Structural insights into *kinetoplastid* coronin oligomerization domain and F-actin interaction

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**ABSTRACT**

The two-domain actin associated protein coronin interacts with filamentous (F-) actin, facilitating diverse biological processes including cell proliferation, motility, phagocytosis, host-parasite interaction and cargo binding. The conserved N-terminal β-propeller domain is involved in protein: protein interactions, while the C-terminal coiled-coil domain mediates oligomerization, transducing conformational changes. The *L. donovani* coronin coiled-coil (LdCoroCC) domain exhibited a novel topology and oligomer association with an inherent asymmetry, caused primarily by three α residues of successive heptads. In the *T. brucei* homolog (TbrCoro), two of these α residues are different (Val 493 & 507 replacing LdCoroCC Ile 486 and Met 500 respectively). The elucidated structure possesses a similar topology and assembly while comparative structural analysis shows that the *T. brucei* coronin coiled-coil domain (TbrCoroCC) too possesses the asymmetry though its magnitude is smaller. Analysis identifies that the asymmetric state is stabilized via cyclic salt bridges formed by Arg 497 and Gln 504. Co-localization studies (LdCoro, TbrCoro and corresponding mutant coiled coil constructs) with actin show that there are subtle differences in their binding patterns, with the double mutant V493I/V507M showing maximal effect. None of the constructs have an effect on F-actin length. Taken together with LdCoroCC, we therefore conclude that the inherent asymmetric structures are essential for kinetoplastids, and are of interest in understanding and exploiting actin dynamics.

1. Introduction

Leishmaniasis and trypanosomiasis are parasitic vector borne diseases affecting millions of people in tropical and sub-tropical regions, caused by the kinetoplastid protozoans *Leishmania* and *Trypanosoma species* respectively. A characteristic feature of these parasites is the presence of flagella bound to the membrane network, consisting of actin filaments that are vital for the organism. Depletion of this cyto-skeletal protein in *T. brucei* resulted in subsequent death (Reisler, 1993). Actin exists in two forms: a globular monomeric form known (G-actin) and as filamentous polymer F-actin. The actin modulation machinery entails filament assembly induced by ATP binding to globular actin (G-actin) forming F-actin, while the release of inorganic phosphate upon hydrolysis results in its dis-assembly, forming ADP-G-actin. These complex arrays of dynamic changes are regulated by actin associated proteins including coronin, coflin, profilin and the Arp2/3 complex (De Hostos et al., 1993; Ayscough et al., 1998). The widely expressed eukaryotic conserved, F-actin binding protein coronin, first identified in *D. discoidium* where lack of coronin led to cell migration defects, is a key regulator of actin assembly-dis-assembly dynamics that plays important roles in cell motility, phagocytosis, cytokinesis, immune regulation (Maniak et al., 1995; Hacker et al., 1997; Nagasaki et al., 2001; Bharathi et al., 2004; Morgan and Fernandez, 2008; Xavier et al., 2008). The association of coronin with the Arp2/3 protein complex mediates actin filament branch generation, while the ADP-bound actin filaments are dismantled by coronin and coflin (Humphries et al., 2002). In *Leishmania*, *S. cerevisiae* and *D. melanogaster*, coronin has been shown to bind with microtubules and cross-link to actin filaments (Goode et al., 1999; Yan et al., 2005) in *Toxoplasma* it facilitates host cell invasion (Shina et al., 2011; Xavier et al., 2012; Tchang et al., 2013) while in *Plasmodium* sporozoite mutant coronin exhibits defect in motility (Bane et al., 2016) in *Leishmania*, the actin filaments are distinct from most organisms, being primarily...
multi-branched and comparatively short, and overexpression of coronin in *L. donovani* increases the occurrence of actin filaments (Nayak et al., 2005; Srivastava et al., 2015). The kinetoplastid coronins, are short (Type I, 400–600 residues) and consist of an N-terminal WD40 repeat domain (~350 residues), and a C-terminal coiled-coil domain (50–80 residue), connected by a variable linker that has unique and conserved regions (de Hostos, 1999; Mc Ardle and Hofmann, 2008; Appleton et al., 2006; Eckert et al., 2011; Nayak et al., 2016). The Coiled coil (CC) domain, made of two or more helices forming a supercoil by mutually burying their hydrophobic regions, are ubiquitous structural motifs, constituting 3–5% of the whole genome, that are primarily involved in homo- and hetero-oligomerization (Wolf et al., 1997; Liu et al., 2006; Rackham et al., 2010). CCs are associated with specific sequence patterns corresponding to the supercoil handedness, with the left-handed supercoil having a heptad repeat abcdefg where hydrophobic residues a, d associate in a knobs into holes pattern, with electrostatic interactions between e, g providing additional stability (O'Shea et al., 1993; Krylov et al., 1994; Monera et al., 1994; Zeng et al., 1997; Kohn et al., 1998; McClain et al., 2001). The CC is dynamic, with changes as subtle as mutation of a single residue leading to drastic consequences, including changes in the oligomer assembly, topology of the helical bundle (Harbury et al., 1993, 1998; Stetefeld et al., 2000; Spoerl et al., 2002; Kammerer et al., 2005; Deng et al., 2008; Kumar et al., 2018).

