Expression and copper binding characteristics of *Plasmodium falciparum* cytochrome c oxidase assembly factor 11, Cox11

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**Abstract**

**Background:** Copper is an essential metal for living organisms as a catalytic co-factor for important enzymes, like cytochrome c oxidase the final enzyme in the electron transport chain. *Plasmodium falciparum* parasites in infected red blood cells are killed by excess copper and development in erythrocytes is inhibited by copper chelators. Cytochrome c oxidase in yeast obtains copper for the CuB site in the Cox1 subunit from Cox11.

**Methods:** A 162 amino acid carboxy-terminal domain of the *P. falciparum* Cox11 ortholog (*PfCox11Ct*) was recombinantly expressed and the rMBP*PfCox11Ct* affinity purified. Copper binding was measured in vitro and in *Escherichia coli* host cells. Site directed mutagenesis was used to identify key copper binding cysteines. Antibodies confirmed the expression of the native protein.

**Results:** rMBP*Cox11Ct* was expressed as a 62 kDa protein fused with the maltose binding protein and affinity purified. rMBP*Cox11Ct* bound copper measured by: a bicinchoninic acid release assay; atomic absorption spectroscopy; a bacterial host growth inhibition assay; ascorbate oxidation inhibition and in a thermal shift assay. The cysteine 157 amino acid was shown to be important for in vitro copper binding by *PfCox11* whilst Cys 60 was not. The native protein was detected by antibodies against rMBP*Cox11Ct*.

**Conclusions:** *Plasmodium* spp. express the *PfCox11* protein which shares structural features and copper binding motifs with Cox11 from other species. *PfCox11* binds copper and is, therefore, predicted to transfer copper to the CuB site of *Plasmodium* cytochrome c oxidase. Characterization of *Plasmodium* spp. proteins involved in copper metabolism will help scientists understand the role of cytochrome c oxidase and this essential metal in *Plasmodium* homeostasis.

**Keywords:** Plasmodium, Malaria, Cox11, Cytochrome c oxidase, Copper

**Background**

Mitochondria with the associated electron transport chain are present in most apicomplexans, have small mitochondrial genomes and appear to be present throughout all the stages of their life cycle [1]. Their electron transport chains have sufficient differences to the electron transport chains of their mammalian hosts to make them promising drug targets [1, 2]. Human malaria caused by *Plasmodium* spp. parasites is the most prevalent apicomplexan infection. The parasite is passed by the bite of an infected mosquito in human infections to the blood of the human host, infects the liver and then red blood cells, which coincides with clinical symptoms. *Plasmodium* parasites in red blood cells depend predominantly on glycolysis for the generation of ATP and electron micrographs show the presence of a single mitochondrion [3, 4]. Despite a reduced

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dependence on mitochondria, the parasites are sensitive to drugs targeting the electron transport chain during this stage of development indicating the importance of the pathway for the parasite [1].

Proteins participating in the electron transport chain contain iron and copper to transport electrons leading to the subsequent generation of ATP. Copper is essential for most living organisms as the metal can cycle between the reduced (Cu\(^{+}\)) and more stable oxidized (Cu\(^{2+}\)) form under physiological conditions and so participate in redox reactions and electron transfer. Copper can be toxic due to its ability to undergo Fenton reactions leading to the generation of toxic reactive oxygen species [5]. Copper can be toxic, can displace metals in proteins, is rarely free in cells and is, therefore, tightly regulated and bound to proteins, like copper chaperones.

During the erythrocytic stage Plasmodium spp. can, like yeast and mammalian cells, import copper via the copper transport protein (Ctr1) expressed initially on the red cell membrane and then the parasitophorous vacuolar membrane [6]. The parasite also obtains copper from the digestion of ingested host erythrocyte Cu/Zn superoxide dismutase [7, 8]. Plasmodium spp. do not express an ATOX-1 (Antioxidant Protein 1) or CCS (Copper Chaperone for Superoxide Dismutase) chaperone homologue that shepherd copper to ATPases in the Golgi apparatus or to Cu/Zn superoxide dismutase respectively in mammalian cells [9]. ATPases are involved in the export of copper from cells and Plasmodium express a copper P-ATPase [8] that is important for parasite fertility [10]. Excess copper is toxic for Plasmodium spp. [11] and chelating copper in vitro growth media with neocuproine or tetrathiomolybdate inhibit the growth of intraerythrocytic parasites [12].

Plasmodium spp. mitochondria have the four complexes of the electron transport chain which includes the copper containing cytochrome c oxidase of complex IV [13]. Cytochrome c oxidase is assembled in many steps, requiring over 30 accessory proteins [14]. The Cox17 protein is a key participant in mitochondrial copper transport to proteins for insertion into cytochrome c oxidase [15]. Plasmodium spp. express Cox17 [16]. Cox17 was first observed in the cytoplasm and mitochondria of yeast which suggested a role in copper transport to mitochondria [17], but subsequent data using Cox17 tethered to the mitochondrial matrix in yeast does not support this suggestion [18]. Cox17 transfers copper to Sco1 and Cox11 which in turn insert copper into the CuA and CuB sites of Cox2 and Cox1 of cytochrome c oxidase [18, 19]. Copper appears to be transported to mitochondria by the phosphate carrier Pic2 in yeast [20] and SLC25A3 in mammalian cells [21].

