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Parasitophorous vacuole poration precedes its rupture and rapid host erythrocyte cytoskeleton collapse in Plasmodium falciparum egress

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In the asexual blood stages of malarial infection, merozoites invade erythrocytes and replicate within a parasitophorous vacuole to form daughter cells that eventually exit (egress) by sequential rupture of the vacuole and erythrocyte membranes. The current model is that PKG, a malarial cGMP-dependent protein kinase, triggers egress, activating malarial proteases and other effectors. Using selective inhibitors of either PKG or cysteine proteases to separately inhibit the sequential steps in membrane perforation, combined with video microscopy, electron tomography, electron energy loss spectrosco py, and soft X-ray tomography of mature intracellular Plasmodium falciparum parasites, we resolve intermediate steps in egress. We show that the parasitophorous vacuole membrane (PVM) is permeabilized 10–30 min before its PKG-triggered breakdown into multilayered vesicles. Just before PVM breakdown, the host red cell undergoes an abrupt, dramatic shape change due to the sudden breakdown of the erythrocyte cytoskeleton, before permeabilization and eventual rupture of the erythrocyte membrane to release the parasites. In contrast to the previous view of PKG-triggered initiation of egress and a gradual dismantling of the host erythrocyte cytoskeleton over the course of schizont development, our findings identify an initial step in egress and show that host cell cytoskeleton breakdown is restricted to a narrow time window within the final stages of egress.

Significance

Malaria parasites develop within red blood cells inside a membrane-enclosed parasitophorous vacuole. An essential step in their life cycle is the exit of mature parasites from the blood cell, a multistage process termed egress. To do this, the parasites orchestrate a highly regulated sequence of membrane permeabilization and breakage steps culminating in the explosive release of parasites for a new round of infection. Here, we describe a previously unidentified permeabilization of the vacuolar membrane at the start of egress, preceding membrane rupture, suggesting a new initiation step in egress. We also show that, in the final minutes of egress, the blood cell membrane abruptly loses its structural rigidity and collapses around the parasites, showing a precise timing for cytoskeletal breakdown.

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SERA proteins is temporally associated with egress (20, 23–27), and at least one *P. falciparum* SERA family member, SERA6, is likely a cysteine protease that is activated by SUB1 (16). As well as PKG, a parasite calcium-dependent kinase called CDPK5 has been implicated in egress (28). However, in CDPK5-deficient parasites, SUB1 discharge and MSP1/6/7 and SERA processing are unaffected, suggesting that CDPK5 action is either independent from or downstream of the PKG/SUB1/SERA pathway (28).

Events downstream of SUB1 activation and MSP/SERA processing are less well understood. SUB1 has multiple substrates in addition to the SERAs and MSP1/67 (29), and other proteases have been proposed to play a role in egress, including the erythrocyte protease calpain-1 (30–35). Proteomic studies based on analysis of parasite populations normalized by physical techniques have indicated that erythrocyte membrane and cytoskeletal proteins are proteolyzed during or before egress (13, 36). It has been suggested on the basis of that work and atomic force microscopy (AFM) that the host cell cytoskeleton is progressively degraded over several hours of the ∼48 h-long erythrocytic life cycle of *P. falciparum* (13). However, in these studies, the developmental status (time from egress) of the individual parasitized cells examined was unknown. Other effector proteins involved in membrane breakdown are not fully characterized. In the related apicomplexan parasite *Toxoplasma gondii*, the pore-forming perforin-like protein 1 (PLP1) mediates rupture of vacuole and host cell membranes (37), whereas in *Plasmodium* the perforin-like proteins PLP2 and PLP1 have been implicated in egress of gametes (38, 39) and asexual blood forms, respectively (40).

