Heparan Sulfate Proteoglycans Provide a Signal to Plasmodium Sporozoites to Stop Migrating and Productively Invade Host Cells

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SUMMARY

Malaria infection is initiated when Anopheles mosquitoes inject Plasmodium sporozoites into the skin. Sporozoites subsequently reach the liver, invading and developing within hepatocytes. Sporozoites contact and traverse many cell types as they migrate from skin to liver; however, the mechanism by which they switch from a migratory mode to an invasive mode is unclear. Here, we show that sporozoites of the rodent malaria parasite Plasmodium berghei use the sulfation level of host heparan sulfate proteoglycans (HSPGs) to navigate within the mammalian host. Sporozoites migrate through cells expressing low-sulfated HSPGs, such as those in skin and endothelium, while highly sulfated HSPGs of hepatocytes activate sporozoites for invasion. A calcium-dependent protein kinase is critical for the switch to an invasive phenotype, a process accompanied by proteolytic cleavage of the sporozoite’s major surface protein. These findings explain how sporozoites retain their infectivity for an organ that is far from their site of entry.

INTRODUCTION

Infection of the mammalian host with malaria is initiated when Plasmodium sporozoites are injected into the skin as the mosquito probes for blood (reviewed in Sinnis and Coppi, 2007). Live imaging studies show that a proportion of the injected sporozoites actively move within the dermis in a random fashion and eventually contact blood vessels which they penetrate to enter the blood circulation (Amino et al., 2006). Once in the circulation, sporozoites arrest in the liver, cross the sinusoidal barrier and invade hepatocytes where they develop into exoerythrocytic forms (EEFs).

Sporozoites must traverse several cell barriers as they make their way from the skin to the liver. Previous studies have shown that sporozoites can interact with cells in one of two ways: they can productively invade a cell, forming a parasitophorous vacuole in which they will replicate, or they can migrate through a cell, breaching the cell’s plasma membrane in the process (Mota et al., 2001). The ability to traverse cell barriers likely enables sporozoites to reach the liver from their injection site in the dermis. Indeed, mutants lacking this ability have reduced infectivity in vivo but not in vitro when they are placed directly on hepatocytes (Bhanot et al., 2005; Ishino et al., 2004).

The molecular signals that allow sporozoites to know where they are and “decide” whether to continue to migrate or prepare for cell invasion are not known. That the sporozoite recognizes different cell types was suggested by recent studies on the proteolytic cleavage of the sporozoite’s major surface protein, the circumsporozoite protein (CSP; Coppi et al., 2005). CSP is processed by a parasite cysteine protease and cleavage is specifically associated with productive invasion and not cell traversal. Since sporozoites contact cells during both invasion of and migration through cells, these data suggested that sporozoites recognize different cell surface molecule(s) on permissive versus non-permissive cells. Sporozoites are known to bind heparan sulfate proteoglycans (HSPGs; reviewed in Sinnis and Coppi, 2007). Using the rodent malaria model, Plasmodium berghei, we show that contact with cells expressing highly sulfated HSPGs activate sporozoites to begin the invasion process. In contrast, contact with less sulfated HSPGs results in continued migration through cells. Furthermore, we show that the transition from a migratory to an invasive phenotype is mediated by a signaling pathway that includes a new member of a family of calcium-dependent protein kinases.

RESULTS

A New Sporozoite Migration Assay

We began by developing a quantitative migration assay since the currently used assay, in which fluorescently labeled dextran is taken up by cells that are wounded during sporozoite migration, allows one to observe wounded cells but does not quantify the frequency with which a cell is wounded (Mota et al., 2001). When added to cell
monolayers, sporozoites move in circles so that sporozoites that migrate for a long time wound a few cells over and over again whereas those that migrate for a short time wound the same few cells but less frequently. This difference in migratory behavior can be measured by determining the extent of release of an intracellular dye such as calcein green-AM. When Hepa 1-6 cells were loaded with calcein green-AM, incubation with freshly isolated sporozoites caused measurable release of the dye, while uninfected salivary gland material had no effect (Figure 1). This result was dependent on sporozoite motility as treatment with cytochalasin D, an inhibitor of motility, caused minimal release of calcein.

We next compared sporozoite migratory activity in the presence or absence of E-64d, a cysteine protease inhibitor that inhibits CSP processing and productive invasion (Coppi et al., 2005). Treatment with E-64d caused a fivefold increase in calcein release, suggesting that when sporozoites are inhibited from productively invading a cell, they continue to migrate (Figure 1). In contrast, the dextran-uptake assay does not distinguish between E-64d treated and untreated sporozoites (Figure 1). Because repeated wounding of a cell may lead to its rupture with release of all intracellular calcein, the calcein assay may not be completely linear, although it would still reflect the sporozoites’ overall migratory activity. Importantly, E-64d does not interfere with the resealing of the wound in Hepa 1-6 cells as cells mechanically wounded in the absence or presence of E-64d showed no difference in the number of dextran-positive cells (Figure 1).

