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A malaria membrane skeletal protein is essential for normal morphogenesis, motility, and infectivity of sporozoites

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Membrane skeletons are structural elements that provide mechanical support to the plasma membrane and define cell shape. Here, we identify and characterize a putative protein component of the membrane skeleton of the malaria parasite. The protein, named PbIMC1a, is the structural orthologue of the Toxoplasma gondii inner membrane complex protein 1 (TgIMC1), a component of the membrane skeleton in tachyzoites. Using targeted gene disruption in the rodent malaria species Plasmodium berghei, we show that PbIMC1a is involved in sporozoite development, is necessary for providing normal sporozoite cell shape and mechanical stability, and is essential for sporozoite infectivity in insect and vertebrate hosts. Knockout of PbIMC1a protein expression reduces, but does not abolish, sporozoite gliding locomotion. We identify a family of proteins related to PbIMC1a in Plasmodium and other apicomplexan parasites. These results provide new functional insight in the role of membrane skeletons in apicomplexan parasite biology.

Introduction

Apicomplexan parasites cause major diseases in humans and animals. Amongst these is featured malaria, which remains the most important parasitic human disease responsible for millions of deaths per year. Malaria parasites have a complex life cycle involving many life stages as well as mosquito and vertebrate hosts. Sporozoites infect both hosts and play a vital role in malaria transmission. They possess a specialized cortical structure, the pellicle, which is composed of the plasma membrane and a tightly associated inner membrane complex (IMC). Underneath the IMC lie subpellicular microtubules (SMT), which extend for about two thirds of the cell starting from the apical end. The sporozoite pellicle and associated cytoskeletal elements are conserved in the other invasive stages of Plasmodium, ookinete and merozoites, as well as in invasive stages of other apicomplexan parasites (for review see Morrisson and Sibley, 2002). Recently, the subpellicular network (SPN) was identified in Toxoplasma gondii tachyzoites (Mann and Beckers, 2001). The SPN is a two-dimensional network of filaments situated on the cytoplasmic side of the IMC that takes the shape of the cell and is resilient to detergent extraction, indicating that it acts as a membrane skeleton (Mann and Beckers, 2001). Two structurally unrelated proteins of the tachyzoite SPN, named TgIMC1 and TgIMC2, were identified. TgIMC1 shares sequence homology with articulins, proteins of membrane skeletons of free-living protists, and has been implicated in SPN maturation during development of daughter parasites (Mann and Beckers, 2001; Hu et al., 2002; Mann et al., 2002). The existence of a similar filamentous network in other apicomplexan parasites has been described previously (D’Haese et al., 1977). Moreover, a homologue of TgIMC1 exists in the human malaria species Plasmodium falciparum (Mann and Beckers, 2001), indicating that the SPN is not confined to T. gondii tachyzoites, but constitutes a conserved structure across the Apicomplexa.

Despite the central role of membrane skeletons in cell development, shape, and structural integrity, little is known of the molecular composition and function of the membrane skeleton in malaria parasites. In this paper we identify and characterize the structural orthologue of the membrane skeletal protein TgIMC1 in the rodent malaria parasite Plasmodium berghei and shed light on its role in the parasite life cycle.