The *L. donovani* coronin coiled-coil domain (LdCoroCC), solved earlier in the laboratory showed that the structure, unlike the mammalian homologues (parallel trimer), assembles with a novel topology and oligomeric state (anti-parallel tetramer), with an inherent asymmetry, arising from steric clashes between a residues (Ile 486, Leu493 and Met 500) of three successive heptads (Nayak et al., 2016; Karade et al., 2020). Sequence analysis of the homologous *T. brucei* coronin coiled-coil domain (TbCoroCC) indicates that two of the a residues are different, from *L. donovani* with Val 493 and 507 replacing Ile 486 and Met 500 respectively (Fig. 1A). To understand and to investigate the role of these changes, the structures of *T. brucei* coronin coiled coil domain (TbCoroCC) constructs were elucidated and the effects of the constructs along with LdCoro mutants (I486A and I486A-L493A (I-L) M500V) were functionally characterized in vivo.

### 2. Methods

#### 2.1. Bacterial strain and growth condition

The *E. coli* strains DH5α and RPL were grown in Lysogeny Broth (LB) medium at 37°C and 25°C.

#### 2.2. Sequence analysis

The coronin gene sequences of *L. donovani* and *T. brucei* homologues were retrieved from GeneDB (http://www.genedb.org) and Uniprot (www.uniprot.org). Sequence alignments of coronin homologues were done by Clustal omega (http://www.ebi.ac.uk/Tools/msa/clustalo/). Coiled coil domain was predicted with MARCOIL web server (http://too.kit.tuebingen.mpg.de/marcoil), and the alignment visualized using ESPript (https://espript.ibcp.fr/) (Robert and Gouet, 2014).

#### 2.3. Cloning, point mutation, protein expression and purification

The *T. brucei* Coronin Coiled-coil domain was amplified using forward and reverse primers GAATTCCTGCGAGTGTTAGCCTTGCGCTCG and CTCGAGGGCAAGGGGCTTTATTTG CGAT containing BamHI and EcoRI restriction sites (Bold) designed manually from *T. brucei* strain (927/4 GUTat10.1) and *L. donovani* strain (BPK282A1) and the PCR product ligated with T/A vector pTZ57 R/T (Ins TA clone™ PCR cloning kit, Fermentas International Inc.) TA clone digested and sub clone into pET28 (a) (Novagen) expression vector. For study the digested product ligates and expressed in pLEXSY integrative vector system in *Leishmania*. We have created single point mutation at V507M using appropriate forward and reverse primers (CTATTGGGGCAGCAA- CAGGCGGAGATCCAG and TGAAGGATCTCTCAGGAAACCTTGGA TCT, for double mutant for the V493I (ATGG and CTTTATCTTTGCGATTGTCTCCAT-) and V507M:GACCTGCAGAA respectively using single mutant as template. All the point mutations were functionally characterized in vivo.

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**Fig. 1. Characterization of TbrCoroCC** (A) Pairwise sequence alignment of *L. donovani* and *T. brucei* coronin coiled-coil domains, with a, d residues highlighted in yellow and red boxes respectively. Residues implicated in the structural asymmetry in LdCoroCC are colored yellow (I486, L493 and M500) (B) Size exclusion chromatography profile of TbrCoroCC shows the protein construct elutes as a tetramer (~32 kDa) the inset shows molecular weights of standards, while the SDS-PAGE showing the protomer of ~8 kDa (C) Experimental scattering curve of TbrCoroCC with the Guinier Region and linear regression (solid line) for Rg evaluation shown in inset (D) Pair distance distribution function p(r) at 10 mg/ml concentration reveals longest dimension (Dmax) of ~11.5 nm with the unstructured residues from the expression vector possibly accounting for the additional length, in insetKratyplot for the protein TbrCoroCC shows broadness of curve of a protein indicates rod like protein. (E) Different orientations of the low-resolution structure of the protein (envelope), generated by using the in-solution scattering of X-ray intensity data. The TbrCoroCC structure model was fitted well inside the envelope.
mutation created by site directed mutagenesis method and confirmed by sequencing.