Highly Sulfated HSPGs Inhibit Migration and Promote Invasion by Sporozoites

Previous work has shown that sporozoites bind to the glycosaminoglycan chains (GAGs) of HSPGs (reviewed in Sinnis and Coppi, 2007). In addition, the sulfate moieties of GAGs are critical for sporozoite binding and a high overall density of sulfation is required (Pinzon-Ortiz et al., 2001). To determine whether highly sulfated heparan sulfate might direct sporozoites to stop migrating and actively invade hepatocytes, we treated Hepa 1-6 cells with chlorate, a metabolic inhibitor of sulfation that decreases the extent of GAG sulfation (Humphries and Silbert, 1988), and monitored sporozoite migration. Previous studies in hepatoma cells indicated that treatment with 10 mM and 30 mM chlorate decreased incorporation of $^{35}$SO$_4$-sulfate into proteoglycans by 60% and 75% respectively, with no effect on protein synthesis or cell growth (Pinzon-Ortiz et al., 2001). When sporozoites were added to cells pretreated with chlorate, their migratory activity increased in a manner dependent on the dose of chlorate (Figure 2A). At 20 mM chlorate, their migratory activity was equivalent to that of parasites treated with E-64d. Importantly, chlorate-treatment had the opposite effect on sporozoite invasion with a dose-dependent inhibition of this process (Figure 2A and Figure S1 in the Supplemental Data available with this article online). The effect was specific to inhibition of sulfation by chlorate, as adding back sulfate to the chlorate-treated cells restored the ability of sporozoites to productively invade (Figure S1). These findings indicate that the sulfation state of HSPGs dramatically changes the nature of the interaction between sporozoites and host cells.

Since reliable markers for the early parasitophorous vacuole (PV) in sporozoite infection are lacking, it is difficult to distinguish invading from migrating sporozoites. To address this issue, we performed invasion assays in the presence of E-64d, which inhibits productive invasion of cells (Coppi et al., 2005) and therefore allows us to approximate the number of intracellular parasites that are in the process of migrating. As shown in Figure 2A, there are few intracellular sporozoites in the presence of E-64d, indicating that the majority of intracellular sporozoites in the absence of E-64d have productively invaded. Chlorate
had no effect on the percentage of intracellular sporozoites when sporozoites were preincubated with E-64d (Figure S1). These data suggest that sulfation-dependent invasion and the triggering of CSP processing observed when sporozoites contact hepatocytes (Coppi et al., 2005) may share a similar mechanism.

Since chlorate is a general inhibitor of macromolecular sulfation, we also performed experiments with a CHO cell mutant specifically deficient in sulfation of heparan sulfate (HS) chains. The mutant, CHO pgsE, is derived from the parent cell line CHO K1 and has a mutation in N-deacetylase/N-sulfotransferase (Ndst1), which results in the formation of HS with less overall sulfation (Esko et al., 1985). Sporozoite migratory activity was increased in CHO pgsE cells compared to the parental wild-type cell line (Figure 2B). However, sporozoite migration

Figure 2. Highly Sulfated HSPGs Arrest Sporozoite Migration and Trigger Invasion and CSP Processing
(A) Effect of chlorate treatment on migration, invasion, and CSP processing. Hepa 1-6 cells were grown in the presence of the indicated amounts of chlorate and washed, and P. berghei sporozoites preincubated ±E-64d were added for 1 hr (migration and invasion assays) or 3 min (CSP cleavage assay). To measure migration (gray bars), chlorate-treated Hepa 1-6 cells were preloaded with calcein green, sporozoites were added, and the fluorescence released into the supernatant was measured. Shown are the means of duplicates which did not vary more than ±5%. Invasion (black bars) was quantified by fixing, staining, and counting the numbers of intracellular and extracellular sporozoites. Shown are means ± SD of triplicates. Proteolytic cleavage of CSP (pie charts) was quantified by adding GFP-expressing sporozoites to cells for 3 min and fixing and staining with antiserum specific for full-length CSP. One hundred sporozoites per coverslip were counted, and the pie charts show the percentage of sporozoites with cleaved (white) or full-length (hatched) CSP. Shown are the means of triplicates, which did not vary more than ±5%. All experiments were performed at least three times, and shown are representative experiments.

