------------------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|----------|
| 61 21 | aag K | caao Q | P | aaca N | acat T | F | ttç L | v | gag E | F | F | acga T | iaaa K | iaac K | P | jaad E | L | tcg S | gaaç E | v V |
| 121 41 | gag E | gaaa E | I | gact D | L L | igca S | iaga K | aact N | taca Y | I | ggg G | aaco N | gtg R | gta G | I | ttç L | acci A | L | ctcç L | D D |
| 181 61 | gta V | atta I | agc S | gago E | L | cat P | gct C | F | cgc R | F | L | aact N | gct C | caa S | naco N | aaa Q | aaao K | L | tata Y | nac N |
| 241 81 | acg T | gact D | L L | aato N | gagg E | D D | s S | v v | agaq R | ggt: G | aac N | gcca A | icaa T | I | D | gca R | I | gtc V | gacç D | v |
| 301 101 | ttc: F | aaat K | tca S | cato H | ccaa P | icaç T | A A | aaco N | gca A | L | gat D | ctca L | s S | aca H | N N | P | I | tca S | aact N | at Y |
| 361 121 | gct A | ggco G | cgt: R | agat R | ttat L | tgt L | L | L | T | caaa Q | aat N | aaca N | ago K | gca R | I | gco C | cgto R | gtg V | gaat E | tg L |
| 421 141 | gtg V | gaca D | act T | cgca R | I | pact D | F | gaad E | L | cgc R | tct S | cgca R | itta I | tcgc T | aac Q | aat Q | tgto C | gaa E | aaga K | nat N |
| 481 161 | acci T | atco I | gct: A | I | tggg W | agt E | s S | caaç Q | gca A | caa Q | gaa E | aago K | gagg E | agg E | aad E | gcç R | A | F | ggto G | jaa E |
| 541 181 | agt S | gtca V | acg T | tggo W | gtco V | caa P | icad T | aaa Q | acg T | s | gcg A | gact D | tga L | r T | A A | I | ggto G | ggc G | ggco G | gg R |
| 601 201 | aag K | cgao <u>R</u> | cgt: R | acta T | acaç T | v v | gaq R | ggto G | gago E | gga: G | I | gato D | ctg P | aga E | iago K | A | aaat K | tca S | tato Y | v V |
| 661 221 | gca A | ccat P | tat Y | ttco F | gaga E | iaga K | s S | gaag E | gaco D | gaa: E | acg T | gcad A | tca L | I I | tga L | agt K | L | ttg: L | acgt T | at Y |
| 721 241 | aac N | gta V | ctc L | ttc1 F | tcct S | F | tgç L | gact D | tca S | cgc R | gat D | ctta L | ntga M | t T | ytgo V | acgo A | ggtg G | gcc: A | atgt M | gg W |
| 781 261 | cgt R | gtg V | gag E | ttca F | aago K | agg Q | pato D | D D | tgc: C | I | atg M | gago E | acgo A | ggo G | aga Q | icaa T | acat T | tgt C | gaca D | iaa K |
| 84 281 | ctg L | tata Y | I | I | caag Q | D D | gta G | aaag K | gcag A | gat: D | I | I | aaag K | aag E | gad G | aga Q | aago K | gta V | tato Y | tc L |
| 901 301 | aag K | gtag V | gaa E | ggta G | acgg T | A A | ytag V | ggag G | gago E | L | gaa E | ctta L | ntgt M | ato Y | aga Q | icad T | ccaa P | T | gttg V | icc A |
| 961 321 | acg T | gtga V | aa⊡ K | gtti V | tgca C | T | P | gago E | L | I | gca A | tggg W | A | L | D | gco R | gaca D | acg T | tato Y | gt R |
| 1021 341 | cac H | L | gtg: V | atgo M | ggta G | s S | A | I | cgco R | cgc R | cgt R | gaga E | t T | ata Y | I | Q | F | L | acaa T | N |
| 1081 361 | att | ccg P | F | ctca L | agto S | gco G | ttç L | gaca D | N | tato Y | gag: E | aago K | tgc L | aac Q | ttç L | acgo A | gato D | gcc A | ctca L | igc S |
| 1141 381 | agc S | gaco D | gag E | F | gaad E | cgg P | gt: G | D | taca Y | I | I | cgct R | acg Y | gag G | agg E | gagg E | ggco G | gaa E | tggo W | tg L |
| 1201 401 | tac: Y | atca I | I | ttgg L | gaag E | gtt G | s | v v | D | ytg V | gtt V | gggo G | ggg R | D D | jaco D | D | ggaa G | N | gaaa E | ag K |
| 1261 421 | cat H | gtt: V | tgg W | gaat E | ttcç F | gga G | iago K | gggg G | D D | cat H | gtg V | ggtg G | gago E | tgç L | jaat E | tco F | L | aac: N | aato N | ac H |
| 1321 441 | gcc: A | aato N | gta V | gcaç A | gatç D | V | v V | gcaa A | aaaa K | acg T | H | ytto V | ytca V | t T | A | k K | L | aac N | cgta R | ngg R |
| 1381 461 | cac H | F | gaa E | atgi M | tgco C | ttç L | gao G | P | ytc: V | I | gat D | gtad V | tga L | iaac K | :gaa R | t T | s S | cag Q | cago Q | ca P |
| 1441 481 | aac N | tato Y | gag [.] E | tact Y | taco Y | agt Q | cga S | aaao K | L | aaa K | T | acti T | taa L | nggg R | A | gago E | ggga G | agg R | aagt K | ag * |