The genes, cloned in pET28a containing N- and C- terminal His tags were transformed and over-expressed in competent E.coli strain RPL cells and, allowed to grow at 37 °C until OD_600 ~ 0.6 prior to induction with 1 mM isopropyl β-D-thiogalactopyranoside (IPTG), harvested after 6–8h at 25 °C by centrifugation at 6500 rpm for 15 min. Cells were resuspended in the lysis buffer (50 mM TRIS-Cl pH = 7.5, 200 mM NaCl) and subsequently lysed by sonication with a 10s on/10s off pulse for 30 min. On centrifugation at 10,000 rpm for 45 min the cell debris was removed and the supernatant was loaded on an IMAC column (Qiagen, Germany) pre-equilibrated in the buffer, (50 mM TRIS-Cl pH = 7.5, 200 mM NaCl) washed with buffers containing 10 mM and 30 mM imidazole respectively before eluting with 250 mM imidazole in the same buffer. The eluted protein was dialyzed overnight to remove imidazole against a buffer containing 20 mM TRIS-Cl pH 7.5, 100 mM NaCl, 1 mM EDTA and 3 mM β-me, concentrated using 10 kDa cutoff centricon (Amicon).

2.4. Gel filtration analysis

Size exclusion chromatography was carried out using S75 10/300 prepacked columns connected to AKTA pure (GE Healthcare, CytiVA USA). The purified concentrated protein was loaded on pre equilibrated Column. The protein elution volume of 11.6 mL was compared with molecular weight standards (GE catalogue 28403841), the molecular weight and oligomer state estimated using the following standards from GE Healthcare: Ovalbumin (45 kDa), carbonic anhydrase (29 kDa), ribonuclease A (13.7 kDa) and Aprotinin (6.5 kDa). The eluted protein was collected and concentrated for prior to use.

2.5. Western blotting

The proteins from whole Leishmanial cell lysates were separated by SDS-PAGE and blotted to nitrocellulose membrane. Protein bands were detected by incubating with anti His polyclonal primary mouse antibody (Santa Cruz) for 3 h at room temperature subsequently incubation with horseradish peroxidase (HRP)-conjugated anti-mouse IgG, for 2 h at room temperature. The HRP substrate luminol and H_2O_2 mixed with 1:1 ratio was spread over the blot which resulted in chemiluminescence and detected in Chemidoc equipment (GE Biosciences).

2.6. Chemical cross linking

Cross linking reactions were set with a 30 mM HEPES (pH 7.5) and 100 mM NaCl with glutaraldehyde concentrations (0.025 to 0.5%) at protein sample was taken. The spectra measured [spectropolarimeter (Applied Photophysics) (SAIF facility in CSIR-CDRI, Indore, INDIA (TbrCoroCC) or on beamline XR2D, Elettra Sincrotrone, Trieste, Italy (V507M and double mutant)].

2.7. Circular dichroism spectroscopy

CD spectroscopy assays performed using buffer 20 mM TRIS pH 7.5, 100 mM NaF with protein concentration 5 µM on a J810 ChirascanTM CD spectropolarimeter (Applied Photophysics) (SAIF facility in CSIR-CDRI, Lucknow) calibrated with ammonium (+/-)10- camphorsulfonate. The average of four spectra (190–250 nm, scan speed 10 nm/min) of each protein sample was taken. The spectra measured [O]_222 as a function of temperature under the same conditions. Spectra were collected in a temperature-controlled quartz cuvette, the successively 10 °C interval in 20–90 °C scanning window at a ramp rate of 2 °C/min.

2.8. Crystallization and data collection

Preliminary crystallization trials were initiated using commercial screens (Crystal Screens 1, 2; PEG ion screen) and around the homologous LdCoroCC conditions with a protein concentration of 10 mg/mL mixed with equal volume of the reservoir solution, equilibrated against 0.5 ml reservoir solution. An initial hit was observed in a Crystal Screen 1 (condition 0.1M HEPES 7.5 and 4.3 M NaCl) in ~10–14 days and the condition optimized, leading to the growth of diffracting crystals from a hanging drop vapor diffusion where 2 µl of the protein (8 mg/mL) was mixed with equal volume of the precipitate containing 0.1M HEPES pH 7.5 and 3 M NaCl, equilibrated against a 0.5 ml reservoir solution. Crystals were flash frozen with 20% glycerol as a cryo-protectant and data collected (to 2.06–3.02 Å) on the PX-BL21 beamline at INDES-2, Raja Ramanna Centre for Advances Technologies, Indore, INDIA (TbrCoroCC) or on beamline XR2D, Elettra Sincrotrone, Trieste, Italy (V507M and double mutant).