Results and discussion

Identification of a TgIMC1 orthologue in P. berghei

We used BLAST homology searches with the TgIMC1 amino acid sequence to identify orthologous genes in Plasmodium. The
Figure 1. Sequence and structure of PbIMC1a. (A) Diagram of the intron/exon structure of the PbIMC1a gene. Introns are indicated with v-shaped lines with sizes in nucleotides. (B) ClustalW multiple alignment of PbIMC1a (Pb), PyIMC1a (Py), PfIMC1a (Pf), PvIMC1a (Pv), and TgIMC1 (Tg). Indicated are conserved domains (shaded); terminal cysteine motifs (bold and in italics); gaps introduced to allow optimal alignment (hyphens). Conserved amino acid identities (asterisks) and similarities (colons and points) are shown underneath. (C) Schematic diagram of the protein structure of TgIMC1 and its structural homologues in Plasmodium spp. The proteins shown are based on predicted proteins of P. yoelii, GenBank/EMBL/DDBJ accession nos. EAA16469 (PyIMC1a), EAA15257 (PyIMC1b), EAA16185 (PyIMC1c), EAA17029 (PyIMC1d), EAA15249 (PyIMC1e), EAA15609 (PyIMC1f), and EAA20426 (PyIMC1g). Boxes mark domains corresponding to the conserved amino-terminal domain (hatched), central domain (gray), and carboxy-terminal domain (open). Conserved terminal cysteine motifs are indicated with open circles.
highest sequence homology (33% amino acid identity) was found with a predicted protein of *Plasmodium yoelii*. We designed primers to this gene to PCR amplify the cognate gene in *P. berghei*. Sequence analysis of two distinct PCR products amplified from genomic and cDNA, respectively, allowed us to map four introns in the *P. berghei* gene, named *PbIMC1a* (Fig. 1 A). The sequence data have been submitted to the GenBank database under accession no. AF542052. The orthologous genes in *P. yoelii* (*PyIMC1a*), *P. falciparum* (*PfIMC1a*) and *Plasmodium vivax* (*PvIMC1a*) are also predicted to contain four introns each at similar positions to those found experimentally in *PbIMC1a*.

The *PbIMC1a* mRNA encodes a predicted cytoplasmic protein of 784 amino acids with a calculated mass of 91,699. Amino acid alignment of *PbIMC1a* with its orthologues reveals three domains of high sequence conservation (Fig. 1 B). These three domains are separated by regions of variable amino acid composition and length. The amino-terminal and central conserved domains, albeit distinct, are both rich in valine (V) and proline (P) and have homology to the VP-rich core domain of articultins from free-living protists. The proteins are further characterized by conserved cysteine motifs at both the amino- and carboxy-terminal ends (Fig. 1 B). *PbIMC1a* and *PyIMC1a* share 80% amino acid identity, reflecting the close ancestry of these two rodent malaria parasite species. Lower, 44–45% identity is found between orthologues of the rodent and human malaria species, whereas the two human malaria species share 51% identity in this protein.

We identified another seven conserved *Plasmodium* proteins with significant sequence homology to *PxIMC1a* (*PxIMC1b* to *h*) containing up to three of the conserved domains (Fig. 1 C). Various homologues can also be identified in the genome sequences available from *Cryptosporidium*, *Toxoplasma*, *Babesia*, and *Eimeria* species (unpublished data), suggesting these molecules belong to a new gene family confined to the Apicomplexa.

Expression and localization of *PbIMC1a*

We studied transcription of the *PbIMC1a* gene by RT-PCR analysis in different life stages of the parasite using cDNA-specific primers. *PbIMC1a*-specific mRNA was detected in samples of asexual blood stages, gametocytes, oocinetes, and sporulating oocysts. The highest amount of *PbIMC1a*-specific mRNA with respect to the reference gene *tubulin-1* was present in sporulating oocysts, whereas *PbIMC1a* mRNA transcription was down-regulated in samples of sporozoite-infected mosquito salivary glands (Fig. 2 A).

To investigate *PbIMC1a* protein expression we raised polyclonal antisera to a region of the protein predicted to have high antigenicity and unique epitopes. Immunofluorescent antibody (IFA) staining with the anti-*PbIMC1a* immune serum labeled sporozoites, but failed to label asexual and sexual life stages including mature schizonts (containing merozoites), gametocytes, and ookinetes (Fig. 2 B). These results indicate that *PbIMC1a* is predominantly expressed during sporogony. Sporozoites isolated from sporulating oocysts and from salivary glands stained positive for *PbIMC1a* (Fig. 2 B). Salivary gland sporozoites typically showed peripheral staining. Moreover, these sporozoites failed to label in live staining (unpublished data). This localization pattern is thus consistent with the position of the IMC/SPN.