Cox11 in eukaryotes is attached to the inner mitochondrial membrane via a single transmembrane domain with the N-terminus in the mitochondrial matrix and the C-terminus in the intermembrane space [22–24]. The protein metalates the CuB site of Cox1 with copper in cytochrome c oxidase [25]. Cox11 copper is coordinated by two cysteines in a CFCF motif found in the C-terminal domain [23]. Cox11 is essential in yeast for the assembly of cytochrome c oxidase and the activity of cytochrome c oxidase in Arabidopsis [23, 26].

The present study extended the understanding of Plasmodium spp. copper metabolism and the synthesis of cytochrome c oxidase by studying recombinant P. falciparum Cox11 and showing that the recombinant protein binds copper in vitro and in vivo when expressed in Escherichia coli host cells.

**Methods**

**Ethical clearance**

All procedures involving animals were approved by the University of KwaZulu-Natal, Animal Research Ethics Committee (approval number: 0045/15/Animal).

**Identification of the gene and analysis of Plasmodium spp. Cox11 protein sequences**

Plasmodium falciparum cytochrome c oxidase assembly protein **PfCox11** (PF3D7_1475300) was identified [6] using a BLASTp search of the Plasmodb.org database with the Human Cox11 protein amino acid sequence (UniProtKB-Q9Y6N1 COX11_HUMAN).

Transmembrane domains in the PfCox11 amino acid sequence (PF3D7_1475300) were identified using the TMHMM 2.0 programme [27]. Multiple sequence alignment was generated with Clustal Omega (http://ebi.ac.uk/Tools/msa/clustalo) [28]. Theoretical molecular weight and isoelectric point were determined using PROtparam (http://web.expasy.org/protparam). The P. falciparum Cox11 structure (Fig. 1) was modelled on the Sinorhizobium melloti SmtCox11 homologue NMR structure template (PDB: 1so9) template [24] using the Swiss-Pdb DeepView program [29].

**PCR amplification, mutagenesis and cloning of the C-terminal domain of Plasmodium falciparum PfCox11 and mutants**

The PfCox11 gene portion coding for the predicted C-terminal domain and equivalent to that found in the mitochondrial intermembrane space for the yeast protein [22] was amplified using the following primers: rPfCox11Ct-fwd ccGTCGACAAATTATTTTGTCAATCCACAGG and rPfCox11Ct-rev 5’ ttCGTCAGTCA GGAATAVGCCTTGGAG with added SalI and PstI restriction endonuclease sites shown in bold. Primers for site-directed mutagenesis [30] for the C60A mutation...
were rP/COX11Ct<sub>C60A</sub>-F1 GAAAGACGCAGCAGAC TAATTGCAGCTC, rP/COX11Ct<sub>C60A</sub>-R1 GCCGAG TGCCAAGCTTGCTGCAG, and rP/COX11Ct<sub>C60A</sub>-F2 GTGC-ACCAATTTGCGCAATCCAGG, rP/COX11Ct<sub>C60A</sub>-R2 CTGTGGATTGCGCAATAT-TGGTCGAC. The C157A mutant was amplified using the following primers: rP/COX11Ct<sub>C157A</sub>-F1 GAATT CGATGCCTCTTAGTC rP/COX11Ct<sub>C157A</sub>-R1 TGC GAAGCTTGCTGCAG and rP/COX11Ct<sub>C157A</sub>-F2 CAA TGTTTTCCTTGAAGAC, rP/COX11Ct<sub>C157A</sub>-R2 GTTCTTCA-AAGGCAAAACATTG. The polymerase chain reaction was conducted with a reaction mix (20 µl) containing 1–100 ng template DNA, 0.5 µM of each primer, 2.5 mM MgCl<sub>2</sub>, 0.2 mM dNTPs, 1 U of Phusion DNA polymerase (New England Biolabs) for PCR and site directed mutagenesis, and 1× Taq buffer in a T100™ thermocycler (Bio-Rad, USA). An initial denaturation step (94 °C; 3 min), followed by 35 cycles of denaturation (94 °C; 30 s), annealing (temperature determined by the Tm of the primers, 30 s), and final elongation (72 °C; 30 s) steps. Amplicons of the rP/COX11-Ct were cloned into the pGEMTeasy vector and the plasmid propagated in JM109 E. coli host cells. The plasmid with insert was isolated and digested with SalI and PstI restriction endonucleases. The insert was gel purified and ligated in frame with an N-terminal maltose
binding protein (rMPB) in the pMal-c2x expression vector digested with the same two restriction endonucleases (New England Biolabs, USA). The sequence of the cloned gene was confirmed by DNA sequencing (Central Analytical Facility, Stellenbosch University, South Africa). The mutated gene fragments were PCR amplified, digested with Sall and Pst1 restriction endonuclease and ligated into the pMal-c2x vector as described above.