Despite the above insights into the molecules that regulate egress, many details remain obscure concerning the timing, order, and nature of the membrane perturbations involved. The selective PKG inhibitors compound 1 (C1) and compound 2 (C2) reversibly inhibit egress before the rounding up stage (10, 14, 28, 41). In contrast, treatment with the broad-spectrum cysteine protease inhibitor E64 allows PVM rupture but selectively prevents erythrocyte membrane rupture, resulting in merozoites trapped in the blood cell after rupture of the PVM (30, 42). Here we use single cell tracking by video microscopy, electron tomography, and X-ray tomography of C1- or C2-treated or E64-treated *P. falciparum* schizonts to capture and discriminate intermediate stages in egress to an unprecedented degree of temporal accuracy. We show that although the PKG inhibitors prevent PVM rupture, there is a previously undetected, initial step in egress which permissively induced PVM rupture occurs 10–30 min before its complete rupture into multilayered vesicles. Just before or at the point of PVM rupture, the blood cell cytoskeleton undergoes a sudden breakdown causing the red blood cell membrane to collapse around the intracellular parasites. Finally, the blood cell membrane becomes permeable seconds before merozoite escape.

**Results**

**Electron Tomography of *P. falciparum* Schizonts Reveals the Fate of the PVM at Different Steps in Egress.** As asexual blood stages of *P. falciparum* invariably replicate asynchronously in vitro, resulting in “mixed” cultures that contain parasites at various stages of maturation, we enriched mature schizonts by centrifugation over cushions of Percoll. We then cultured these preparations in the absence of PKG inhibitors also to collapse around the intracellular parasite contents (Table S1). Analysis of the C1-treated schizonts showed that contrast equalization was evident in ∼50% of fully segmented parasites but was only observed in one example of a partially segmented schizont out of 80 analyzed. Importantly, examination of schizonts prepared in the absence of PKG inhibitors also revealed contrast equalization in 16% of segmented schizonts. These results suggested that contrast equalization occurs largely or entirely following merozoite segmentation and moreover confirmed that the observed contrast equalization before PVM rupture was not an artifact of C1 treatment.

In contrast to the PKG inhibitors, the broad-spectrum cysteine protease inhibitor E64 does not prevent PVM rupture but potently prevents final rupture of the erythrocyte membrane. Consistent with this, tomography of schizonts arrested by E64 showed a single membrane surrounding the parasites and loss of the PVM and blood cell contents (Fig. 1C). The presence of knob structures on the remaining enclosing membrane (Fig. 1C, Top) and staining under the membrane consistent with the presence of the erythrocyte cytoskeleton confirmed that this was the blood cell membrane. The empty appearance of the blood cell cytosol in the E64-arrested cells was consistent with previous descriptions of erythrocyte membrane poration preceding rupture (9, 40). In contrast to earlier parasite stages, the erythrocyte membrane shape was observed to closely follow the contours of the parasites in these cells (Fig. 1C). Also present were whorls of membrane vesicles, presumably the remains of the ruptured PVM. Tomography of the E64-treated schizonts enabled 3D visualization of the extensive, multilamellar nature of these vesicles (Fig. 1D).

**Time Course of Membrane Permeabilization and Breakage During Egress.** To monitor the vacuole and blood cell cytosol compartments during the course of egress in live schizonts, we used a transgenic *P. falciparum* parasite line (called 3D7_mCherryEXP1) that expresses mCherry fused to the secretory signal sequence of the PV protein EXP1. This targets the fluorescent fusion protein to the PV lumen. Examination of mature, C1-treated mCherryEXP1 schizonts revealed that although in some cells the fluorescence signal was restricted to the PV, in others it extended into the erythrocyte cytosol (Fig. 2A), suggesting permeability of the PVM to proteins of the molecular mass of the mCherryEXP1.
fusion (∼35.2 kDa) and consistent with the equalization of contrast observed by EM. These observations suggest that the PVM becomes permeable before activation of PKG, previously thought to be the first step in egress.

To further investigate the temporal relationship between PVM permeabilization and egress, mature mCherryEXP1 schizonts cultured for 3–4 h in the presence of C2 were washed to remove the inhibitor, allowing the cells to progress to egress. The schizonts were imaged during this process by time-lapse video microscopy to monitor changes in the distribution of fluorescence. Movies S1 and S2 and Fig. 2B and C show that PVM rupture and egress were always preceded by leakage of the fluorescence into the host cell cytosol. Of the cells that egressed during the time-lapse experiments, half were already leaky at the start or became leaky during recording. Of the cells that did not egress, only 18% were observed to be leaky during the recording. Furthermore, 73% of these leaky cells that did not rupture rounded up, which occurs a few minutes before egress (5), indicating that they would have likely have undergone egress shortly after the 30-min recording time. All of the cells that were not leaky at the start of the videos but egressed during the recording became leaky 10–30 min before final escape of parasites (tracking of 14 cells). Leakage occurred several minutes before rounding up and while the PVM still appeared intact (Fig. 2C and Movies S1 and S2). These observations confirmed the EM and EELS data and showed that permeabilization of the PVM occurs only 10–30 min before PVM rupture and egress.

