Localisation of a candidate anion transporter to the surface of the malaria parasite

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Abstract

PfSulP, encoded by the human malaria parasite *Plasmodium falciparum*, is a member of the sulphate permease family of anion transporter proteins. By transfecting the parasite with an epitope-tagged version of PfSulP, and detecting via western blot and indirect immunofluorescent assay microscopy, we show that PfSulP is localised to the surface of the intraerythrocytic parasite, where it is postulated to play a role in the flux of anions across the parasite plasma membrane.

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The malaria parasite, a unicellular eukaryote which, in the course of its complex lifecycle, invades the erythrocytes of its vertebrate host, is responsible for up to three million deaths and close to five billion episodes of clinical illness annually [1]. As it develops within the infected blood cell the parasite is reliant on membrane transport proteins to mediate the uptake of nutrients, the efflux of metabolic wastes, and the maintenance of ion homeostasis. Analysis of the genome of the human malaria parasite *Plasmodium falciparum* has led to the annotation of over a hundred different membrane transport proteins [2]. These proteins are of interest as potential drug targets in their own right, as well as for their possible role(s) in antimalarial drug resistance [3]. However, in all but a few cases, their subcellular localisation, substrate specificity, or physiological role(s) remain to be determined.

The transport of inorganic anions across cellular membranes can affect factors such as the pH, volume and membrane potential of a cell, and in doing so can affect a vast array of cellular processes. The *P. falciparum* protein PF14_0679 (gi:23497754), designated here as PfSulP, is a member of the SulP (sulphate permease) family of secondary active anion transporters and is a likely candidate for the transport of inorganic anions in the malaria parasite. The corresponding mRNA is expressed throughout the intraerythrocytic lifestages of the parasite [2]. Here we present a brief bioinformatic analysis of the PfSulP protein and report its cloning and subcellular localisation.

Materials and methods

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Materials and methods

Bioinformatics. A protein alignment of SulP family members, in which two proteins of the distantly related ‘anion exchanger’ (AE) family (NCBI database Accession Codes 114787 and 10883391) were included as an outgroup, was generated and edited in MacVector™ 7.1.1. Regions that could not be aligned unambiguously were excluded prior to analysis. A phylogenetic tree was estimated using the Neighbour-Joining method [4] and uncorrected (‘p’) amino acid distances. Ties in the tree were resolved randomly and a bootstrap analysis [5] was performed with 1000 pseudoreplicates.

The putative transmembrane topology of PfSulP was established from hydropathy plot analyses using the web-based programs TMHMM v 2.0 (CBS; Denmark), TMPred (EMBnet-CH), and PredictProtein [6], and from the analysis of a protein sequence alignment of the PfSulP family. The resulting figure was generated using the TOPO2 transmembrane protein display tool (SACS, UCSF; http://www.sacs.ucsf.edu/TOPO-run/wtopo.pl).

Subcellular localisation. The subcellular localisation of PfSulP in human erythrocytes infected with mature trophozoite-stage *P. falciparum* (3D7) infected erythrocytes was investigated by transfecting parasites with
a vector encoding a haemagglutinin (HA)-tagged form of the PiSulP protein (PiSulP-HA), with three sequential copies of the HA tag fused to the N-terminus. The full sequence of PiSulP was amplified from *P. falciparum* 3D7 cDNA using appropriate primer sets, and inserted into the vector pDONR®P2R-P3 (Invitrogen) via a BP clonase recombination reaction. A transfection vector, in which expression of the HA-tagged protein is under the control of the ‘chloroquine resistance transporter’ promoter was generated using a vector set developed by van Dooren et al. [7] as described previously [8,9]. The construct was electroporated into 3D7 parasites as described by Wu et al. [10] and transfectants were selected with WR99210 (5 mM).

Expression of the HA-tagged protein was confirmed by Western blot analysis of protein samples prepared as described previously [11]. The proteins were separated on a denaturing gel, labelled with a mouse monoclonal anti-HA antibody (1/5000) and a goat anti-mouse IgG peroxidase conjugate secondary antibody (1/10,000; MP Biomedicals), and detected using a chemiluminescent peroxidase substrate.

For immunofluorescence assays parasitised erythrocytes were fixed and labelled using a protocol described elsewhere [11]. Primary antibodies [a mouse anti-HA antibody, and rabbit PfENT1 antisera (a kind gift from B. Ullman, Oregon Health and Science University)] were used at a 1 in 200 dilution. Secondary antibodies (AlexaFluor 488 goat anti-mouse and AlexaFluor 594 goat anti-rabbit; Molecular Probes) were used at a 1 in 1000 dilution. Nuclei were stained with DAPI. Fluorescence and DIC (differential interference contrast) images were collected using a Leica laser scanning confocal microscope with a 63× water lens.