Fig. 1. Gene and amino acid sequence of the *T. equiperdum* R-like protein.

The single nucleotide difference between the *T. equiperdum* and the *T. brucei* PKA R-like protein genes is highlighted in black (nucleotide 969). Highlighted in grey is the sequence information gathered by MS/MS analysis of the tryptic peptides from the recombinant Teq R-like protein (Table SI). A potential PKA consensus phosphorylation sequence that resembles the inhibitor or pseudosubstrate site is double underlined. The putative CNB domains and PBC sites are boxed and underlined, respectively.

| Δ | | 0 | RIα |
|---|---|-----|--------------------|
| ~ | | 0 | RIß |
| | | 0 | Rlla |
| | | 0 | RIIß |
| | MSEKGTSLNLFLAACQKEGVKQPNTFLVEFFTKKPELSEVEEIDLSKNYIGNRGILALLD | 60 | Teq R-like |
| | | 3.8 | RIα |
| | Wispercesepercesever workstoo | 3.8 | RIG |
| | | 37 | Blla |
| | | 3.6 | RIIG |
| | UT OPI, DO PD PI, NO ON A TI, NO POUD ANA TI DI UNA SUPERA SUN DI ONV | 120 | Teg P_like |
| | | 100 | redivince |
| | CTADDEDDWAPLDEV PEDLEKEPAKOTONLOKBCTETOSDEDETSDED | 8.6 | Pla |
| | CTCP | 86 | DIR |
| | TDLDFADDASVLDATDOSLGHDDFDGDDDVADAGOSPSPEDD | 85 | Dlla |
| | TRIOOFNERKGINERGERGRUNDIGALAGGERSKGUNFAFFFMOSDSFDGEFFFAA | 9.4 | PIIR |
| | AGRRLLLLTQNNKRICRVELVDTRIDFELRSRITQQCEKNTIAIWESQAQEKEEERAFGE | 180 | Teq R-like |
| | 1 | | |
| | PNPVVKGRRRRGAISAEVYTEEDAASYVRKVIPKDYKTMAALAKAI | 132 | RIα |
| | PNPVVKARRRRGGVSAEVYTEEDAVSYVRKVIPKDYKTMTALAKAI | 132 | RIß |
| | LEVPVPSRFNRRVSVCAETYNPDEEEEDTDPRVIHPKTDEQRCRLQEAC | 134 | RIIα |
| | PADAGAFNAFVINRFTRRASVCAEAYNPDEEEDDAESRIIHFKTDDQRNRLQEAC | 149 | RIIß |
| | SVTWVFTQTSADLTAIGGGRKRRTTVRGEGIDPEKAKSYVAPYFEKSEDETALILKLL | 238 | Teq R-like |
| | | 107 | DIa |
| | EKNYLESHEDDNEKSDIFDAMESYSFIAGEIVIQQDE-GDNFIVIDQGEIDVIVN | 107 | DIO |
| | SKNVBFAREDDNEKSDIFDAMFFVIRIAGEIVIJQGAD-GDNEVVDQGEVDVIVN | 102 | RIP |
| | RDILBERNEDQEQESQVEDAMFERIVENDEUVIDGED-GENEVIERGIIDIUVIRAQ | 193 | DHQ |
| | | 200 | Tog D like |
| | IIN BEST BUSKDERIVAGARWERVERKUDDEINERGUITEDKEITIQUGKADIIKE | 231 | req R-like |