(B) A CHO cell mutant in HSPG sulfation enhances sporozoite migratory activity and inhibits invasion and CSP cleavage. Sporozoites were incubated with the indicated cell type, and migration (gray bars), invasion in the absence (black bars) or presence (white bars) of E-64d, and CSP cleavage (pie charts) were quantified as described above. For the invasion assay, shown are means ± SD of triplicates. Migratory activity is measured as fluorescence a.u., and shown are the means of duplicates, which did not vary more than ±5%. Pie charts show the means of triplicates, which did not vary more than ±5%. All experiments were performed three times, and shown is a representative experiment.

(C) Quantification of CSP cleavage induced by highly sulfated HSPGs. Sporozoites were metabolically labeled with 35S[SO4]-Cys/Met and kept on ice (lane 1) or chased for 1 hr (lanes 2–4). They were then spun onto coverslips without cells (lanes 1 and 2) or with cells grown in the absence (lane 3) or presence (lane 4) of chlorate and brought to 37°C for 3 min. Cells and sporozoites were then lysed, and CSP was immunoprecipitated and analyzed by SDS-PAGE and autoradiography. Band intensity was quantified by densitometry and is displayed graphically on the right; gray bars represent uncleaved CSP, and black bars represent cleaved CSP.

Heparan Sulfate Regulates Sporozoite Invasion

Table 1. Disaccharide Analysis of Heparan Sulfate on Hepa 1-6 Cells, CHO Cells, Mouse Dermal Fibroblasts, and the Endothelial Cell Line HBMVEC

<table>
<thead>
<tr>
<th>Sample</th>
<th>ΔUA-GlcNAc</th>
<th>ΔUA-GlcNS</th>
<th>ΔUA-GlcNAc6S</th>
<th>ΔUA-GlcNS6S</th>
<th>ΔUA2S-GlcNAc6S</th>
<th>ΔUA2S-GlcNS6S</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hepa 1-6</td>
<td>33.21</td>
<td>30.73</td>
<td>10.05</td>
<td>5.52</td>
<td>12.70</td>
<td>0.24</td>
</tr>
<tr>
<td>CHO K1</td>
<td>46.17</td>
<td>13.28</td>
<td>5.52</td>
<td>2.32</td>
<td>22.47</td>
<td>0.00</td>
</tr>
<tr>
<td>CHO pgsE</td>
<td>67.07</td>
<td>14.95</td>
<td>1.20</td>
<td>2.55</td>
<td>6.51</td>
<td>0.00</td>
</tr>
<tr>
<td>MDF</td>
<td>46.08</td>
<td>29.63</td>
<td>10.32</td>
<td>5.67</td>
<td>4.55</td>
<td>0.04</td>
</tr>
<tr>
<td>HBMVEC</td>
<td>69.86</td>
<td>20.08</td>
<td>5.04</td>
<td>2.46</td>
<td>0.78</td>
<td>0.13</td>
</tr>
</tbody>
</table>

Values are expressed as molar percent and normalized to total protein.

through wild-type CHO cells was increased compared to Hepa 1-6 cells, a difference that is not unexpected, as the HS chains on CHO cells are less sulfated than those found on hepatoma cells (Table 1). Invasion efficiency in these cell lines was inversely correlated to sporozoite migratory activity, with fewer parasites invading the CHO cells with reduced HS sulfation (Figure 2B). Treatment of sporozoites with E-64d, before addition to CHO cells, demonstrated that the majority of intracellular sporozoites had indeed productively invaded (Figure 2B).

Sporozoites Preferentially Migrate through Dermal Fibroblasts and Endothelial Cells and Do Not Efficiently Invade These Cells

The finding that sulfation of HS chains plays a role in the sporozoite’s “decision” to migrate or invade a cell is likely to be relevant in vivo, where the density of HS sulfation differs among organs. HSPGs on hepatocytes are among the most highly sulfated in the mammalian host, and those found on endothelial cells and in the dermis are, in comparison, less sulfated (Lindblom and Fransson, 1990; Lyon et al., 1994). Our data, therefore, suggested that sporozoites use the sulfation density of HSPGs to ascertain where they are and whether they should continue to migrate or prepare for cell invasion.