Expression and purification of recombinant MBPPCox11Ct and mutants

Recombinant pMal-c2c vectors containing the PfCox11Ct insert were transformed into competent E. coli BL21 host cells. A single E. coli transformant was picked from an agar plate and grown overnight (16 h) in 10 ml 2 × YT medium (30 °C, 200 rpm) containing 100 µg/ml ampicillin. A 1:100 dilution of the culture was added to 400 ml 2 × YT medium with 100 µg/ml ampicillin and grown (30 °C, 200 rpm) to an OD_{600} between 0.5 and 0.6. Recombinant protein expression was induced with 0.5 mM IPTG with additional ampicillin. The bacterial cells were pelleted by centrifugation (4000 × g, 4 °C, 10 min) and the supernatant discarded. The pellet was resuspended in 10% of the original culture volume of buffer (20 mM Tris–HCl, 200 mM NaCl, 1 mM EDTA, pH 7.4) and was added to a 1 ml amylose resin (New England Biolabs, USA) equilibrated with Tris–HCl buffer (20 mM Tris, 200 mM NaCl, pH 7.4) and lysed with 9 freeze–thaw cycles (−196 °C and 37 °C) followed by sonication 3 cycles, 30 s/ burst, 30 s on ice between sonication). The lysates were centrifuged (12,000 × g, 4 °C, 25 min). The supernatant was added to a 1 ml amylose resin (New England Biolabs, USA) equilibrated with Tris–HCl buffer (20 mM Tris, 200 mM NaCl, pH 7.4) and was washed with wash buffer (100 mM Tris, 200 mM NaCl, pH 7.4). Bound protein was eluted with the wash buffer containing 0.3 mM maltose and eluted fractions were collected and concentrated with an Amicon Ultra-15 Centrifugation Filter Unit (10 kDa MWCO) by centrifugation (5000 × g, 4 °C) [31, 32]. Samples were dialysed at 4 °C against 3 changes (100 mM Tris–HCl buffer pH 7.4). The Bradford dye binding assay [33] was used to estimate the protein concentration of samples as adapted [34]. Proteins were evaluated on a 12.5% reducing SDS-PAGE gel stained with Coomassie Blue [35].

BCA copper release assay

Purified recombinant rMBP-PfCox11Ct and mutants were tested for their ability to bind copper after purification (in vitro) and during expression (in vivo). Escherichia coli cells expressing the protein intended for in vitro or in vivo copper binding studies were grown in a medium containing 0 or 0.5 mM CuCl₂, respectively, during induction of protein expression [16, 36]. The protein was affinity purified as described above. In the in vitro studies, 10 µM purified rMBP/PfCox11Ct was incubated with 20-fold molar excess of CuCl₂ (15 min; 23 °C). Ascorbate (0 or 10 mM) was included in the reaction mix to reduce free copper. Excess unbound copper was removed by dialysis (2 × 2 h; 23 °C followed by 16 h; 4 °C). The dialysed sample was mixed with 30% (w/v) trichloroacetic acid (3:1 v/v) and centrifuged (12,000 × g; 2 min; 23 °C). Released copper was detected by mixing with bicinchoninic acid (BCA) solution (0.15 mM BCA, 0.9 M NaOH, 0.2 M HEPES) with or without 2 mM ascorbate and the absorbance read at 354 nm [37]. To chelate copper, and thus render it unavailable for binding to a copper binding protein, EDTA was added to the BCA release assay prior to the addition of copper. In the in vivo assay, 10 µM purified rMBP/PfCox11Ct grown in the presence of copper was affinity purified and added to the BCA copper release assay.

Atomic absorption spectroscopy

The amount of copper bound to the recombinant proteins was quantified by atomic absorption spectroscopy (AAS) using an Agilent Varian AA280FS atomic absorption spectrophotometer. The reference solution was a certified atomic absorption standard (Fischer Scientific Co.) prepared at 1.0 ± 0.01 mg/ml in dilute nitric acid. Samples were prepared as described for the in vitro copper binding assessment using the BCA release assay.

Growth of E. coli bacteria expressing recombinant proteins in the presence of toxic concentrations of copper

The influence of rMBPP/Cox11Ct on the growth of E. coli host cells in the presence of toxic concentrations of copper was determined. Overnight cultures of E. coli cells alone or expressing the recombinant MBP or rMBP/Cox11Ct proteins were grown in 2 × YT medium (30 °C, 200 rpm) containing ampicillin (100 µg/ml). At an OD₆₀₀ between 0.5 and 0.6, recombinant expression was induced with 0.5 mM IPTG and additional ampicillin (100 µg/ml) and 0 or 8 mM copper added and bacterial growth (30 °C, 200 rpm) monitored at OD₆₀₀ for 6 h.

Inhibition of copper-catalysed ascorbate oxidation assay

Copper catalyses the oxidation of ascorbate in vitro, and the ability of rMBP/Cox11Ct and mutants to bind copper and inhibit the ascorbate oxidation reaction was assessed. The pH of a 2 mM ascorbate stock solution was adjusted to pH 4.5 with dilute NaOH. The reaction mixture contained 5 µM protein, 120 µM ascorbic acid and 8 µM CuCl₂ (1 ml total volume). Recombinant proteins were added to the reaction mixture and the absorbance of the solution at 255 nm was followed for 300 s at 25 °C [6, 38].
Diffential scanning fluorescence to determine rMBPfCox11Ct thermal melt temperature

The assay described by Niesen et al. [39] with minor modifications was followed. Protein (0.5 µg) in phosphate buffer pH 7.4 and 10 × SYPRO Orange in 25 µl. The assay was run in Xtra-clear qPCR tubes (Star labs) in a Rotor-Gene 6000 RT-PCR machine (Corbett). The condition parameters were high resolution melt (HRM) which has an λ excitation at 470 nm and an λ emission at 570 nm, which was used to measure the SYPRO Orange fluorescence probe. A temperature range of 25–90 °C was used with a ramp temperature of 0.3 °C/s. The change in fluorescence/change in temperature (dF/dT) was calculated and the dF/dT data was plotted against temperature and the highest dF/dT point on each peak was referred to as the T_m.