Soft X-Ray Cryotomography and Scanning Electron Microscopy Reveal Dramatic Changes in Membranes and Cell Shape During Egress. To obtain further insights into the fate of the membranes in intact, C2-, or E64-treated cells without dehydration or sectioning, we used soft X-ray cryotomography, as the greater penetration depth of soft X-rays compared with electrons makes it possible to generate 3D reconstructions of intact, frozen-hydrated cells (reviewed in ref. 43). Although the resolution of this method is lower than for EM, the individual parasites and some of their organelles were clearly visible in the images (Fig. 3A–F). In the E64-arrested schizonts, most of the contents of the erythrocyte cytosol were absent, confirming the tomographic data (Figs. 1B and C and 3B and E). In addition, the 3D overview of whole cells provided by these data revealed striking differences in shape and mechanical properties between the C2-blocked and E64-arrested schizonts. In the former case, with the intact vacuole present, the erythrocyte membrane formed a globular enclosure around the vacuole with an extending remnant of the original biconcave shape (Fig. 3C). In

Fig. 1. Tomography of C1 and E64 stalled schizonts reveals the fate of the PVM at different stages in egress. (A) Schematic indicating the steps at which egress is stalled by the inhibitors C1/C2 and E64. (B) Life cycle stage-specific equalization of vacuole and erythrocyte contents in C1-treated schizonts. Tomogram slices from freeze-substituted sections of a partially segmented schizont showing a contrast difference across the PVM (Top) and a fully segmented schizont with an equalization of contrast across the PVM (Bottom). Each image is the average of 10 central slices from a tomogram and is shown with and without outlining of key features. Red, blood cell membrane; yellow, PVM; cyan, merozoite plasma membrane; green, merozoite nuclei; purple, apical organelles. (Scale bar, 500 nm.) (C) Slices from tomograms of freeze-substituted sections of schizonts arrested by E64. Each slice is shown with and without outlining of key features. Brown, digestive vacuole; other colors as above. In Top, an asterisk marks a knob structure on the erythrocyte membrane. (Scale bar, 500 nm.) (D) Slices in the xy, xz, and yz planes from tomograms of membrane whorls in E64 stalled schizonts, demonstrating that they are spherical multilamellar vesicles. Lines in the xy image show the positions of the other planes. (Scale bar, 100 nm.)
contrast, after PV breakdown in the E64-arrested schizonts, the erythrocyte membrane had collapsed around the parasites, following their contours (Fig. 3F). This collapse was also clearly present in the electron tomography sections (Fig. 1B and C) and was seen in optical microscopy as a compaction and rounding up of the schizont (Fig. 2C, compare 6- and 26-min time points). Scanning EM of these cells confirmed the change in shape and membrane collapse (Fig. 3G-J). In C2, 74% of the cells (790 cells examined) retained a vestige of the original biconcave disk shape, whereas in E64 48% (of a total of 759) showed the erythrocyte membrane collapsed around the parasites (Table S2). Because the rounding up and collapse occurred only a few minutes before final egress (Fig. 2C and Movies S1 and S2), these findings show that loss of mechanical integrity in the red cell membrane and cytoskeleton takes place only in the final stage of the egress pathway.

Discussion

We have used selective pharmacological inhibition of steps in egress coupled with tomographic and live cell imaging to show that the PVM becomes permeable before its lysis, allowing contents of the PV lumen and erythrocyte cytosol to mix. Importantly, the occurrence of PVM poration in C1-arrested cells shows that it occurs upstream of or independent of PKG signaling and SUB1 discharge into the PV. Our observation reveals a previously unidentified prior step in egress, in which the PVM is permeabilized before being disrupted. Intriguingly, our findings mirror those of Sturm et al., who noted gradual permeabilization of the PVM before its complete rupture in liver stage schizonts of the rodent malarial species *Plasmodium berghei* (44). This suggests similarities in the fate of the PVM in these otherwise very different developmental stages of the parasite life cycle.