| | NEWATSVGEGGSFGELALIYGTPRAATVKAKT-NVKLWGIDRDSYRRILMGSTLRKRKMY | 246 | RIα |
| | GEWVTNISEGGSFGELALIYGTPRAATVKAKT-DLKLWGIDRDSYRRILMGSTLRKRKMY | 246 | RIß |
| | TRSVGQYDNRGSFGELALMYNTPRAATIVATS-EGSLWGLDRVTFRRIIVKNNAKKRKMF | 252 | RIIα |
| | GRCVGNYDNRGSFGELALMYNTPRAATITATS-PGALWGLDRVTFRRIIVKNNAKKRKMY | 267 | RIIß |
| | GQKVYLKVEGTAVGELELMYQTPTVATVKVCTPELIAWALDRDTYRHLVMGSAIRRRETY | 354 | Teg R-like |
| | | | |
| | EEFLSKVSILESLDKWERLTVADALEPVQFEDGQKIVVQGEPGDEFFIILEGSAAVLQRR | 306 | RIa |
| | EEFLSKVSILESLEKWERLTVADALEPVQFEDGEKIVVQGEPGDDFYIITEGTASVLQRR | 306 | RIß |
| | ESFIESVPLLKSLEVSERMKIVDVIGEKIYKDGERIITQGEKADSFYIIESGEVSILIRS | 312 | Rlla |
| | ESFIESLPFLKSLEFSERLKVVDVIGTKVYNDGEQIIAQGDSADSFFIVESGEVKITMKR | 327 | RIIB |
| | IQFLTNIPFLSGLDNYEKLQLADALSSDEFEPGDYIIRYGEEGEWLYIILEGSVDVVGRD | 414 | Teq R-like |
| | | | |
| | SENEEFVEVGRLGPSDYFGEIALLMNRPRAATVVARGPLKCVKLDRPRFERVLG | 360 | RIG |
| | SPNEEYVEVGRLGPSDYFGEIALLLNRPRAATVVARGPLKCVKLDRPRFERVLG | 360 | RIB |
| | RTKSNKDGGNQEVEIARCHKGQYFGELALVTNKPRAASAYAVGDVKCLVMDVQAFERLLG | 372 | RIIQ |
| | KGKSEVEE-NGAVEIARCSRGQYFGELALVTNKPRAASAHAIGTVKCLAMDVQAFERLLG | 386 | RIB |
| | DDGNEKHVWEFGKGDHVGELEFLNNHANVADVVAKTHVVTAKLNRRHFEMCLG | 967 | leq R-like |
| | | | |
| | PCSDILKRNIQQYNSFVSLSV 381 RIG | | |
| | PCSEILKRNIQRYNSFISLTV 381 RIB | | |
| | PCMDIMERNISHYEEQLVEMPGSSVDLGNLGQ 404 RHa | | |
| | PCMEIMKRNIATYEEQLVALFGTNMDIVEPTA 418 RHB | | |
| | EVIDVLERTSQQPNYEYYQSKLKTTLRAEGRK 499 TeqR-like | | |
| | | | |
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| D | | | |
| в | | P | AR subunits |
| | Inhibitor site | | |
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| | | _ | Variation and |
| | CNBA CNBB | Te | qR-like protein |
| | 4 | | |

Potential phosphorylation site

Fig. 2. Comparison of the primary structure and domain organization of the *T. equiperdum* R-like protein and the four human PKA R subunit isoforms. A, Alignment of amino acid sequences of the R-like protein from *T. equiperdum* and the