To test this hypothesis, we studied the interaction of sporozoites with cell types they are likely to encounter on their way to the liver, namely, dermal fibroblasts and endothelial cells. First, we examined the sulfation of HS produced by mouse dermal fibroblasts (MDF) and endothelial cells (HBMVEC) compared to Hepa 1-6 cells and CHO cells (Table 1). The overall extent of sulfation, as measured by summing the amount of N-sulfated glucosamine, 6-O-sulfated glucosamine, and 2-O-sulfated uronic acids, increased from 0.4 sulfate residues/disaccharide in HBMVEC to 0.6, 0.8, and 1 sulfate/disaccharide in MDF, CHO, and Hepa 1-6 cells, respectively (Table 1). Thus, hepatocytes produce HS with a higher content of sulfate compared to other cells a sporozoite may encounter in the mammalian host.

Sporozoite migration experiments in dermal fibroblasts, endothelial cells, and hepatocytes showed that migratory activity correlated inversely with overall sulfation of the cell type encountered (Figure 3A and Table 1). To insure that wounding by sporozoites had the same effect in these different cell lines, we incubated each cell type with maximally migrating, i.e., E-64d-treated sporozoites. Similar amounts of calcein were released by each cell line, indicating that inherent differences among them do not lead to significant differences in the damage caused by migrating sporozoites (Figure S2).

In contrast to the migratory activity of sporozoites in dermal fibroblasts and endothelial cells, productive invasion and CSP processing were decreased in these cells (Figure 3A). In addition, when sporozoites were placed on freshly excised mouse skin they retained full-length...
CSP on their surface, whereas those placed on liver sections had predominantly cleaved CSP on their surface (Figure S3).

The number of intracellular sporozoites in MDFs and HBMVECs was higher than that observed with E-64d-treated parasites, suggesting that sporozoites can productively invade these cells. Further studies showed that sporozoites could develop in dermal fibroblasts and endothelial cells, albeit at a low efficiency (Figure 3B). Overall, these data suggest that, after their injection into the mammalian host, sporozoites preferentially migrate through those cell types in which they cannot efficiently replicate and are activated, by the high level of HS sulfation, to productively invade hepatocytes.

**Soluble Heparin Induces CSP Cleavage and Augments Invasion of Cell Lines with Low Levels of Sulfated HSPGs**

To determine whether the effect of highly sulfated HS chains on sporozoites was dependent upon cell contact, we tested the effect of soluble heparin (a highly sulfated form of heparan sulfate) on CSP cleavage and sporozoite invasion. In the absence of cells, 5 μg/ml of soluble heparin induced CSP cleavage in 40% of sporozoites (Figure 4A). We then tested whether a short preincubation with this concentration of heparin could enhance sporozoite infectivity for cells that have undersulfated HSPGs. Indeed, soluble heparin could partially compensate for the lack of sulfated HSPGs on these cells and enhanced sporozoite infectivity for these normally less permissive cell lines (Figure 4B). Interestingly, the dose-response curve for CSP cleavage had an inverted bell shape. One explanation for this type of dose response is that heparin at low dose allows CSP multimers or CSP/protease complexes to form, which facilitate cleavage and invasion. At high doses of heparin, complexes fail to form due to titration of the individual factors. Activation of growth factor signaling by heparin shows a similar response, which has been interpreted as binding of heparin to both the growth factor and the receptor (Pellegrini, 2001).
Heparan Sulfate Regulates Sporozoite Invasion

**Sporozoite Binding to Heparin Induces a Signaling Cascade**

The ability of highly sulfated HSPGs on the host cell surface (and soluble heparin) to trigger CSP cleavage and productive invasion of cells suggests that downstream signaling events occur following the sporozoite’s interaction with optimally sulfated HSPGs. Therefore, we tested the effect of staurosporine, a broad-spectrum protein kinase inhibitor, previously shown to inhibit *Plasmodium* invasion but not motility (Mota et al., 2001; Ward et al., 1994), on sporozoite migration, invasion, and CSP cleavage. Pretreatment with staurosporine increased sporozoite migration, while invasion and CSP cleavage were inhibited, indicating that a parasite protein kinase was potentially involved in this process (Figure 5).

Although staurosporine is not specific for a particular family of protein kinases, it is slightly more active against kinases of the protein kinase C (PKC) family, which in higher eukaryotes are activated by calcium and play a central role in calcium-mediated signaling. No obvious PKC-like kinases have been found in the *Plasmodium* genome database (Ward et al., 2004); however, *Plasmodium* has a family of calcium-dependent protein kinases (CDPKs; Ward et al., 2004) that function to transduce calcium-mediated signals (Billker et al., 2004; Ishino et al., 2006; Siden-Kiamos et al., 2006). Since calcium signaling plays a central role in the regulation of cell invasion in both *Plasmodium* and the related parasite, *Toxoplasma gondii* (Carruthers and Sibley, 1999; Mota et al., 2002), we hypothesized that, after sporozoite binding to highly sulfated HSPGs, one of the CDPKs may play a role in the sporozoite’s transition from a migratory to an invasive phenotype. Selective CDPK inhibitors have not been described, but the calmodulin antagonist W-7 targets the calmodulin-like domain of plant CDPKs, and KN-93 inhibits the structurally related animal calmodulin-dependent protein kinase II. Like staurosporine, these inhibitors enhanced migration but inhibited cell invasion and CSP processing (Figure 5). Purvalanol A, an inhibitor of cyclin-dependent kinases, was used as a control and had no effect on sporozoite migration, invasion, or CSP cleavage. Together, these data suggest that binding of sporozoites to highly sulfated HS chains triggers a signaling cascade involving one of the *Plasmodium* CDPKs.