**Results and discussion**

**Bioinformatics**

PlasmoDB predicts a six-exon structure for the PiSulP gene. However, examination of the genomic sequence surrounding the gene revealed an alternative model with a five-exon configuration, in which the fifth exon is longer (by 42 bases) than predicted. In PCR analyses using primer pairs specific for each gene model, and genomic DNA-free *P. falciparum* cDNA as template, only the coding sequence of the five-exon gene model was able to be amplified, consistent with this (five-exon) model representing the true coding sequence.

PiSulP bears good sequence similarity (BLASTP ‘E’ values ≥ e⁻⁶²) to bacterial members of the SulP family and a phylogenetic analysis ([Fig. 1A](#fig1){refig}) places PiSulP, together with SulP homologues from other apicomplexan species, within a large group containing bacterial homologues.

The SulP family is so-called because the first members to be characterised, from yeast and plants, were identified as SO₄²⁻/H⁺ symporters. However, it has since been demonstrated that SulP proteins can facilitate a variety of anion transport processes. Mammalian SulPs have been shown to catalyse anion exchange processes including, but not limited to, SO₄²⁻/HCO₃⁻ exchange, Cl⁻/HCO₃⁻ exchange, SO₄²⁻/Cl⁻ exchange and Cl⁻/formate exchange. One cyanobacterial homologue (LtnT) has been implicated in nitrate transport [12] while another (BicA) has been shown to catalyse HCO₃⁻/Na⁺ symport [13]. A bioinformatic analysis revealed that many bacterial SulP homologues are fused to homologues of the enzyme carbonic anhydrase [14], prompting the suggestion that they might transport HCO₃⁻ as their primary substrate. By contrast, a SulP homologue from the bacterium *Chloroflexus aurantiacus* is fused to a domain of rhodanese, a thiosulphate:cyanide sulfotransferase [15], consistent with a role as a sulphate permease.

PiSulP is predicted, on the basis of hydrophobicity analysis, to have 12 transmembrane helices ([Fig. 1B](#fig1){refig}), which are predicted (on the basis of the ‘positive inside’ rule [16,17]) to reside on the cytosolic side of the membrane. The cytosolic C-terminus of PiSulP, like that in other SulP homologues, is large (approximately...
140 amino acids long). In many SulP proteins the C-terminus houses a STAS domain (Pfam 01740.12); i.e. a domain with similarity to a class of soluble bacterial proteins called antisigma-factor antagonists [18]. Although PfSulP has not been assigned a Pfam designation of this domain, a BLAST search conducted using only the C-terminus of PfSulP brought up numerous STAS domain proteins including STAS-containing SulPs and antisigma-factor antagonist proteins. The functional role of the STAS domain in SulP proteins has not yet been resolved; however, random mutagenesis of an Arabidopsis homologue has shown that the domain is important for both trafficking and function of the transporter [19].

### Subcellular localisation

To investigate the localisation of PfSulP in the *P.falciparum*-infected human erythrocyte parasites were transfected with a vector encoding an HA-tagged form of the PfSulP protein, with three sequential copies of the HA tag fused to the N-terminus. This approach has been employed previously to localise several other parasite-encoded transporters in the infected cell [8,9].

Fig. 2A shows a Western blot of protein prepared from the transfected parasites and from a non-transfected control, probed with an anti-HA antibody. The protein preparation from the transfectants exhibited a single band that corresponded approximately to the predicted size of PfSulP-HA (~76 kDa). There was no band in the non-transfected control.

In order to determine the subcellular localisation of PfSulP, transfectant cells were fixed, immunostained with fluorophore-conjugated antibodies and observed by fluorescent microscopy. The upper panel of Fig. 2B shows the pattern of HA-associated fluorescence in the transfectants. Fluorescence was localised to the vicinity of the parasite surface. Some fluorescence was also observed around the parasite digestive vacuole; however this was probably due to the reflective properties of this compartment rather than HA-specific staining, as similar vacuole fluorescence was observed in non-transfected control cells (data not shown). No fluorescence was observed at the erythrocyte plasma membrane.

The lower panel of Fig. 2B shows the PfSulP-HA-associated fluorescence co-localised with fluorescently labelled PfENT1, a transport protein shown previously to reside on the plasma membrane of the parasite [20]. HA-associated fluorescence and PfENT1-associated fluorescence localised together at the parasite surface.

Attempts to gain insight into the transport function of PfSulP using the *Xenopus laevis* oocyte expression system proved unsuccessful. Oocytes injected with cRNA encoding PfSulP showed no significant change in the rate of uptake of $^{36}$Cl$^-$, $^{35}$SO$_4^{2-}$, or $^{33}$P. Whether this was due to a lack of functional expression of the protein in the heterologous system, or due to an inappropriate choice of substrate was not investigated.

### Conclusion

In summary, the PfSulP protein belongs to a family of anion transporters. An HA-tagged form of the protein localised to the surface of the parasite within trophozoite-infected erythrocytes, consistent with it being present on the parasite’s plasma membrane. It is therefore a good candidate for playing a role in the movement of anions between the intraerythrocytic parasite and its host cell.
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References