The sporozoite-specific expression of *PbIMC1a* suggests the parasite uses different proteins to fulfill functionally equivalent roles in the other invasive stages. The homologues of *PbIMC1a* are prime candidates. Indeed, we detected *PbIMC1b* mRNA in oocinetes samples only (unpublished data), suggesting that *PbIMC1b* could carry out the equivalent role of *PbIMC1a* in this life stage. Mature schizonts of *P. falciparum* were reported to express a protein recognized by anti-*TgIMC1* antiserum, indicating that a homologue of *TgIMC1* is also expressed in the merozoite stage (Mann and Beckers, 2001).

*PbIMC1a* gene disruption

To study the role of *PbIMC1a* we generated *PbIMC1a*-gene-disrupted parasites. A modified *T. gondii* dihydrofolate reductase/thymidylate synthase (*DHFR/TS*) gene cassette (confering resistance to pyrimethamine) was inserted into the
PbIMC1a gene by double homologous recombination replacing nucleotides 550–2,404 of the PbIMC1a gene (Fig. 3 A). Southern analyses of SphI-digested genomic DNA from the resulting parasite (named PbIMC1a-KO) confirmed correct integration into the target locus: A probe corresponding to the central PbIMC1a region gave rise to a single band in parental wild-type (WT), but no signal in PbIMC1a-KO parasites. Conversely, a probe corresponding to the DHFR/TS gene gave rise to three DHFR/TS-specific bands in PbIMC1a-KO, but no signal in WT parasites (Fig. 3 B). In addition, PbIMC1a-specific mRNA was readily amplified by RT-PCR from total RNA samples of WT, but not from PbIMC1a-KO blood stage parasites, confirming disruption of the PbIMC1a gene (Fig. 3 C).

PbIMC1a-KO parasites developed normally in mice and were morphologically indistinguishable from WT parasites in Giemsa-stained blood films. Normal exflagellation and differentiation of gametocytes into ookinetes both in vitro and in vivo was observed (unpublished data). WT and PbIMC1a-KO parasite-infected Anopheles stephensi mosquitoes developed comparable numbers of oocysts in controlled experiments (Table I), indicating that the PbIMC1a-KO ookinetes are capable of normal midgut invasion and subsequent ookinete to oocyst transition. Oocysts in PbIMC1a-KO parasite-infected mosquitoes appeared to develop normally and formed large numbers of sporozoites (Table I). However, closer examination of these sporozoites revealed that they were of abnormal shape and some 20–30% smaller in size (Fig. 4, A and B). Each sporozoite possessed a single enlarged, protruding area associated with the position of the nucleus. The position of these protrusions varied between sporozoites from being located at the posterior end to being positioned near the middle of the sporozoite. Viability of oocyst and hemolymph sporozoites was comparable between WT and PbIMC1a-KO parasites (97–99%; n = 100), indicating that the abnormal cell shape was not the result of premature cell degeneration or death. This is supported by the fact that we did not observe sporozoites with normal morphology.

Western blot analysis of WT oocyst sporozoites gave rise to a major band of ~90 kD (Fig. 4 C), which likely corresponds to the full-length protein. No signal was detected in PbIMC1a-KO oocyst sporozoites (to compensate for the smaller size we loaded at least 50% more of the PbIMC1a-KO sporozoites). PbIMC1a-KO sporozoites were also negative in IFA staining (Fig. 4 D), demonstrating the specificity of the immune serum for PbIMC1a.

Ultrastructural examination of sporulating oocysts showed PbIMC1a-KO sporozoites being less homogeneous in shape.