**CDPK-6 Signaling Is Required for the Switch from Migration to Invasion**

CDPKs in *Plasmodium* are present as a multigene family, and initial mining of the *Plasmodium falciparum* genome database suggested that this family had five members (Ward et al., 2004). With the exception of CDPK-2, all of the CDPK genes have clear orthologs in the rodent malaria parasite genomes (Hall et al., 2005). Our analysis indicates that a sixth conserved gene, which we call CDPK-6, should be grouped within the CDPK family. CDPK-6 is predicted to encode an atypically large protein, characterized by a CDPK-like kinase domain, an incomplete carboxy-terminal calmodulin-like domain, and an unusually large amino-terminal extension. The *P. falciparum* transcriptome shows that transcription of PICDPK-6 (PF11_0239) is significantly upregulated in the sporozoite stage (Le Roch et al., 2003). To determine if this kinase has a role in the switch from migration to invasion, we generated a CDPK-6 gene knockout by replacing the coding region of *P. berghei* CDPK-6 (PB001122.01.0) with a resistance marker via double homologous recombination (Figure 6A). Resistant parasites were selected and cloned, and integration of the targeting construct into the CDPK-6 locus was verified by pulse-field gel electrophoresis and Southern blot analysis (Figure 6B).

CDPK-6 mutant parasites produced fewer salivary gland sporozoites (R.T., A.C., P.S., and O.B., unpublished...
data); however, we were able to obtain sufficient numbers for the present study. As shown in Figure 6C, CDPK-6 mutant sporozoites displayed enhanced migratory activity and were significantly less infective for hepatocytes. Importantly, preincubation of CDPK-6 mutant sporozoites with soluble heparin could not enhance their invasive capacity (Figure 4B), further supporting our hypothesis that CDPK-6 transduces the signal resulting from sporozoite contact with highly sulfated HSPGs. We have confirmed these findings in vivo, where there was significant delay in the time to detectable blood-stage infection with CDPK-6 mutant sporozoites compared to wild-type (data not shown). In addition, the majority of CDPK-6 mutant sporozoites did not cleave CSP upon contact with hepatocytes (Figure 6C, pie charts), and pulse-chase metabolic labeling experiments confirmed that they were deficient in their ability to cleave CSP (Figure 6D).

The finding that CDPK-6 mutant sporozoites can productively invade Hepa 1-6 cells to a small extent (Figure 6C, CDPK-6 KO ± E-64d) suggested that CDPK-6 was not the only kinase involved in the transition to an invasive phenotype. To test this, we performed invasion assays with wild-type and CDPK-6 mutant sporozoites in the presence of the protein kinase A (PKA) inhibitor H-89, since PKA inhibitors have been shown to affect activation of sporozoites (A. Rodriguez and L. Cabrita-Santos, personal communication). This inhibitor decreased invasion by wild-type sporozoites by approximately 30% and further decreased invasion of the CDPK-6 mutant sporozoites (Figure 6C). In agreement with this finding, H-89 enhanced migration of wild-type sporozoites and further enhanced the migratory activity of CDPK-6 mutant sporozoites. Together, these data suggest that CDPK-6 works in concert with other signaling molecules to induce an invasive phenotype. These data were confirmed with a second CDPK-6 mutant generated by an independent transfection experiment (Figure S4).

DISCUSSION

In this report we show that HSPGs provide an environmental signal that modulates the behavior of Plasmodium sporozoites. Using chlorate-treated Hepa 1-6 cells and a CHO cell mutant defective in an enzyme required for HS sulfation, we found that sporozoites preferentially migrate in the presence of HSPGs with low levels of sulfation, whereas contact with cells expressing highly sulfated HSPGs triggers CSP cleavage and productive invasion. These findings are of relevance in the mammalian host, where we found that sporozoites preferentially migrate through dermal fibroblasts and endothelial cells and productively invade hepatocytes. Importantly, the HS chains expressed on the surface of these different cell types parallel the HSPG sulfation level of the organs from which they are derived. These data, therefore, suggest a mechanism by which sporozoites navigate in the mammalian host and retain their infectivity for a cell that is some distance from where they are deposited by infected mosquitoes.