Antibody production

The amino acid sequence of rMBPfCox11Ct was used to identify immunogenic epitopes with Predict7 software [40]. The peptide sequence KIQF(Abu)F(Abu)EEQMLNAKEEM where internal cysteines were replaced with alpha aminobutyric acid (Abu), was synthesized by GeneScript, USA. The C-terminal cysteine residue was added for m-maleimidobenzyol-N-hydroxysuccinimide ester (MBS) coupling to rabbit serum albumin [41]. Two Hy-line brown laying hens were immunized with the peptide-carrier conjugate equivalent to 200 µg peptide or 50 µg affinity purified rMBPfCox11Ct per immunization, emulsified with Freund's complete adjuvant for three subsequent immunizations at 2-week intervals. Chicken IgY from the yolks of eggs collected 4–16 weeks after the first immunization, was isolated using the PEG6000 precipitation method [42]. The anti-peptide IgY and anti-rMBPfCox11Ct IgY was affinity purified using a peptide or rMBPfCox11Ct affinity column prepared by coupling the respective molecules to a Sulfolink™ or Aminolink™ resin according to the manufacturer’s instructions (Thermo Fisher Scientific).

Plasmodium berghei parasites

Plasmodium berghei parasites (donated by P. Smith University of Cape Town, originally from A.P. Waters Glasgow University) were propagated in male BALB/c mice by intraperitoneal injection of the parasite stabilate (1 × 10^7 parasitized mouse red blood cells) [43]. Parasitaemia was monitored daily on a Giemsa-stained thin blood smear of tail blood [44]. Once parasitaemia reached a suitable level, mice were bled, and the blood collected in a heparinized vacuum test tube.

SDS-PAGE and western blotting

Native and recombinant proteins were detected with affinity purified anti-peptide antibodies in a western blot [45]. Blood from infected mice was washed twice in PBS and resuspended in lysis buffer (20 mM Tris–HCl, 10 mM Na_2EDTA, 1% (v/v) Triton X-100, pH 7.4). The lysates (approximately 10^7 parasites) and recombinant proteins were separated on a 12.5% reducing SDS-PAGE gel, transferred to a nitrocellulose membrane and blocked by incubation in 0.5% (w/v) non-fat milk (Amresco, USA) in Tris-buffered saline (TBS, 20 mM Tris–HCl, 200 mM NaCl, pH 7.4). The membrane was incubated in an appropriate dilution of primary antibody followed by the secondary antibody diluted in 0.5% (w/v) BSA in TBS. The blots were developed by incubation in either 0.06% (w/v) 4-chloro-1-naphthol/0.0015% (v/v) H_2O_2 or Pierce™ enhanced chemiluminescence (ECL) western blotting substrate. Primary antibodies were affinity purified chicken anti- KIQF(Abu)F(Abu)EEQMLNAKEEM peptide IgY (2.5 µg/ml), chicken anti-rPfCox11Ct protein IgY (1 µg/ml) or mouse monoclonal anti-MBP IgG (1:12 000) (New England Biolabs, USA). Detection antibodies were rabbit anti-chicken IgY-horse radish peroxidase (HRPO) conjugate (1:5000) (Jackson Immuno-Research Laboratories, USA) or goat anti-mouse IgG-HPRO conjugate (1:5000) (Roche, Germany).

Results

Bioinformatic

Choveaux et al. [6] identified a putative Cox11 using the human Cox11 amino acid sequence as the input sequence, alongside 13 copper-dependent protein orthologues in the P. falciparum genome from a BLASTp search of the PlasmoDB genome database [46]. The Plasmodium Cox11 sequences lack the 51–61 amino acid N-terminal extension found in the Theileria, Babesia, Arabidopsis, Homo and Mus sequences and as a result have 4, instead of the 5 or 6 cysteines seen in the plant, yeast or mammalian Cox11 sequences (Fig. 1a). The P. falciparum Cox11 gene is found on chromosome 14, and codes for a 228 amino acid protein sequence [47]. The Plasmodium spp. sequences, like all other characterized Cox11 sequences, have a 19 amino acid transmembrane domain (Fig. 1a, b) followed by a conserved cysteine (Cys-60) and 94 amino acids towards the C-terminus an a conserved CFCCF motif (Cys-155 and Cys-157, Fig. 1a, b). There is an 11 amino acid region around the CFCF motif (Cys-60) and 94 amino acids towards the C-terminus a domain (Fig. 1a, b) followed by a conserved cysteine assignment parameters set were high resolution melt (HRM) which has an λ excitation at 470 nm and an λ emission at 570 nm, which was used to measure the SYPRO Orange fluorescence probe. A temperature range of 25–90 °C was used with a ramp temperature of 0.3 °C/s. The change in fluorescence/change in temperature (dF/dT) was calculated and the dF/dT data was plotted against temperature and the highest dF/dT point on each peak was referred to as the T_m.