### Table I. Effects of PbIMC1a knockout on *P. berghei* parasite development in *A. stephensi* mosquitoes

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Parasite [clone]</th>
<th>Mean ± SEM no. of oocysts per mosquito</th>
<th>Percent infection</th>
<th>Mean no. of sporozoites per mosquito&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Midgut</td>
</tr>
<tr>
<td>I</td>
<td>WT</td>
<td>268 ± 34 (30)</td>
<td>100</td>
<td>10,190 (10)</td>
</tr>
<tr>
<td></td>
<td>PbIMC1a-KO [104]</td>
<td>225 ± 30 (29)</td>
<td>100</td>
<td>12,420 (10)</td>
</tr>
<tr>
<td></td>
<td>PbIMC1a-KO [204]</td>
<td>270 ± 33 (30)</td>
<td>100</td>
<td>24,000 (10)</td>
</tr>
<tr>
<td>II</td>
<td>WT</td>
<td>380 ± 53 (10)</td>
<td>100</td>
<td>26,630 (10)</td>
</tr>
<tr>
<td></td>
<td>PbIMC1a-KO [104]</td>
<td>302 ± 69 (10)</td>
<td>100</td>
<td>14,000 (10)</td>
</tr>
<tr>
<td>III</td>
<td>WT</td>
<td>118 ± 19 (20)</td>
<td>100</td>
<td>31,000 (20)</td>
</tr>
<tr>
<td></td>
<td>PbIMC1a-KO [104]</td>
<td>127 ± 18 (20)</td>
<td>100</td>
<td>36,330 (20)</td>
</tr>
</tbody>
</table>

<sup>a</sup>Assessed at 9–11 d after infection.

*References to the tables and figures are corrected to the appropriate sections.*
Figure 4. Characteristics of PbIMC1a-KO sporozoites. (A) Sporozoite shapes as observed by differential interference contrast (DIC). (B) Sporozoite shapes as observed by IFA staining for circumsporozoite protein (CSP, green) and DNA (red). (C) Western blot of oocyst sporozoites from WT and PbIMC1a-KO parasite clones 104 and 204. (D) Negative labeling of a PbIMC1a-KO sporozoite by IFA staining for PbIMC1a. (E) Cross section of sporulating oocysts. (F) Close-up view of E showing budding sporozoites. White arrowheads point at sporozoite budding sites. (G) Cross section of sporozoites. WT: wild-type, KO: PbIMC1a-KO, sb: sporoblast, sz: sporozoite, oc: oocyst wall, nu: nucleus, ap: apicoplast, mi: mitochondrion, pm: plasma membrane, imc: inner membrane complex, smt: subpellicular microtubules. Bars: 5 µm [E], 1 µm [F], 0.1 µm [G].
and size than their WT counterparts and, possibly as a consequence, arranged in a more disorganized fashion inside the oocyst (Fig. 4 E). Developing sporozoites in the process of budding from the sporoblast already appeared to possess the enlarged area around the nucleus (Fig. 4 F), indicating that the effect of the PbIMC1a knockout on cell shape occurs during or before this event. Apart from these visible abnormalities, PbIMC1a-KO sporozoites appeared to possess a complete set of organelles and the assembly of SMT and IMC appeared unaffected by the gene disruption (Fig. 4, F and G).

Given the similarities between Plasmodium sporozoite development by schizogony and Toxoplasma tachyzoite development by endodyogeny, both of which coincide with the assembly of IMC and SMT (Hu et al., 2002; Morriette and Sibley, 2002), a role for PbIMC1a in sporozoite development is conceivable. In tachyzoites, and probably also in sporozoites, this process is concomitant with SPN formation (Mann et al., 2002). In nascent tachyzoites the SPN becomes more stable and rigid upon maturation, a process that is accompanied by proteolytic processing of TgIMC1 for which a conserved carboxy-terminal cysteine is required (Mann et al., 2002). A degree of proteolytic processing of PbIMC1a is also notable in oocyst sporozoites (Fig. 4 C), suggesting that formation of the SPN in sporozoites could involve a similar maturation step.

Infectivity and gliding motility of PbIMC1a-KO sporozoites

We were unable to detect PbIMC1a-KO sporozoites within or associated with the salivary glands (Table I). By contrast, substantial numbers of sporozoites were found in the hemolymph collected from PbIMC1a-KO parasite-infected mosquitoes (Table I). Arguably, these numbers were high enough to allow for detectable salivary gland invasion if the PbIMC1a-KO sporozoites were capable of normal motility, attachment, and Invasion. We were also unable to infect naïve mice by injecting oocyst sporozoites of PbIMC1a-KO parasites under conditions where oocyst sporozoites of WT parasites were infective [4 infected mice out of 5 injected with WT sporozoites; 0 infected mice out of 5 injected with PbIMC1a-KO sporozoites]. These combined observations point to an inability of the mutant sporozoites to invade cells.