Sporozoites recognize these highly sulfated HS chains, likely via binding of their major surface protein, CSP. Previous studies have shown that CSP has two heparin-binding sites, one in the NH2 terminus and one in a cell-adhesive motif, the type 1 thrombospondin repeat (TSR), found in the COOH-terminal portion of the protein (Rathore et al., 2002; Sinnis et al., 1994). Our model is that the NH2-terminal portion of CSP binds to and is cross-linked by highly sulfated HSPGs, resulting in a signaling cascade that is mediated, at least in part, by CDPK-6. Activation of CDPK-6 leads to secretion of the protease that cleaves CSP, thereby exposing the TSR, which binds with high affinity to HSPGs, resulting in productive invasion of the hepatocyte (Figure 7). Thus, this model encompasses glycosaminoglycan-dependent induction of a key signaling response in a pathogen.
Several lines of evidence indicate that, in Plasmodium, CDPKs translate calcium signals into a wide variety of stage-specific end points. In gametocytes, CDPK-4 transduces the calcium signal in male gametocytes that leads to exflagellation (Billker et al., 2004). In ookinetes, mobilization of intracellular calcium and transduction of this signal through CDPK-3 enables the ookinete to glide and penetrate the mosquito midgut wall (Ishino et al., 2006; Siden-Kiamos et al., 2006). Previous studies have shown that calcium is an important second messenger regulating invasion of Plasmodium sporozoites and Toxoplasma tachyzoites (Mota et al., 2002; Carruthers and Sibley, 1999), and here we show that this signal is, in part, transduced by CDPK-6. Together, these studies suggest that Plasmodium CDPKs are a family of protein kinases that translate the ubiquitous second messenger calcium into different stage-specific cellular responses.

The function of cell traversal by sporozoites has been the subject of much debate. Although this study does not directly address the role of migration, our finding that a sporozoite’s migratory activity is inversely correlated to how efficiently it invades a particular cell line demonstrates that migration per se is not sufficient to promote productive invasion. A previous study has suggested that cell traversal activates sporozoites for productive infection via a hepatocyte cytosolic factor (Mota et al., 2002). However, our data demonstrate that sporozoite activation is primarily due to contact with highly sulfated HSPGs on the cell surface and can, at least in part, be induced with soluble heparin in a cell-free system. Nonetheless, our data as well as that of others, indicate that a small amount of migration takes place within hepatocytes (Figures 2, 3, and 5 and Mota et al., 2002), suggesting that sporozoite activation is a progressive process that is initiated by contact with highly sulfated HSPGs. Continued migration through hepatocytes, therefore, may lead to exposure of additional adhesins that are required for productive invasion. The idea that activation is a progressive...
process is consistent with previous data suggesting that highly sulfated HSPGs of hepatocytes protrude through the open fenestrations of the liver sinusoids and arrest circulating sporozoites (reviewed in Sinnis and Coppi, 2007). The sporozoite’s switch to “invasion mode” may therefore begin in the liver sinusoid and progress as the sporozoite crosses this barrier and enters the liver parenchyma.

The switch to “invasion mode” is no doubt accompanied by a multitude of events, many of which have not yet been elucidated. Recently, the secretion of two proteins with 6-cys motifs, Pbs36p and Pbs36, has been associated with productive invasion (Ishino et al., 2005). However, whether they play a role in initiating invasion or at some later point is not yet known (van Dijk et al., 2005). In this study we found that proteolytic cleavage of CSP is an early event in this process. Although the responsible protease is not yet known, our data suggest that the switch to invasion mode results in either secretion of the protease onto the sporozoite surface or activation of a protease that may already be present on the parasite surface. Nonetheless, our findings elucidate both the host cell molecule initiating proteolytic processing of a Plasmodium adhesin and the kinase that transduces this signal, perhaps providing a model for adhesin-specific processing in other Plasmodium life-cycle stages and other Apicomplexa. Further studies to determine additional events that occur upon CDPK-6 signaling in response to contact with highly sulfated HSPGs are ongoing.