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The sequence coding for 162 amino acids of the carboxy-terminal domain of the PfCox11 gene was cloned and expressed as a recombinant maltose binding fusion protein and called “rMBPPfCox11Ct”. A His tag fusion protein was insoluble and was not used. The identity of the cloned sequence was confirmed by DNA sequencing. The recombinant MBP fusion protein was affinity purified from E. coli host cell lysates using an amylose affinity matrix (Fig. 2a). A 62 kDa protein that corresponds to the 62.745 kDa, predicted from the cloned gene sequence, and consists of 19 kDa of PfCox11 and 43 kDa of the maltose binding protein fusion partner, was expressed and detected in a western blot by anti-MBP antibodies (Fig. 2b). Truncated versions of the recombinant protein, including a prominent 45 kDa protein, corresponding to the size of the rMBP fusion partner, were also detected by the anti-MBP antibodies. About half a milligram of protein (0.49 mg) was obtained from each gram of wet bacterial pellet (Table 1).

rMBPPfCox11Ct and mutants bind Cu$^{2+}$ in vitro

Recombinant copper chaperones, PfCox17 and the C-terminus of recombinant PfCtr-1, along with copper chaperones from different sources have been shown to bind copper in vitro. Purified rMBPPfCox11Ct was exposed to both Cu$^+$ and Cu$^{2+}$ in vitro before the protein was denatured and the oxidative state of the released Cu determined by the BCA assay in the

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**Fig. 2** Evaluation of rMBPPfCox11Ct on SDS-PAGE and detection with anti-MBP antibodies on a western blot. Proteins were separated on a 12.5% reducing SDS-PAGE gel and stained with Coomassie blue (A) and (B) the same proteins on an identical gel electrophoretically transferred to nitrocellulose and probed with mouse anti-MBP monoclonal antibody and detected with goat anti-mouse IgG-horse radish peroxidase (HRPO) conjugate and enhanced chemiluminescence (ECL). Mw: molecular weight marker; lane 1, soluble E. coli lysate, and fractions from an amylose affinity matrix: unbound (lane 2); wash (Lane 3) and eluted in the presence of maltose (lanes 4–8)
presence or absence of the reducing agent, ascorbate. The rMBP/PfCox11Ct bound cuprous ions (Fig. 3). To identify which particular cysteine residues are involved in copper binding, the PfCox11 Cys-60 and Cys-157 residues were substituted with alanine by site-directed mutagenesis [51]. Single C60A, and C157A mutants and a double C60A/C157A mutant were generated. The amino-acid sequences of the mutants were confirmed by sequencing the respective DNA sequences in the host plasmid. The remaining cysteine (Cys-155) was not mutated as the DNA sequence around the cysteine is AT rich and primer sequences would lack specificity for the targeted site in the experimental approach used. There was no significant difference in the binding of copper in vitro by the rMBP/PfCox11Ct protein or its derivatives containing the three different mutations in the BCA release assay (Fig. 3). The proteins containing the C157A and the C60A/C157A double mutations, like the wild type protein, bound copper in the assay, but appeared to bind marginally less copper than the wild type protein (Fig. 3). The rMPB fusion partner did bind some copper, as has been described before, and was included as a control in all experiments [6].

**EDTA inhibition of copper binding and detection of copper with atomic absorption spectroscopy**

EDTA has been reported to chelate copper and remove copper from the media and the chelated copper is unavailable to bind to molecules [52]. Therefore, it was expected that copper chelated to EDTA would no longer be available for binding to the rMBP/PfCox11Ct. The binding of Cu\(^{2+}\) to rMBP/PfCox11Ct was inhibited by EDTA as measured by the BCA-release assay (Fig. 4a). The binding of copper measured by the BCA-release assay was confirmed (Fig. 4b) with atomic absorption spectroscopy [37]. The rMPB fusion partner bound small amounts of copper as seen in the assays above. All the above assays indicate both rMPB and to a greater extent rMBP/PfCox11Ct bound Cu\(^{2+}\).

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### Table 1 Purification table for rMBP/PfCox11Ct

| Sample                         | Total vol (ml) | Total protein\(^a\) (mg) | Yield (%) | Yield (mg/g pellet) |
|-------------------------------|---------------|--------------------------|-----------|---------------------|
| Soluble E. coli lysate\(^b\)  | 30.15 ± 1.91c | 122.41 ± 2.20            | 100       | 42.07 ± 0.76        |
| Unbound affinity column fraction | 27.46 ± 2.06 | 103.33 ± 0.70            | 84.41 ± 2.09 | 35.51 ± 0.24     |
| Washes                        | 58.50 ± 7.78  | 11.08 ± 1.10             | 9.04 ± 0.74 | 3.81 ± 0.38        |
| Affinity purified rMBP/PfCox11Ct | 23.59 ± 1.47 | 1.43 ± 0.19              | 1.17 ± 0.18 | 0.49 ± 0.06        |

\(^a\) Protein determined by the Bradford protein assay (see Expression and purification of recombinant MBPPfCox11Ct and mutants)

\(^b\) Original cell pellet = 2.91 ± 0.22 g

\(^c\) Data presented are Mean ± SD values from duplicate purifications

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**Fig. 3** Copper binding to rMBP/PfCox11Ct and mutants in vitro measured by the BCA release assay. Affinity purified rMBP/PfCox11Ct or mutants C60A, C157A, C60A–C157A or rMPB were incubated with CuCl\(_2\) in the absence (−) or presence (+) of ascorbic acid in vitro. Copper was detected with the BCA release assay without (open bars) or with (closed bars) the addition of ascorbic acid. Copper chloride was equimolar to protein. Results are the means ± SE of triplicate measurements from duplicate samples. A two-way ANOVA with Bonferroni multiple comparison tests were conducted on the results of samples incubated with CuCl\(_2\) in the presence of ascorbate. Comparisons are between rMBP/PfCox11Ct and mutants and rMPB where *\(P<0.001\)**
Binding of copper to \textit{rMBPPfCox11Ct} and \textit{rMBPPfCox11Ct} mutants in vivo in \textit{E. coli} host cells

The ability of \textit{rMBPPfCox11Ct} and the protein with mutated cysteine residues to bind \( \text{Cu}^+ \) in an in vivo cellular environment was evaluated with two different approaches. Initially, \textit{E. coli} host bacteria with plasmids containing the \textit{PfCox11Ct} DNA sequence or the three mutated sequences were grown in the presence of \( \text{CuCl}_2 \) at non-toxic concentrations of copper and recombinant fusion protein expression induced, followed by protein isolation. Copper binding to \textit{rMBPPfCox11Ct} expressed in copper rich medium was determined with the BCA-release assay. In the second approach the plasmid containing host cells were grown in the presence of increasing levels of copper and host bacteria growth patterns monitored.