Gliding locomotion is necessary for cell invasion in sporozoites (Sultan et al., 1997). Therefore, we examined the ability of the sporozoites to glide in vitro. Observing live gliding of oocyst sporozoites on glass slides clearly showed that locomotion was compromised by the PbIMC1a knockout: null mutant sporozoites were able to attach normally to the glass surface and undergo circular gliding, but their gliding speed was fivefold slower than that of WT sporozoites (WT 2.5 ± 0.4 circles min⁻¹; PbIMC1a-KO 0.5 ± 0.1 circles min⁻¹; n = 7) (Fig. 5 A). Possibly as a consequence of the reduced speed, the percentage of attached PbIMC1a-KO sporozoites observed to be gliding within a set time frame was just half that of WT sporozoites (WT 32%; PbIMC1a-KO 17%; n = 53). These results indicate that PbIMC1a is not a vital component of the motility machinery of the cell, which is closely associated with the pellicle structure (for review see Kappe et al., 2004). Possibly, the reduced motility of the knockout sporozoites is an effect of their abnormal cell shape or size. Alternately, PbIMC1a may interact directly or indirectly with components of the gliding machinery. The molecular interactions involved in gliding are known to be complex and involve multiple protein components besides actin and myosin, such as myosin A tail domain-inter-
acting protein (Bergman et al., 2002), aldolase (Buscaglia et al., 2003; Jewett and Sibley, 2003), and membrane receptors (Gaskins et al., 2004). Given the opportunities for multiple protein interactions, as well as the existence of multiple PbIMC1a homologues, some functional redundancy may exist, resulting in reduced rather than abolished motility. Given the essential connection between gliding motility and cell invasion, the reduced motility of PbIMC1a-KO sporozoites could be partly or solely responsible for their lack of infectivity.

### Mechanical stability of PbIMC1a-KO sporozoites

Assessment of the SPN as done for tachyzoites is difficult to do with sporozoites due to their small size and the lack of a culture system to produce large numbers of cells. These problems preclude a straightforward assessment of the presence and integrity of a SPN in WT and PbIMC1a-KO sporozoites. Given data from *T. gondii* studies and the known functions of membrane skeletons in general (Pumpin and Bloch, 1993), sporozoites with a disrupted SPN are likely to be compromised in their ability to withstand mechanical pressures from inside and outside the cell. This scenario conforms to the observed loss of function phenotypes. First, PbIMC1a-KO sporozoites are of abnormal shape and most notably possess a protruding area around the nucleus. The shape of the nucleus in these sporozoites is rounded, whereas in WT sporozoites the shape of the nucleus is elongated (Fig. 4 B), indicating that they are unable to confine the nucleus and its associated structures within the normally narrow sporozoite frame. Second, PbIMC1a-KO sporozoites appear incapable of cell invasion. Considerable mechanical pressures are applied to sporozoites when they invade cells. Hence, a weakened membrane skeleton could prevent PbIMC1a-KO sporozoites from successfully completing the process of cell invasion.

To assess whether PbIMC1a knockout had affected the mechanical stability of the parasite, we subjected sporozoites to hypo-osmotic shock. Hypo-osmotic conditions cause cells to draw in water and swell, and the degree of hypo-osmotic stress a cell can tolerate is a measure of its mechanical strength (Menke and Jockusch, 1991). Under hypo-osmotic conditions both WT and PbIMC1a-KO sporozoites swelled in one particular area of the cell, namely around the nucleus (Fig. 5 B). Exposure of the cells to 5 min of hypo-osmotic stress caused only a small degree (6.5%) of cell death in WT sporozoites. By contrast, the same treatment caused more than sevenfold higher (47%) cell death in PbIMC1a-KO sporozoites practically halving the population of viable sporozoites (Fig. 5 C). These results show that PbIMC1a-KO sporozoites have considerably lower resistance to osmotic shock, indicating that their ability to withstand mechanical pressure is adversely affected by the *PbIMC1a* gene disruption. This is consistent with PbIMC1a being a membrane skeleton component.