In conclusion, our data suggest that migration through cells is the default behavior of the newly injected sporozoite and that this behavior is maintained as long as the sporozoite encounters cells expressing HSPGs with low levels of sulfation. Once in the liver, where the HSPGs are highly sulfated, the sporozoite undergoes activation via a calcium-dependent kinase, resulting in controlled proteolysis of surface CSP and a shift to invasion mode to begin the next stage of malaria infection (Figure 7). These findings begin to unravel how Plasmodium sporozoites optimize their infectivity for an organ that is a significant distance from the skin, where their journey begins.

**EXPERIMENTAL PROCEDURES**

**Parasites**

Three- to five-day-old Anopheles stephensi mosquitoes were fed on Swiss Webster mice infected with either P. berghei (ANKA or NK65 strain), GFP-transgenic P. berghei (Janse et al., 2006), or the CDPK-6 mutant. For experiments with CDPK-6 mutant parasites, controls were the ANKA strain. On days 18 to 20, post-infective blood meal, mosquitoes were anesthetized on ice, rinsed in 70% ethanol, and washed in Dulbecco’s modified Eagle’s medium (DMEM), and the salivary glands were removed. Tissue was gently ground in a tissue homogenizer and centrifuged at 80 × g for 3 min to remove mosquito debris, and sporozoites were counted in a hemocytometer.

**Antibodies**

mAb 3D11 is directed against the repeat region of P. berghei CSP and, for some experiments, was conjugated to sepharose as described (Coppi et al., 2005). Polyclonal antisera to the NH2-terminal third of CSP was generated in rabbits as previously described and is used to detect full-length CSP (Coppi et al., 2005).

**Cells**

All cells were maintained in DMEM supplemented with 10% fetal calf serum and 1 mM glutamine (DMEM/FCS), except CHO cells, which were grown in Ham’s F-12 medium supplemented with 7.5% FCS, and HBMVEC cells, which were grown in EGM-2MV media (Cambrex,
Cell Host & Microbe

Heparan Sulfate Regulates Sporozoite Invasion

Rockland, ME) supplemented with 5% FCS. CHO cell mutant pgslE defective in GlcNAc-2-deacetylase/N-sulfotransferase 1 (Ndst1) was derived from wild-type CHO K1 cells and was described previously (Barne and Esko, 1989). Other cell lines were obtained from the following sources: Hepa 1-6 cells (CRL – 1830; ATCC, Rockville, MD); HBMVEC, a human endothelial cell line (CC-2811; Cambrex); mouse dermal fibroblasts were isolated from Balb/C mice as previously described (Freshney, 2000 and Supplemental Data).

Chlorate Treatment of Cells

Hepa 1-6 cells were seeded on glass coverslips (2.5 x 10^5/well) and grown overnight in low-sulfate medium (Ham’s F-12, 1 mM glutamine, and 2% FCS that had been dialyzed extensively versus 150 mM NaCl, 10 mM HEPES [pH 7.3]) with the indicated concentration of sodium chlorate. An appropriate amount of medium was replaced with water to maintain normal osmolarity. On the following day, cells were washed twice with DMEM not containing chlorate, and sporozoites were added in DMEM/FCS.

Metabolic Labeling

P. berghei sporozoites were metabolically labeled in DMEM with [35S]Cys/Met for 1 hr at 28°C. P. berghei in DMEM/FCS. To maintain normal osmolarity. On the following day, cells were washed and fixed, and sporozoites were stained with a double-staining assay that distinguishes intracellular from extracellular sporozoites as previously outlined (Coppi et al., 2005). For experiments in which development into EEFs was tested, sporozoites were added to the indicated cell line, and after 2 days, cells were fixed and stained, and EEFs were counted.

Dextran Migration Assay

Sporozoites (2 x 10^5) in DMEM with 0.2% BSA were added to monolayers of Hepa 1-6 cells in the presence of 1 mg/ml FITC-dextran (10,000 MW; Molecular Probes) for 1 hr at 37°C. Cells were washed with DMEM and fixed with 4% PFA, and the number of FITC-positive cells in 100 fields was counted. In some cases, sporozoites were preincubated with 10 μM E-64d or 1 mM cytochalasin D (CD) for 10 min at 28°C and added to cells in the continued presence of E-64d or CD. As a control for the effect of E-64d on wound healing, monolayers of Hepa 1-6 cells were preincubated ±10 μM E-64d for 15 min at 37°C. Following this, 1 mg/ml FITC-dextran was added to the cells, which were then wounded by scratching the coverslip five times with a single-edge razor blade. After 1 hr at 37°C in the continued presence of E-64d, cells were washed and fixed with 4% PFA, and the number of FITC-positive cells in 100 fields was counted.

