High resolution microscopy reveals an unusual architecture of the *Plasmodium berghei* endoplasmic reticulum

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Summary

To fuel the tremendously fast replication of *Plasmodium* liver stage parasites, the endoplasmic reticulum (ER) must play a critical role as a major site of protein and lipid biosynthesis. In this study, we analysed the parasite’s ER morphology and function. Previous studies exploring the parasite ER have mainly focused on the blood stage. Visualizing the *Plasmodium berghei* ER during liver stage development, we found that the ER forms an interconnected network throughout the parasite with perinuclear and peripheral localizations. Surprisingly, we observed that the ER additionally generates huge accumulations. Using stimulated emission depletion microscopy and serial block-face scanning electron microscopy, we defined ER accumulations as intricate dense networks of ER tubules. We provide evidence that these accumulations are functional subdivisions of the parasite ER, presumably generated in response to elevated demands of the parasite, potentially consistent with ER stress. Compared to higher eukaryotes, *Plasmodium* parasites have a fundamentally reduced unfolded protein response machinery for reacting to ER stress. Accordingly, parasite development is greatly impaired when ER stress is applied. As parasites appear to be more sensitive to ER stress than are host cells, induction of ER stress could potentially be used for interference with parasite development.

Introduction

*Plasmodium* parasites are transmitted by the bite of an infected female *Anopheles* mosquito, which injects a moderate number of sporozoites into its host (about 100 in the case of *P. berghei*) (Frischknecht et al., 2004). About a third of the injected sporozoites find a blood vessel and travel through the bloodstream until they reach the liver. Here they invade and infect hepatocytes, thereby forming a parasitophorous vacuole membrane (PVM) (Bano et al., 2007). Within the parasitophorous vacuole (PV), parasites undergo extensive replication by schizogony; a single sporozoite can differentiate into up to 30,000 merozoites, depending on the species (Menard et al., 2013). In parallel with nuclear replication, the parasite has to expand its other vital organelles and ultimately provide sufficient membranous material to surround single daughter parasites (Sturm et al., 2009; Stanway et al., 2011). *Plasmodium* replication and growth, particularly in the liver, are very fast and resource-demanding processes. Despite the scavenging of host lipids and nutrients, the parasite must still produce enormous amounts of protein and lipid within minimal time (Mikolajczak et al., 2007; Blume et al., 2011; Slavic et al., 2011; Itoe et al., 2014). In addition to specialized organelles like micronemes, rhoptries and dense granules, *Plasmodium* possesses a rather classical set of secretory organelles, such as an ER and a single Golgi apparatus (Striepen et al., 2007). So far, description of their morphology and biogenesis is rudimental and restricted to *Plasmodium* blood stage parasites (van Dooren et al., 2005; Struck et al., 2005, 2008b).

The ER is an essential organelle in eukaryotic cells. It typically makes up almost half of the cell’s membranous material and is crucial for lipid biosynthesis, protein folding, calcium storage and protein secretion. Interestingly, the architecture of the ER has been recently described to have a greater influence on its function than previously anticipated (Westrate et al., 2015). ER sheets, or cisternae, are often found in a stacked conformation, coated with ribosomes, their main function being protein synthesis, folding, and post-translational modification.

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ER tubules are generally organized in dynamic, constantly rearranging and extending networks that function primarily in lipid biosynthesis. The ER can form direct contact with other membranes at membrane contact sites (MCS) (Phillips and Voeltz, 2015). In yeast such contacts occur, for example, with the mitochondrion, where they are important for lipid biosynthesis and calcium signalling (Kornmann and Walter, 2010).

Major components of membranes, such as phospholipids and proteins, are sorted within the ER and transferred to specialized regions called ER-Golgi intermediate compartments (ERGICs), which are in close proximity to the Golgi apparatus. Cargo from the ER is packed into COPII-coated vesicles and leaves the ER at distinct, long-lived transitional ER (tER) sites, also called ER exit sites (Szul and Sztul, 2011). It is proposed that secretory material first passes through the ERGIC before fusing with the cis-Golgi (Hammond and Glick, 2000). It has been suggested that P. falciparum parasites possess the machinery for COPII-coated vesicle transport as parasites encode components of this machinery and have distinct tER sites that are closely juxtaposed to the cis-Golgi, following the dynamics of the cis-Golgi (Struck et al., 2008b). The Plasmodium ER has been proposed to make contacts with the inner membrane complex (Hanssen et al., 2013), but whether the Plasmodium ER forms contacts with other organelles or the plasma membrane (PM), and whether proteins are transported through an ERGIC has not been described.

High demands for protein synthesis will cause overloading of the ER with unfolded and misfolded proteins, as space and chaperone numbers become restricted, which leads to an increased error rate in protein folding. If too many unfolded or misfolded proteins are present in the ER, an ER stress response known as the unfolded protein response (UPR) is mounted, which seeks to restore ER homeostasis (Wu et al., 2014). Apicomplexan parasites appear to harbour a dramatically reduced UPR machinery, with only Binding immunoglobulin Protein (BiP) and PERK homologues identified in the genome. If this is true, Plasmodium parasites would only be able to globally downregulate translation upon ER stress but would lack the transcriptionally mediated UPR (Gosline et al., 2011; Joyce et al., 2013) (Fig. 1).