2.9. Data processing, structure determination and refinement

The X-ray diffraction data of TbCoroCC were processed using the HKL2000 suite of programs (Otwinowski and Minor 1997; Minor et al., 2006). Indexing the data suggested that the crystal belongs to body centered tetragonal space group 1 4 2 2, with unit cell dimensions a = b = 93.3, c = 82.9 Å and the data integrated and scaled to 2.06 Å. Solvent content analyses indicate the asymmetric unit contains two protomers with a solvent content of 50% and Vm of 2.11. The qualities of data were assessed by phenix.Xtriage (Zwart et al., 2005), which confirmed the absence of twinning or diffraction anisotropy. The V493I and double mutant structures were processed using iMosfil & Aimless, from the CCP4 suite of programs (Evans, P. 2014). Data reduction statistics are summarized in Table 1.

The crystal structure was solved by molecular replacement methods using the Phaser program from the phexin suite (Adams et al., 2010) with a canonical dimer of the L. donovani coronin coiled-coil domain structure as the template (PDB ID 5CX2, chains A&B), which gave a unique solution (TFZ 12.5; LLG 196). Refinement of the structure was carried out using the phenix.refine (Afonine et al., 2012) program from the phexin suite. The refinement strategy was an initial poly alanine rigid body refinement followed by restrained refinements. Electron density maps were computed after each round of refinement, and side chains progressively fitted in COOT (Emsley et al., 2010) and towards the end of the refinement cycles, solvent and other organic groups from the crystallization conditions were added. The crystallographic R-factor and R-free, which were 31.1% and 34.5% after the first rigid body refinement, subsequently converged to 22.6% and 25.7% respectively. The final model contains 99 amino acid residues (52 in chain A and 47 in chain B), 68 solvent molecules.

2.10. Small-angle X-ray scattering (SAXS)

SAXS measurements were done at in-house SAXS instrument SAXSpace (Anton Paar) at Central Drug Research Institute, Lucknow, and samples with 5–15 mg/ml concentration were applied to the system in 30 mM Tris-Cl pH 7.5 and 100 mM NaCl buffer. The protein and buffer frames were selected for processing using Prisms (Petrovokh et al., 2012). The buffer subtracted protein frames were scaled and averaged. DAMMIF used to generate ab initio models were averaged using DAMAVER (Volkov et al., 2003) and the most typical model was selected. The analogy of models theoretical scattering and experimental scattering superposition was performed with Gasbor, (Svergun et al., 2001) the graph plotted with the help of GraphPad prism 7.0. The processing software was part of the ATSAS package version 3.

2.11. Genetic manipulations and parasite culture

For in vivo study, the amplified gene products were cloned in pLEXY-hyg2.Integrative vector system (Jena Biscience). The resulting pLEXY constructs were linearized by Swal or Smiel restriction enzyme and approximately 10 µg of purified DNA products were transfected into...
dehyde and these wash with 5% (w/v) glycine in PBS was given to quench extra formaldehyde. The Penicillin-Streptomycin preparation was added. For each transfection, 10 μl serum (Gibco) along with 40 µl DMEM (Gibco) supplemented with 10% heat inactivated fetal bovine serum was added. Blocking was done using 1% BSA in PBS with 0.02% sodium azide for 1 h at room temperature. Primary antibody treatment with anti-actin antibodies (rabbit) (1:500) and anti-His antibodies (mouse) (1:200) was given in same blocking buffer at 4 °C, overnight.

Cover slips were washed for about 7–8 times with blocking buffer which was followed by secondary antibody treatment with Alexa Fluor 488 (anti-rabbit) and Alexa Fluor 546 (anti-mice) for 4 h at 4 °C. Cover-slips were then washed again for 7–8 times with blocking buffer and then mounted onto glass slides in mounting media (Calbiochem, Germany) and were stored at 4 °C in the dark until scanning. Confocal images were collected on Leica SP8 confocal microscope using 63X oil plan apochromate lens (1.4 NA) at 3X digital zoom. Images were processed and arranged for presentation in Adobe Photoshop (Creative Suit 6).

2.13. Statistical analysis

The results were presented as the pooled data of three independent experiments expressed as mean ± standard deviation (SD). Analysis was done using unpaired t-test with equal SD using PRISM software (Version 6.2).