### Materials and methods

**Parasite maintenance, culture, and purification; mosquito infections; and Southern and Western blotting were performed as described previously (Dessens et al., 1999; Sindén et al., 2002).** Sporozoite gliding was assessed as described previously (Vanderberg, 1974; Sultan et al., 1997).

**Sporozoite preparation, viability, and osmotic shock.** Oocyst, hemolymph, and salivary gland sporozoites were isolated as described previously (Kariu et al., 2002) for viability assessment, sporozoites were incubated for 10 min in medium supplemented with 5 mg/l propidium iodide and 1% Hoechst 33258, and were examined by fluorescence microscopy. Sporozoites whose nucleus stained positive for both propidium iodide and Hoechst were scored as nonviable, those staining positive only for Hoechst were scored as viable. Sporozoites were subjected to hypo-osmotic shock (0.33× normal osmotic strength) by adding two equal volumes of water. After 5 min the osmotic strength of the media was returned to normal by adding an appropriate amount of 10× PBS and cell viability was scored.

**Cloning of PbIMC1a.** Parasite genomic DNA and cDNA was prepared as described previously (Dessens et al., 1999). For cloning of PbIMC1a, primers P1F and P1R, designed to the predicted 5’ and 3’ coding sequence of *PyIMC1a*, were used in PCR amplification on *P. berghei* cDNA and genomic DNA. PCR products were analyzed by automated sequencing and primer walking. Stage-specific transcription of *PbIMC1a* was determined by RTPCR using cDNA-specific primers P7F and P7R on different life stages as described previously (Claudians et al., 2002).

**Construction of transgenic parasites.** A 5’ portion of *PbIMC1a* was amplified with primers P13F-KpnI and P13R-HindIII, digested with KpnI and HindIII, and ligated into KpnI/HindIII-digested pBSS-DHFR (Dessens et al., 1999) to give pMC1-KH. A 3’ portion of *PbIMC1a* was amplified with primers P14F-XbaI and P14R-NotI, digested with XbaI and NotI, and ligated into XbaI/NotI-digested pMC1-KH to give the transfection plasmid pMC1a-KO. 50 μg KpnI/NotI-digested pMC1-KO was transfected into purified schizonts followed by pyrimethamine selection and limiting dilution cloning as described previously (Waters et al., 1997).

**Antibody production.** The *PbIMC1a* coding sequence corresponding to residues 456–632 was PCR amplified with primers P22F-BamHI and P23R-XhoI, digested with BamHI and XhoI, and ligated into BamHI/XhoI-digested pGEX-4T-1 (Amersham Biosciences) to give pGEX-PbIMC1a4. Expression and purification of the resulting GST-*PbIMC1a* fusion protein and antiseraum production in mice was as described previously (Dessens et al., 2003).

**Immunofluorescence and electron microscopy.** Parasites were spotted on multwell microscope slides, air dried, and heated on a hot plate at 98°C for 1 min. IFA staining was performed as described previously (Claudians et al., 2002) using anti-*PbIMC1a* immune serum as primary antibody diluted 1:100, and goat anti–mouse FITC-conjugated secondary antibody (Sigma-Aldrich) diluted 1:100. Samples were mounted in Vectashield with DAPI (Vector Laboratories) and examined at RT on a DMR microscope (Leica) with a 100× oil immersion objective lens. Images were acquired using a camera (DC500; Leica) and IM1000 image manager software (Leica). Mosquito midguts with stained oocytes were prepared for transmission electron microscopy as described previously (Sinden et al., 1985) and were examined with a transmission electron microscope (EM 400T; Philips).

**Online supplemental material.** Table S1 shows sequences of oligonucleotide primers. Online supplemental material available at http://www.jcb.org/cgi/content/full/jcb.200406068/DC1.

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