\textit{rMBPPfCox11Ct} isolated from \textit{E. coli} grown in the presence of \( \text{CuCl}_2 \) bound copper in the in vivo copper enriched environment. The bound copper was in the \( \text{Cu}^+ \) oxidation state, as inferred from the BCA release assay in the presence and absence of ascorbic acid (Fig. 5). Unlike in the in vitro BCA release assay (Fig. 3), there was more copper bound to the \textit{rMBPPfCox11Ct} and there was a significant difference in copper binding between the \textit{rMBPPfCox11Ct} native sequence and the proteins with mutated cysteines. The \textit{rMBPPfCox11Ct} containing the C60A mutation, bound the same levels of copper as the protein without the mutation (Fig. 5). However, the \textit{rMBPPfCox11Ct} protein with the C157A mutation bound significantly less copper and the protein with the double C60A/C157A mutant bound the least amount of copper (Fig. 5).

In the second assay 8 mM copper chloride inhibited the growth of \textit{E. coli} without a plasmid as seen by a plateau in the \textit{E. coli} growth curve after copper addition (Fig. 6a). When expression of \textit{rMBPPfCox11Ct} was induced it was proposed that the protein would bind copper and enable the host bacteria to continue to grow in the presence of otherwise toxic levels of \( \text{CuCl}_2 \). The bacteria expressing \textit{rMBPPfCox11Ct} continued to divide and grow, though at a lower rate compared to bacteria in the absence of \( \text{CuCl}_2 \) (Fig. 6b). When bacteria expressing the \textit{rMBPPfCox11Ct} protein with the C60A mutant were exposed to 8 mM \( \text{CuCl}_2 \) like the non-mutant (wild type) sample, they grew but at a lower rate than the non-mutant. Bacteria expressing the \textit{rMBPPfCox11Ct} protein with the single C157A or the double C60A/C157A mutation stopped dividing, indicating that the mutants did not bind copper and did not rescue the host cells in the presence of these toxic concentrations of copper (Fig. 6b).

The data from the two in vivo experiments suggests that Cys-60 appears not to be involved in the binding of copper, whilst Cys-157 appears to be involved in copper binding.
Analysis of copper binding by rMBPPfCox11Ct using the rate of ascorbate oxidation

Proteins binding copper in vitro can influence the rate of copper-catalyzed oxidation of ascorbate and this has been used to determine copper binding by PfCox17, PfCtr-1 and other proteins [6, 16, 37]. The rate of copper-catalyzed ascorbate oxidation was monitored at 255 nm in the presence or absence of the native and mutated rMBPPfCox11Ct proteins. The addition of 8 mM CuCl₂ caused a rapid oxidation of ascorbate indicated by a decrease in absorbance at 255 nm (Fig. 7). Oxidation of ascorbate was inhibited by rMBPPfCox11Ct and inhibited to a similar extent by the C60A mutant protein. Both the C157A and the C60A/C157A mutant proteins inhibited ascorbate oxidation to a lesser extent than the native rMBPPfCox11Ct protein.
The inhibition of ascorbate oxidation is interpreted as the binding of copper by the proteins. These results follow the same pattern as the growth inhibition assay and support the importance of the Cys-157 for the binding of copper by PfCox11.

**Thermal shift analysis of rMBP-PfCox11-Ct**

Thermal shift analysis measures the thermal denaturation of a protein [53]. The denaturation profile of a protein can be influenced by the presence of metal ions, drug–protein or protein–protein interactions [39]. The differential scanning fluorescence (DSF) analysis presented in...
Fig. 8 showed a small decrease in the T_m of rMPB alone in the presence (49.12 ± 0.25 °C) or absence of copper (49.61 ± 0.35 °C). There was a more pronounced decrease in the T_m of rMBPP/Cox11Ct in the presence of copper from 55.68 ± 0.12 °C to 52.42 ± 0.49 °C, a ΔT_m of −2.26 °C. The decrease in the thermal transition temperature for rMBPP/Cox11Ct supports the binding of copper to the protein (Fig. 8).

Chicken IgY antibodies against rMBPP/Cox11Ct and the peptide detect the recombinant and native PfCox11 protein

Antibodies were raised in chickens against the affinity purified rMBPP/Cox11Ct fusion protein and against a peptide in the carboxy-terminus of the protein, coupled to a rabbit albumin carrier. The peptide (KIQF(Abu)F(Abu)EEQMLNAKEEM) was chosen based on hydrophathy, antigenicity and surface probability parameters, conservation of the sequence in different species of human infecting Plasmodium spp. and the presence of the conserved CFCF motif (see Fig. 1b). The antibodies were isolated and affinity purified [54, 55] using the rMBPP/Cox11Ct protein or peptide coupled to an affinity matrix. The affinity purified rMBPP/Cox11Ct protein was previously shown to be detected by anti-MBP antibodies (Fig. 2b). The affinity purified rMBPP/Cox11Ct protein was separated on an SDS-PAGE gel (Fig. 9a) and detected by IgY antibodies against the recombinant protein (Fig. 9b) and antibodies against the synthetic KIQF(Abu)F(Abu)EEQMLNAKEEM peptide (Fig. 9d). The anti-peptide antibodies raised against a synthetic peptide derived from the Plasmodium falciparum Cox11 amino acid sequence confirmed the identity of the recombinant rMBPP/Cox11Ct protein (Fig. 9d). All the antibodies detected the 62 kDa recombinant fusion protein and did not detect any E. coli host cell proteins or red blood cell proteins (Fig. 9b, d).