Calcein Migration Assay

2.5 x 10^5 cells/well were seeded in 96-well tissue culture-treated plates in medium ± 20 mM chlorate and allowed to grow overnight until semiconfluent. On the day of the experiment, cells were washed with DMEM without phenol red and incubated in this medium with 10 μM calcein green-AM (Molecular Probes) for 1 hr at 37°C, then washed three times. Between 5 x 10^4 and 1 x 10^5 sporozoites in culture medium were centrifuged (300 x g at 14°C) onto the calcein-loaded cells and incubated for 1 hr at 37°C. Supernatants containing the released calcein were transferred to a ThermoElectron Microfluor 96-well plate, and fluorescence was read in a Labsystems Fluoroscan II using excitation and emission wavelengths of 485 nm and 538 nm, respectively. For experiments using staurosporine, W-7, KN-93, H-89, or purvalanol A, sporozoites were preincubated in the concentrations used above for 10 min at 28°C in DMEM and diluted 15-fold with the appropriate culture medium before being added to the calcein-loaded cells. Controls included addition of the same number of homogenized uninfected salivary glands and sporozoites preincubated with 10 μM E-64d or 1 μM CD for 10 min at 28°C and added to cells in the continued presence of these inhibitors. Unlike the dextran assay, which is a visual assay and therefore enables few events to be scored, the calcein assay requires larger numbers of sporozoites for signal detection.

Heparin and CSP Cleavage

Sporozoites were incubated in DMEM containing 0–100 μg/ml heparin for 5 min at 28°C, washed twice, fixed with 4% PFA, blocked with 1% BSA/PBS, and stained with polyclonal antiserum that recognizes full-length CSP followed by anti-rabbit Ig conjugated to FITC. One hundred to two hundred sporozoites were counted using phase and fluorescence microscopy, and the percentage of total sporozoites staining for full-length CSP was determined.

Disaccharide Analysis

Confluent cultures of each cell line were washed three times with PBS and lysed with 0.1 N NaOH for 15 min. An aliquot was removed for protein analysis, and the remaining lysate was neutralized with 10 N acetic acid and stored at −20°C until the analysis was performed. HS was isolated from 10^6 cells and purified as described for its isolation from tissues (Grobe et al., 2005; for details see Suppential Data). HS composition was determined by disaccharide analysis (for details see Supplemental Data), and molar percentages were calculated based on the relative area under each peak. These results were confirmed by a recently developed mass spectrometry-based technique (R.L., R. Cummings, and J.D.E., unpublished data). Results are presented as total disaccharides containing a particular modification.
**Construction of CDPK-6 Mutant Parasites**

CDPK-6 mutant *P. berghei* (ANKA) parasites were generated by double homologous recombination using the targeting vector pBS DHFR, in which the *T. gondii* dihydrofolate reductase/thymidylate synthase gene (*DHFR/TS*) is flanked by the upstream and downstream control elements from *P. berghei* DHFR/TS. This cassette confers resistance to pyrimethamine and was targeted to the CDPK-6 locus (PB01122.01.0) using CDPK-6 5′ and 3′ UTR sequences (for details see Supplemental Data). A second mutant parasite was made using the same region of homology except that the targeting construct was generated using a two-step PCR method as described in Ecker et al. (2006) (for details see Supplemental Data). Transgenic parasites were generated using the Amaxa Nucleofector (program U33) and 5 μg of DNA as previously outlined (Janse et al., 2006). Selection and cloning by limiting dilution for the generation of CDPK-6 mutant clones in which the selection cassette replaced the CDPK-6 coding sequence were carried out as previously described (Janse et al., 2006).

**Genotypic Analysis of CDPK-6 Mutant Parasites**

Cloned parasites were genotyped by pulse-field gel electrophoresis (PFGE) and Southern blot analysis. For PFGE, chromosomes were separated on an LKB 2015 PulsoPhor system using a linear ramp of 60–500 s for 72 hr at 4V/cm. The gel was blotted and hybridized with a probe that binds to the 3′UTR of DHFR/TS, which detects both the wild-type DHFR/TS locus on chromosome 7 and, in the case of the CDPK-6 mutant, the locus at which the transfected DNA integrated on chromosome 9. For Southern analysis, wild-type and genomic DNA from mutant parasites was digested with EcoRV, and the fragments were separated on a 0.8% agarose gel, blotted onto nylon membrane, and probed with a fragment of CDPK-6 3′UTR.

**Supplemental Data**

The Supplemental Data include Supplemental Experimental Procedures and four supplemental figures and can be found with this article online at [http://www.cellhostandmicrobe.com/cgi/content/full/2/5/316/DC1/](http://www.cellhostandmicrobe.com/cgi/content/full/2/5/316/DC1/).

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