3. Results

3.1. TbrCoroCC structure determination and analysis

The 47 residue native T. brucei coronin coiled-coil domain (Residues 477–523 of T. brucei coronin, UNIPROT ID Q57W63), and the single and double mutant constructs V507M and V493I–V507M were over-expressed and purified to homogeneity as described in Experimental procedures. The size exclusion profile shows the elution volume corresponds to a tetramer (Fig. 1A and B, Supplementary Fig. S1), further validated by solution X-ray scattering studies (Fig. 1C–E) and glutaraldehyde crosslinking studies (Supplementary Fig. S2). The crystal structure was determined by molecular replacement using a canonical dimer of LdCoroCC (AB dimer of PDB: 5CX2) as a template. The asymmetric unit contains an anti-parallel dimer, with the tetramer generated by the crystallographic two -fold axis. The biological assembly has dimensions of ~70 × 25.7 × 20.9 Å3 (Fig. 2 A, B). Structures of the single (V507M) and the double mutant (V493I-V507M) constructs were determined using TbrCoroCC as the template and the mutations validated by omit maps (Fig. 2C). Data collection and refinement statistics for the apo and mutant constructs are summarized in Table 1.

The individual protomers A, B are essentially identical in the three TbrCoroCC structures with root mean square displacement (RMSD) values ~0.5 Å. The protomers associate as a coiled coil between residues 479–514 are stabilized by ad interactions in a knobs-into-holes packing. Significantly, in TbrCoroCC 3 d residues are polar (Gln 489, Lys 503 and Thr 510), and these polar groups provide additional stability: side chains of Gln A 489:Thr B 510 (and B489:A510) form a hydrogen bond (2.8 Å) while Lys 503 forms a salt bridge with Glu 492 of the opposite chain (2.78 Å) (Fig. 2A). The corresponding residues in LdCoroCC are hydrophobic except Lys 496 (equivalent to Lys 503), which forms an ionic interaction with Glu 485 of the partner chain. The salt bridges forming residue Arg 497 and Glu 504 strengthen the stability of structure (Fig. 2B).

Oligomerization analysis using PISA (Krissinel, E., and Henrick, K. 2007) confirms the biological assembly to be a tetramer, consistent with solution experiments. In left-handed coiled-coils with three or more helices, an additional residue, typically e or g is furthermore buried in the interface, referred to as ada/adg packing: core packing analysis of the TbrCoroCC tetramer using the helical analysis programs (Dunin-Horkawicz, S. and Lupashin, A. N. 2016; Pratap et al., 2013) indicates that TbrCoroCC adopts the ada core packing, similar to LdCoroCC (Fig. 3A and B C D). The canonical dimer A/B and C/D show identical knobs into holes interactions (Fig. 3A) while there is difference at non-canonical dimeric interaction between A/D and B/C (Fig. 3B and C); while the B/C dimeric interface extends to the length of the coiled-coil region, helices A and D associate only at either termini (Fig. 3B). As seen in Fig. 3C, the A/D and B/C interfaces are not identical; Residue Ala 518 interacts with residue

| Table 1 Data collection, processing and structural refinement statistics (Values in parentheses are for highest resolution shell). |
|-----------------|-----------------|-----------------|
|                  | TbrCoroCC       | V493I–V507M     | V507M        |
| **Unit cell**    |                 |                 |               |
| a [Å]            | 82.9            | 80.7            | 82.9          |
| b [Å]            | 90              | 90              | 90            |
| c [Å]            | 93.3            | 95.02           | 90.65         |
| α [°]            | 90              | 90              | 90            |
| β [°]            | 90              | 90              | 90            |
| γ [°]            | 90              | 90              | 90            |
| **Completeness (%)** | 99.14 (98.95) | 95.02 (99.12) | 90.65 (95.82) |
| **Resolution range** | 2.13 (2.1) | 2.41 (2.4) | 2.41 (2.4) |
| **Unique reflections** | 13712 (1308) | 6752 (681) | 3477 (345) |
| **Multiplicity** | 8.9 (8.4) | 14 (13.9) | 2.0 (2.0) |
| **Rmerge** | 0.11 (0.61) | 0.053 (0.374) | 0.040 (0.19) |
| **Rfree** | 0.098 (0.451) | 0.013 (0.101) | 0.017 (0.027) |
| **Reflections used in refinement** | 556 (64) | 297 (27) | 140 (15) |
| **Rwork** | 0.220 (0.334) | 0.236 (0.265) | 0.223 (0.311) |
| **Rfree** | 0.240 (0.419) | 0.255 (0.434) | 0.264 (0.409) |
| **CC1/2 (R-work)** | 0.971 (0.716) | 0.972 (0.875) | 0.975 (0.564) |
| **CC1/2 (R-free)** | 0.971 (0.716) | 0.972 (0.875) | 0.975 (0.564) |
| **Residuals** | 0.51 (0.419) | 0.515 (0.434) | 0.518 (0.409) |
| **CC(work)** | 0.971 (0.716) | 0.972 (0.875) | 0.975 (0.564) |
| **CC(free)** | 0.971 (0.716) | 0.972 (0.875) | 0.975 (0.564) |
| **Number of non-hydrogen atoms** | 807 | 752 | 736 |
| **macromolecules** | 763 | 748 | 731 |
| **solvent** | 94 | 4 | 5 |
| **Protein residuals** | 99 | 98 | 95 |
| **RMS(bonds)** | 0.006 | 0.008 | 0.007 |
| **RMS(angles)** | 0.77 | 0.97 | 1.07 |
| **Ramachandran favored (%)** | 100 | 98.94 | 100 |
| **Ramachandran allowed (%)** | 0 | 1.06 | 1.09 |
| | 0 | 0 | 0 |
| **Clashscore** | 1.96 | 2.66 | 6.08 |
| **Average B-factor** | 67.28 | 67.77 | 66.38 |
| **macromolecules** | 67.33 | 67.83 | 66.42 |
| **solvent** | 68.01 | 56.82 | 59.80 |