Affinity purified chicken IgY antibodies against rMBPP/Cox11Ct were used to probe lysates of uninfected and P. berghei infected mouse erythrocytes separated on an SDS-PAGE gel (Fig. 9e) and transferred to nitrocellulose (Fig. 9g). An anti-Plasmodium LDH antibody against a conserved PLDH common peptide served as a positive control for the assay and detected PhLDH at 37 kDa (Fig. 9f). The antibodies did not detect E. coli host cell proteins or mouse red blood cell proteins in the western blot of the gel. The antibodies detected Plasmodium proteins at 28 kDa corresponding to the size predicted from the genomic sequence, a 38 kDa protein and a possible dimer and tetramer of 58 and 106 kDa, and a very large protein of undetermined molecular mass (Fig. 9g). This data suggests that the recombinant protein represents the native protein and confirms in silico data indicating that Plasmodium spp. parasites express a Cox11 protein [47].

Discussion

Cox11 is an essential protein in multiple organisms inserting copper into the CuB site of the Cox1 protein of the terminal mitochondrial respiratory chain enzyme cytochrome c oxidase. Plasmodium spp. Cox11 has not been studied to date and little is known about both the Plasmodium spp. cuprome and assembly of Plasmodium cytochrome c oxidase. Sixteen proteins of the Plasmodium cytochrome c oxidase complex have been identified [2]. Plasmodium spp. Cox1, which contains the CuB
copper moieties, and Cox3, are encoded by a mitochondrial gene, whilst the remaining proteins, like Cox2 with the Cu₄ site and Cox11 are encoded by nuclear genes [2, 47]. Here we have begun to characterize *Plasmodium* Cox11 using a recombinant truncated section of the carboxy-terminus of the protein.

The *Plasmodium* spp. Cox11 protein, like the well-characterized Cox11 protein sequences contains a single transmembrane domain (Fig. 1). Tzagoloff et al. [22] proposed that Cox11 is an intrinsic membrane protein in the inner mitochondrial membrane. The C-terminal region of yeast Cox11 can be removed by proteases from mitoplasts indicating its location in the inner mitochondrial membrane space [56] which was confirmed with antibodies [57] and Khalimonchuck et al. [58] proposed an Nᵢn–Cᵢout topology. Given the location of *Plasmodium* spp. cytochrome c oxidase and the structural features of the *Plasmodium* Cox11 the *Plasmodium* spp. protein is proposed to have the same location and topology as yeast Cox11.

The transmembrane region of *P.f* Cox11 is followed by a conserved Cys (Cys-60) and a highly conserved 11 amino acid sequence which includes Lys-152 and the CFCF quartet (Cys-155 and Cys-157) seen in Fig. 1A. The cysteines in the CFCF motif map to the same surface accessible position on a loop in the model of *P.f* Cox11-Ct and the Banci et al. [24] *S. meliloti* template model (Fig. 1c). The three cysteines and the Lys residue have been described as essential for Cox11 to insert copper into the Cu₄ site of cytochrome c oxidase [23, 48, 59]. The extensive N-terminal domain, found in yeast, mammalian and plant homologues was shown by proteolytic cleavage studies not to be involved in copper binding [56, 58].

*Plasmodium* spp. Cox11 is predicted to have a size of 26.8 kDa, similar to the 28 kDa originally described for yeast Cox11 [22]. The gene encoding 228 amino acids of the carboxy-terminus of the protein, without the transmembrane region, was cloned and recombinantly expressed as a 19 kDa section of a 62 kDa rMPB fusion protein and affinity purified on an amylose resin (Fig. 2). Banci et al. [24] expressed Cox11 as a hexahistidine His tag fusion protein from five organisms with the best expression being that of the *S. meliloti* protein. Carr et al. [23] found that the *S. cerevisiae* Cox11-Ct expressed as a His-Cox11-Ct fusion protein was insoluble, but a thioredoxin fusion protein was soluble. Thompson et al. [59] reported poor yields of a His-Cox11-Ct from *Rhodobacter sphaeroides*, but obtained “reasonable yields” of a thioredoxin-Cox11-Ct fusion protein. During the present study there was difficulty expressing the protein as a His tag fusion protein and so a rMPB fusion protein (rMBP/PfCox11Ct) was engineered, expressed and affinity purified.

To compare the *Plasmodium* spp. protein to yeast and bacterial proteins [23, 59], three mutated forms of the protein, two with single C60A and C157A mutations and a double C60A/C157A mutation were expressed and affinity purified. The *Plasmodium* spp. genomes are AT rich and it was consequently not possible in the present study to design suitable primers to modify the Cys-155 residue.