 R^{I}_{α} , R^{I}_{β} , R^{II}_{α} , and R^{II}_{β} subunits from human [Accesion No. P10644.1, P31321.4, P13861.2 and P31323.3, respectively using the NCBI Protein database (https:// www.ncbi.nlm.nih.gov/protein/)]. Identical residues are shown with an asterisk (*), conservative substitutions are labeled with a colon punctuation mark (:), and semiconservative substitutions are labeled with a period (.). **B**, The standard domain structure of mammalian PKA R subunits is contrasted to the domain organization of the Teq R-like protein. CNBA and CNBB = cyclic nucleotide binding domains A and B, respectively; D/D = dimerization and docking domain.



Fig. 3. Analysis by circular dichroism spectroscopy of the recombinant TeqR-like(His)₈ protein. Comparison of CD measurements obtained for the bovine PKA $R^{I}\alpha$ subunit (A), human PKA $R^{I}\beta$ subunit (B), and *T. equiperdum* R-like protein (C).



Fig. 4. Size of the TeqR-like protein.

A, Separation by molecular exclusion column chromatography of the TeqR-like(His)₈ protein (Teq R-like). Parallel experiments were performed with the $R^{I}\beta_{2}C_{2}$ holoenzyme (Holo $R^{I}\beta$), the $R^{I}\beta_{2}$ subunit ($R^{I}\beta$), and the C subunit (C). **B**, Separation by gel filtration chromatography of TeqR-like(His)₈ in the presence of various molecular weight markers: thyroglobulin (Thyr), immunoglobulin G (IgG), ovalbumin (Ova), myoglobin (Myo) and vitamin B₁₂ (VitB₁₂). The size of the unknown protein was estimated by plotting the *K*av value of each standard *versus* the logarithm of its molecular weight (Mr).

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Fig. 5. Limited proteolysis analyses of the *T. equiperdum* R-like protein.

Assays were carried out at 4 °C and 37 °C, using various amounts of trypsin (**A**), chymotrypsin (**B**), the V8 protease from *S. aureus* (**C**), papaín (**D**), and thermolysin (**E**). Arrows indicate the molecular masses of the undigested TeqR-like(His)₈ protein and of the major proteolytic fragments obtained following the various enzymatic digestions.



Fig. 6. The *T. equiperdum* R-like protein is not capable of binding cyclic nucleotides. A, Cyclic nucleotide binding was determined by equilibrium dialysis. Shown is the [³H] cAMP binding of the PKA bovine R^I α subunit (\blacksquare), the human R^I β subunit (●), and the *T. equiperdum* R-like protein (▲). Also shown is the [³H] cGMP binding of the parasite R-like protein (Inset). B, cAMP binding was also analyzed by chromatography throughout a cAMP-Sepharose affinity column. Gel lanes contained the following samples: 1, original protein; 2, flow through fraction; 3, bound protein. The bovine R^I α subunit and human R^I β subunit were included as controls. Arrows indicate the molecular masses of the three proteins used in the experiment.



Fig. 7. Structural model of the *T. equiperdum* R-like protein.

A, The Phyre2 bioinformatics tool [38] was employed to predict the structure of the *T*. *equiperdum* R-like protein. Ribbon diagram structures of the modeled N-terminal (residues 1–201) and C-terminal (residues 202e499) regions are shown in light grey and dark grey, respectively. **B**, The model retrieved for the trypanosome protein is compared to those retrieved for PKA human R^I β (PDB ID: 4DIN [22]) and rat R^{II} β (PDB ID: 3TNQ [61]). N-terminal and C-terminal regions are shown in light grey and dark grey, respectively.





Fig. 8. View of the predicted structures obtained by molecular docking for the cyclic nucleotide binding pockets of the *T. equiperdum* R-like protein.

Shown is a comparison of the models attained for both cyclic nucleotide binding sites of the *T. equiperdum* R-like protein and the bovine PKA R^Ia subunit. CNB A and CNB B= cyclic nucleotide binding domains A and B, respectively. Highlighted are the functionally important residues involved in cAMP binding in R^Ia, and the corresponding residues in the parasite protein based on amino acid sequence alignment. Docking was performed using both cAMP (**A**) and cGMP (**B**).