Leishmania promastigotes by electroporation using Gene pulser (Bio-Rad) as described earlier (Nayak et al., 2005). The transfectants were selected against 10 μg/ml hygromycin and maintained at 26 °C in high glucose DMEM (Gibco) supplemented with 10% heat inactivated fetal bovine serum (Gibco) along with 40 µg/ml gentamycin and 100 µg/ml Penicillin-Streptomycin preparation.

2.12. Immunofluorescence microscopy

For immunofluorescence microscopy, log-phase Leishmania promastigotes were adhered on poly-L-lysine coated coverslips for 2 min and then fixed with 4% (w/v) paraformaldehyde in PBS for 15 min. A single wash with 5% (w/v) glycine in PBS was given to quench extra formaldehyde and these fixed cells were permeabilized with 1% Triton X-100 in PBS for 20 min. Blocking was done using 1% BSA in PBS with 0.02% sodium azide for 1 h at room temperature. Primary antibody treatment with anti-actin antibodies (rabbit) (1:500) and anti-His antibodies (mouse) (1:200) was given in same blocking buffer at 4 °C, overnight. Cover slips were washed for about 7–8 times with blocking buffer which was followed by secondary antibody treatment with Alexa Fluor 488 (anti-rabbit) and Alexa Fluor 546 (anti-mice) for 4 h at 4 °C. Cover-slips were then washed again for 7–8 times with blocking buffer and then mounted on glass slides in mounting media (Calbiochem, Germany) and were stored at 4 °C in the dark until scanning. Confocal images were collected on Leica SP8 confocal microscope using 63X oil plan apochromate lens (1.4 NA) at 3X digital zoom. Images were processed and arranged for presentation in Adobe Photoshop (Creative Suit 6).
Ala 483 of the partner chain in the B/C interface, while it interacts with Leu 480 on the A/D interface, suggesting that the interacting helices differ by a turn. A similar asymmetry was observed in the LdCoroCC structure too, though the magnitude was larger (2 turns; Nayak et al., 2016; Karade et al., 2020).

3.2. Structural asymmetry and comparison of TbrCoroCC structure with LdCoroCC

Structural superposition of the TbrCoroCC tetramer with LdCoroCC, shows that the structures are similar, the protomers of TbrCoroCC was superposed with LdCoroCC, the structures superpose with a rms deviation of 0.75 Å, while the dimers/tetramers superposing with an overall r.m.s deviation of 1.2–2.2 Å with similar inter-Cβ distances in the A/B, and B/C interfaces but not in the A/D interface, with the D helix is shifted by ~ a turn, smaller than the 2 turns observed in LdCoroCC (Fig. 3A–C, Supplementary Table 1. Mutation of LdCoroCC Ile 486 to alanine, resulted in a symmetric AD/BC interface, but as a consequence, the canonical dimer interface (AB/CD) become asymmetric (Karade et al., 2020). The structural superposition and sequence analysis shows, that there is one heptad shift in case of T.brucei.