*Plasmodium falciparum* Cox11 was shown to bind copper in a BCA release assay, by atomic absorption spectroscopy, a bacterial host growth inhibition assay, in an ascorbate oxidation inhibition assay and in a thermal shift assay (Figs. 3, 4, 5, 6, 7, 8). The in vitro BCA release assay showed no difference in copper binding between the wild type and the C60A mutant, but there were minor, but not significant, differences between the wild type and the C157A or C60A/C157A double mutant (Fig. 3). When the wild type protein was expressed with additional copper in the medium, more copper was bound to the protein as shown by a 50% increase in absorbance values (Fig. 5). The wild type and the C60A mutant showed little difference in copper binding whilst the C157A and the C60A/C157A mutants bound significantly less copper (Fig. 5). The same trend was observed in the growth pattern of *E. coli* host cells expressing the native and mutated proteins in the presence of growth inhibiting (toxic) concentrations of copper and the ascorbate inhibition assay (Figs. 3, 4, 5, 6, 7, 8). The data indicate that Cys-60 is not important for copper binding in vitro or when the protein is in an in vivo environment, while Cys-157 is essential. The same finding has been reported for *R. sphaeroides* Cox11 [59]. However, though the Cox11 Cys-60 amino acid is not necessary for Cu binding, the amino acid is essential for the insertion of Cu into the Cu₄ site of cytochrome c oxidase in yeast and bacteria [23, 59]. Though we were not able to mutate the Cys-155 amino acid it is very likely that the three cysteines, *P. falciparum* Cox11-Ct 60, 155 and 157 residues, like the same cysteines in Cox11 from yeast and bacteria are essential for physiological function [23, 48].

Mutations of several yeast Cox11 amino acid residues have shown that Lys-205, found three amino acid residues N-terminal to the CFCF motif is essential for a functional cytochrome c oxidase [48]. All *Plasmodium* spp. genomes sequenced to date and a large number of other sequences from multiple species (personal observation) have conserved amino acid residues either side of the CFCF motif including the lysine residue (Fig. 1). Met-224, indicated as essential for the yeast Cox11 sequence, is not present in any *Plasmodium* homologues (Fig. 1b), *Babesia bovis* or *Theileria parva* Cox11 proteins (Fig. 1a)
This confirms that the recombinant protein shares structural features with the native protein and provides physical evidence for the expression of the native protein by the parasite. The antibodies detected five proteins in the parasite lysate. The native protein is 26.8 kDa and a protein of 28 kDa was detected in the western blot (Fig. 9g). The larger protein at 58 kDa may represent a possible Cox11 homodimer or a heterodimer of Cox11 and Cox19 (26.8 and 26.1 kDa) as the two have been shown to interact and co-purify [60]. The 108 kDa protein may represent a Cox11 homotetramer, while the largest protein could represent Cox11 interacting with proteins of the cytochrome c oxidase complex. The western blot could be probed with antibodies against the relevant proteins to support or disprove this suggestion.

Yeast Cox11 can obtain copper from Cox17 [15]. Given the structural similarities between the Plasmodium and yeast proteins shown in this study and by Choveaux et al. [16], it is suggested that Plasmodium Cox11 would also obtain copper from Cox17, which will be evaluated. Yeast Cox11 inserts copper to the Cu_b site on the Cox1 protein of the cytochrome c oxidase [24, 48, 61]. Expression data for mRNA indicates that PfCox17, PfCox11 and PfCox1 are expressed during the sporozoite, trophozoite and gametocyte stages of parasite development [49] and interestingly expression levels of PfCox17 and PfCox11 peak at 30 h and PfCox1 at 40 h post erythrocytic invasion [50]. This suggests an important role for complex IV during the development from trophozoites to schizonts in red blood cells to be investigated further.

This study has confirmed: the presence of native Cox11 in Plasmodium spp., that PfCox11 binds copper in vitro or an in vivo E. coli environment and that Cys-157 is essential for copper binding in vitro. The gene for PfCox11 has a mutagenesis index score and a mutant fitness score of 0.145 and −2.887 which are similar to those of two Plasmodium spp. glycolytic enzymes lactate dehydrogenase and glyceraldehyde-3-phosphate dehydrogenase suggesting the essentiality of the protein for optimal parasite growth [62]. Plasmodium spp. obtain most of their metabolic energy from glycolysis which would explain the essentiality of the glycolytic enzymes [63, 64]. There is uncertainty about the role of the Plasmodium spp. electron transport chain and cytochrome c oxidase or complex IV in the metabolism of Plasmodium spp. [4]. What is known is that inhibitors of complex IV depolarize the membrane potential (Δψ) across the inner mitochondrial membrane and reduce oxygen consumption [65, 66]. Further studies of Plasmodium Cox11 will contribute to the understanding of Plasmodium copper metabolism and the assembly and metabolic role of Plasmodium spp. cytochrome c oxidase and the electron transport chain.

Abbreviations
Abu: Alpha amino butyric acid; ATOX-1: Antioxidant Protein 1; BCA: Bicinchoninic acid; CCS: Copper Chaperone for Superoxide dismutase; Cox: Cytochrome c oxidase; Ctr1: Copper transporter 1; ECL: Enhanced chemiluminescence; HRPO: Horse radish peroxidase; rMBP: Recombinant maltose binding protein; rMBP/PfCox11Ct: Recombinant maltose binding protein Plasmodium falciparum Cox11 carboxy-terminus.

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Author contributions
DG conceived the study and obtained the funding, AS conducted the experiments. AS and DG performed the analysis. DG and AS prepared the manuscript. Both authors read and approved the final manuscript.

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Availability of data and materials
The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.
Declarations

Ethics approval and consent to participate
All procedures involving animals were approved by the University of KwaZulu-Natal, Animal Research Ethics Committee (approval number: 0045/15/Animal).

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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