The fact that the leaky PVM is visibly intact in tomograms and contains no breakages or disruptions visible by electron tomography suggests that permeabilization is not due to major membrane breakage but rather to smaller perturbations such as formation of localized pores. The mechanism underlying PVM poration is unknown. In the related apicomplexan *T. gondii*, the pore-forming perforin-like protein PLP1 plays an important role in egress from the host cell by disrupting both the vacuole and the host cell membrane (37). The evidence for an analogous role for a PLP1 ortholog in malarial egress is less clear. Although PLP1 has been indirectly implicated in poration of the erythrocyte membrane in *P. falciparum* (40), genetic knockout of either the PLP1 or PLP2 homolog in *P. berghei* produced no phenotype in the asexual blood stages (39, 45), casting doubts on the essentiality of PLP1 or PLP2 function in these developmental stages.

Live cell video microscopy revealed that, after PVM poration in blood-stage schizonts, there follows a 10–20-min period without further evident structural changes, followed by the sudden rounding up of the blood cell and increased merozoite motility (Movies S1 and S2). EM of E64-stalled schizonts showed a total breakdown of the PVM into multilamellar vesicles (Fig. 1D). EM and X-ray tomography of parasites arrested with E64 at the post-PVM rupture stage showed that the erythrocyte membrane then becomes permeable, indicated by the complete loss of the residual hemoglobin and PV contents and consistent with the findings of others (9, 40). In the live cell video experiments, we observed this loss of blood cell cytosolic contents in about one third of the schizonts just before parasite escape. Given the 5-s increment between frames in our time-lapse movies (a compromise between time resolution and photobleaching), this fraction of detected events is consistent with a previous report that the interval between erythrocyte membrane poration and its rupture is of the order of a few seconds (9).

The second key finding of this study is that a major change in erythrocyte structure occurs rapidly after release of the C2 block (Fig. 3), showing that the erythrocyte membrane in E64-blocked schizonts loses its rigidity and collapses around the parasites. The rounded cells appear smaller in diameter (Fig. 2C). This loss of cell shape and alteration in mechanical properties, both of which have been observed in optical and AFM studies (46), implies that the erythrocyte cytoskeleton undergoes a very sudden, rapid loss of its mechanical integrity just before egress. Importantly, the observation is not in accord with a previous suggestion that the host cell cytoskeleton is progressively dismantled over several hours during schizont maturation (13, 36). In those earlier studies, the timing of
cytoskeleton breakdown in individual cells could not be distinguished, as the analyses were of populations of cells in which egress was not blocked. Under these conditions, cells would gradually egress, accounting for the observed accumulation of cytoskeleton breakdown products. AFM analysis of schizonts from those populations showed cytoskeleton expansion but no loss of erythrocyte mechanical integrity, and does not provide the temporal resolution reported here, as the status of the individual parasites examined was unknown (13). The high degree of parasite synchrony afforded by the PKG inhibitors exploited in the present study, along with the much higher resolution and better cell preservation afforded by the microscopy methods used, gives us a much more accurate picture of these events. Our observations suggest that the effector mechanisms that degrade the host cell cytoskeleton to enable erythrocyte membrane rupture act very rapidly in the final stages of the egress pathway. A few seconds after cell collapse, the erythrocyte membrane breaks, allowing the final, explosive release of parasites. The fact that a similar, two-step breakdown (poration followed by rupture) occurs successively in the PVM and erythrocyte membrane raises the possibility that the effector molecules mediating poration of both membranes may be similar or identical.