To further investigate and identify the cause of asymmetry in TbrCoroCC, the symmetric tetramer was modeled, wherein the D helix is replaced by its symmetry equivalent D*, generated by imposing symmetry at the A/D and B/C interfaces. This was generated by superposing helix A on helix B and using this transformation matrix on helix C to generate the symmetric hypothetical helix D*. Visual analysis of the symmetric tetramer ABCD* reveals that there are steric clashes occurring between helices D* and C, at residues 479 and 482 with 510 and 514 with side chain atoms coming to distances of ~0.8–1.0Å (Fig. 4). The
steric clashes persist, even with alternate rotamer configurations. Significantly, the cyclic salt bridges observed between Arg. 497 and Glu 504 (Fig. 2B) forms only two ionic contacts in the symmetric interface, unlike the eight interactions that stabilize the native structure (Fig. 4). In the elucidated structures, Glu 504 of chain C interacts with Arg 497 of B & D chain with distances 2.8 and 2.6 Å respectively, while Arg B 497 interacts with Glu A 504 (distance 3.1Å) which also interacts with δ-N atom of Arg D 497 (3.2 Å; Fig 4). A similar salt bridge network is also observed in the LdCoroCC structure, with equivalent residues Arg 490 and Glu 497 involved in ionic interactions among three of the protomers, while in the fourth, the side chain points to the solvent (Nayak et al., 2016; Karade et al., 2020).

3.2.1. Sub-cellular distribution and actin coronin co-localization

Wild-type and mutant (L.donovani (I486A, I486A-L493A (I + L), M500V) and T.bruc (V507M & V493I–V507M) coronins were over-expressed in L. donovani promastigotes (Fig. 5). LdAct filaments were analyzed by immunofluorescence microscopy for the number of cells containing LdAct filaments and their lengths. The over-expressed mutant constructs have significantly longer LdAct filaments in comparison to wild type cells. Also, while LdAct filaments were in about 30% wild type cells, the corresponding values for the mutant constructs were ~60%. Interestingly, Leishmania cells overexpressing point mutated versions of T.bruc coronin have more actin filaments when compared to cells expressing native T.bruc coronin. A similar pattern however was not observed for cells expressing point mutated variants of L.donovani coronin, and here the occurrence of LdAct filaments was lesser than to the cells over-expressing native LdCoroCC (Figs. 6 and 7).

The point mutation variants of LdCoro viz. M500V, I486A + L493A and I486A showed similar co-localization pattern as native LdCoro when expressed in Leishmania cells. Intriguingly, the L.donovani cells expressing native Tbr Coro showed similar interaction of coronin and LdAct filaments despite of distinct structure of TbrCoroCC from LdCoroCC. Also the point mutation constructs TbrCoroV493I–V507M and TbrCoro V507M resulted in similar increase in length and occurrence of LdAct filaments when expressed in Leishmania cells though the co-localization was reduced (7A, B) with no co-localization observed in the double mutant (Fig. 7 B bottom right).
4. Discussion

The coiled coil domain of \( T. \) \textit{brucei} coronin showed distinct differences in \( a, d \) residues compared to the \( L. \) \textit{donovani} homolog, whose structure possesses a distinct asymmetry. Structural characterization of \( TbrCoroCC \) shows that the biological molecule is a tetramer, a dimer of dimers, as the homologous \( LdCoroCC \). The oligomer association was additionally confirmed by solution scattering studies and glutaraldehyde cross-linking experiments. Comparative analysis of the \( TbrCoroCC \) with \( LdCoroCC \) shows that the structures are similar, with an r.m.s deviation of ~1.2–2.2 Å, with both having an ade core packing, though small differences are observed in the individual amino acid contributions as well as super-helical parameters.

In the \( T. \) \textit{brucei} coronin CC crystal structures, the asymmetric unit contains a dimer with the tetramer generated by the crystallographic two fold axis. The structure analysis shows that asymmetry causing residues in \( TbrCoroCC \) are distinct from \( LdCoroCC \) with residues Leu 479, Leu 482, Thr 510 and Ile 514 having steric clashes and reduced polar interactions between Arg 497 and Glu 504 in the symmetric model. In the symmetric \( LdCoroCC \) structure also the equivalent residues have steric hindrance, though this region is not in the CC.

\textit{Leishmania} coronin (\( LdCoro \)) upon over-expression increases the length of \( LdCoroCC \) and therefore number of cells showing the filamentous structures of \( LdCoroCC \) also increases (Nayak et al., 2005). It has further been shown that \( LdCoroCC \) interact with the unique region of the linker domain \( LdCoroCC \), though neither the unique domain nor the \( \beta \)-propellor WD40 domain alone is able to promote \( LdCoroCC \) filament formation (Srivastava et al., 2015). This raises a question whether \( \beta \)-propellor domain has an allosteric effect on the quaternary structure of \( LdCoroCC \) due to which \( LdCoroCC \) filament length increases only after full length \( LdCoroCC \) overexpression. Notably, \( T. \) \textit{brucei} coronin (\( TbCoro \)), which has 60% sequence similarity with \( LdCoroCC \), was also able to show patterned co-localization with \( LdCoroCC \) filament and render effects on the filament length, typical to \( LdCoroCC \). The mutant \( TbCoroCC \) structures, generated on the basis of steric hindrance causing residues of \( LdCoroCC \) are identical to the native \( TbCoroCC \) structure, and structural analysis indeed shows that the residues involved in the asymmetry are distinct from \( LdCoroCC \). Therefore, it was obvious to see their \( LdCoroCC \) interaction and effects on filament length. Nevertheless, as an exception, the mutant \( TbCoroCC \) V493I–V507M did not show patterned ________