In conclusion, our results suggest that poration of the PVM is the first detectable step in egress of the asexual blood-stage malaria parasite (Fig. 4). This occurs after parasite segmentation but before PKG-mediated SUB1 discharge. The mechanism underlying PVM poration is unknown, but as the leaky PVM appears intact, permeation is likely to arise from small, localized perforations such as membrane pores, potentially allowing small molecules such as calcium ions and parasite proteins into the residual erythrocyte where they can access the erythrocyte cytoskeleton and membrane. About 10 min after the first detection of PVM poration, the red blood cell rounds up, likely due to loss of mechanical integrity of the erythrocyte cytoskeleton mediated by protease activity. Around the same time, the PVM ruptures into multilamellar vesicles. In the following seconds, the blood cell membrane becomes permeable, rapidly followed by the explosive exit of the parasites. Identification of the effector molecules involved in PVM poration and cytoskeleton collapse may lead to new antimalarial therapies designed to interfere with progression of the malaria parasite blood stage life cycle.

Methods

Parasite Culture and Egress Inhibitor Treatment. P. falciparum clone 3D7 and parasite line 3D7_mCherryEXP1 (see SI Methods: for a description of generation of the latter) were cultured in human red blood cells in RPMI 1640 medium supplemented with Albumax (Invitrogen) as described previously (47, 48). Parasites were synchronized by centrifugation on cushions of 63% (wt/vol) Percoll (GE Healthcare) (47). Where inhibitor treatments were required, schizonts were cultured in medium containing either C1/C2 (2 μM) or E64 (50 μM) for 3–4 h before preparing the cells for imaging.

Live Cell Imaging. Video and fluorescence microscopy were carried out on a Nikon Eclipse Ni-E microscope equipped with a Plan Apo 100×1.45 oil immersion differential interference contrast (DIC) objective, a C11440 ORCA-Flash 4.0 digital CMOS camera (Hamamatsu), and a temperature-controlled stage held at 37 °C. Schizont suspensions in gassed medium were sealed into precooled viewing chambers constructed as described previously (14), immediately transferred to the microscope stage, and images and videos acquired using NIS Elements software (Nikon). Videos were started 5 min after the removal of the C1/C2 block and were acquired over 30 min at 5-s intervals. Images and videos were analyzed using Fiji (49).

Electron Tomography. Mature schizonts were pelleted by centrifugation, mixed with 20% (wt/vol) dextran in RPMI medium with C1/C2 (2 μM) or E64 (50 μM) where necessary and bakers yeast, and frozen using a HP/M10 high-pressure freezer (Baltec). Vitriﬁed cells were freeze-substituted into Lowicryl HM20 resin with uranyl acetate stain and cut into thin sections. Tilt series were collected on a Tecnai F20 200 keV field emission gun microscope (FEI) or a Tecnai T12 120 keV tungsten filament microscope (FEI), equipped with a US4000 CCD camera (Gatan). Dual axis tilt series were typically collected from –66° to +66° with an increment of 2° using the acquisition software SerialEM (50).

Fig. 4. Model of the stages of egress. The PVM undergoes poration before its rupture to form multilamellar vesicles. The cytoskeleton loses mechanical integrity and the erythrocyte membrane becomes porous before final parasite escape.
Soft X Ray Cryotomography. C2- and E64-treated schizonts were fixed in 2% (vol/vol) paraformaldehyde in PBS, added to electron microscopy grids coated with 0.1% (wt/vol) poly-L-lysine (Sigma) and 250 nm gold fiducial markers (BBI solutions), and plunge frozen in liquid ethane. Tilt series were collected on the UltraXRM-S/LS220C X-ray microscope (Zeiss, previously Xradia) at the B24 beamline of the Diamond synchrotron with a 1024 Pixis B CCD camera (Princeton instruments) and a 40 nm zone plate with X-rays of 500 eV. Tilt series were typically collected from −60° to +60° with an increment of 0.5°.

Tomographic Reconstructions. Electron and X-ray tomograms were reconstructed using etomo, part of the IMOD package (51). Postreconstruction, tomograms were denoised by nonlinear anisotropic diffusion filtering in IMOD. Selection of key features was done manually in IMOD.

Scanning EM. Schizonts arrested with either C2 or E64 were fixed in 2.5% glutaraldehyde, washed, osmicated (1% OsO4 for 16 h), dehydrated, critical point dried, and sputter coated with 5 nm gold. Images were collected on a JEOL 3M 7610F with 2.6 kV accelerating voltage.

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