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Fig. 6. Quantitative analysis of \( LdACT \) filament length in the \textit{Leishmania} (A) cells showing a marked increase in the average length of \( LdCoroCC \) filaments in cells overexpressing mutated coronin variants as compared with wild type cells; length of more than 100 filaments was measured from three independently coronin prepared samples. Statistical significance was drawn through unpaired "t" test ***\( P < 0.0001 \). (B) Quantitative analysis of the number of cells containing \( LdACT \) filaments in native \( LdCoro \), native \( TbCoro \) and mutant coronin overexpressing cells compared with wild type cells. Only the cells that contained >2 μm long \( LdACT \) filaments were considered; the data shown are representative of at least three independent experiments (>1000 cells). The values shown are mean ± standard deviation.

Fig. 7. Immunofluorescence confocal images: (A) showing actin filaments in wild type, \( LdCoro \) over-expressing cells and \( LdCoro \) point mutant variants overexpressing cells. Labelling of \( LdACT \) was done by using anti-\( LdACT \) antibodies. Arrowheads mark filamentous structures of \( L. \) \textit{donovani} actin (\( LdACT \)). (B) \( TbCoro \) overexpressing cells and \( TbCoro \) point mutant variants overexpressing cells. Labelling of \( LdACT \) was done by using anti-\( LdACT \) antibodies. Arrow heads mark filamentous structures of \( L. \) \textit{donovani} actin (\( LdACT \)). Bar, 5 μm.
co-localization with LDAct filaments, though it affected an increase in LDAct filament length. Although it is difficult to explain the reason for this drastic change in interaction behavior of TbrCoro V493I-V507M, could be further examined, by generating mutants that disrupt the oligomer assembly, either in LDCC or TbrCoroCC, by the alteration of salt bridge/apolar interactions that stabilize them. Consequently, our perception that interaction of LDCC with LDAct filaments is an important characteristic which promotes LDAct filament formation is now commuting. It appears that coronin (TbCoro or LdCoro) overexpression has a moonlighting effect on LDAct filament length. Although, it is a testable hypothesis, it provides a clue on our previous observations on no increase in LDAct filament lengths upon overexpression of either beta-propeller or unique + coiled coil domains alone (Srivastava et al., 2015). The observed effects of overexpressing TbCoroc on constructs LdAct makes us speculate on whether a similar effect would be observed in T. brucei actin filaments with the LDCC mutant constructs. However, there are reports that the actin filaments in both these kinetoplastids are distinct: trypanosoma do not contain actin filaments as observed in Leishmania, rather displaying a diffused distribution of actin throughout the cells in the insect stage and only in the endocytic pathway in the bloodstream form (Garcia-Salcedo et al., 2004; Sahasrabuddhe et al., 2004; Nayak et al., 2005); hence, a direct comparison might not be feasible. There could be a strong possibility that beta-propeller is a known protein-interaction scaffold, recruits an unknown LDAct interacting partner that actually promotes LDAct filament formation. As the re-introduction of the hypothesized steric hindrance causing residues of LDCC in TbrCoroCC shows no significant structural changes, the analysis implies that the salt bridge forming residues Arg497 and Glu504 might be the major contributor for asymmetry in TbrCoroCC. Based on the observed asymmetries in LDCC and TbrCoroCC, it is also tempting to hypothesize the asymmetry in LdCoroCC and TbrCoroCC are probably essential for the functionality of kinetoplastid coronins.

Data availability

The coordinates have been deposited in the Protein Data Bank (PDB ID: 7DGX, 7DH4 & 7DH8) and SASDBD Id SASDK46.

CRediT authorship contribution statement

Pankaj Singh Parihar: Conceptualization, Investigation, Validation, Formal analysis, Writing – original draft. Aastha Singh: Investigation, Writing – original draft. Sharanasappa Shimrinta Karade: Conceptualization, Formal analysis, Writing – review & editing. Anmogh Anant Sahasrabuddhe: Conceptualization, Methodology, Formal analysis, Validation, Writing – review & editing. Supervision, Project administration. J. Venkatesh Pratap: Conceptualization, Methodology, Validation, Formal analysis, Writing – review & editing. Supervision, Project administration.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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