In this study, we have analysed by microscopy the morphology and biogenesis of the ER in Plasmodium berghei parasites, with a focus on the liver stage, using transgenic parasite lines expressing fluorescently tagged marker proteins. We found that the Plasmodium ER forms extensive accumulations during asexual replication. Utilizing serial block-face scanning electron microscopy (SBFSEM), we were able to define these accumulations as voluminous, intricate and dense networks of ER tubules. We hypothesize that these accumulations might allow the parasite ER to cope with the enormous protein throughput, and hence ER stress, to enable fast and massive parasite replication. Additionally, we found evidence that the parasite ER forms contact sites...
with the PM or PVM and we speculate that these contact sites could allow nonvesicular lipid transfer to feed lipids into these rapidly expanding membranes.

Results

The endoplasmic reticulum of P. berghei liver stage parasites has an unusual architecture

To visualize the parasite ER, we N-terminally tagged the P. berghei homologue of a well described ER marker protein, PbSec61\(\beta\), with both GFP and superfolder GFP (sfGFP) (Couffin et al., 1998; Pédelaqc et al., 2006; Kornmann et al., 2011). The sfGFP offers several advantages over GFP for the tagging of ER transmembrane proteins such as Sec61\(\beta\). GFP has the tendency to weakly dimerize. When this occurs at the cytoplasmic face of the ER membrane, it can induce artificial reorganization of the entire ER architecture, resulting in huge ER patches known as organized smooth ER (Snapp et al., 2003). In contrast, sfGFP is a purely monomeric version of GFP that was designed to exhibit improved folding capacities and to be more resistant to denaturation (Costantini and Snapp, 2013). We did not observe differences in the localization of the two fusion proteins and thus used both for our experiments. Transgenic parasites were generated, expressing either a GFP-PbSec61\(\beta\) or a sfGFP-PbSec61\(\beta\) fusion protein under the control of the constitutive eef1\(\alpha\) promoter, which allowed visualization of the ER throughout the parasite’s life cycle (Fig. 2A). We focused on analysing the parasite ER during liver stage development, but for comparison also imaged other life cycle stages (Supporting Information Figs S1 and S2). Live cell imaging of Pb\(^2\)sfGFP-Sec61\(\beta\) parasites showed that during liver stage schizogony, the parasite ER closely surrounds each parasite nucleus and additionally builds a network of ER membranes that extend through the entire cytoplasm (Fig. 2B and Supporting Information Movie S1). The parasite ER seems to remain as a single interconnected organelle until very shortly before merozoite formation. In merozoites, only the perinuclear ER is present (Fig. 2B and Supporting Information Movie S1). To verify this observation, Pb\(^2\)GFP-Sec61\(\beta\) parasites were fixed towards the end of liver stage development, at 56 h post infection (hpi), and stained with an antisera against PbMSP1 (merozoite surface protein 1, PBANKA.083100), which localizes to the parasite PM of late liver stage parasites. We were able to confirm by visualization of single daughter merozoites, defined by their surrounding PbMSP1-positive PM, that the ER is found mainly around the nucleus with occasional minor protrusions, but is not peripherally localized (Fig. 2C).

Surprisingly, we observed that the parasite ER, in addition to being perinuclear and existing as network-like structures, also forms extensive ER accumulations, which start to appear in trophozoites and become most prominent during the cytomere stage (Fig. 2B, red arrows). Measuring different focal planes of Pb\(^2\)sfGFP-Sec61\(\beta\) parasites (\(n = 3, 9\) parasites per replicate; 5 random focal planes measured per parasite), these ER accumulations were estimated to account for about 24% of the total parasite area during the cytomere stage (Supporting Information Fig. S3). Interestingly, most of the ER accumulations disappear during the formation of individual merozoites and accumulations are completely absent at the end of liver stage development when merozoites have formed. To exclude that these ER accumulations are an artefact of GFP localization to the cytoplasmic face of ER membranes, we generated a specific antiserum against PbBiP, an ER chaperone localized in the ER lumen (Kumar et al., 1991; Snapp et al., 2003). We used this anti-PbBiP antiserum to stain HeLa cells infected with wild-type (WT) parasites (Fig. 3A) and Pb\(^2\)sfGFP-Sec61\(\beta\) parasites (Fig. 3B), both fixed at 48 hpi, which approximates to the cytomere stage. We observed ER accumulations in anti-PbBiP-stained WT parasites, which confirmed that ER accumulations occur naturally during parasite development. Co-localization of the anti-PbBiP staining with the sfGFP-Sec61\(\beta\) was examined by determining the Pearson’s \(R\) value, which is 0.93, implying a high degree of co-localization. Importantly, we were also able to show by intravital microscopy that parasite ER accumulations are present in vivo in Pb\(^2\)sfGFP-Sec61\(\beta\) parasites at 42 hpi, and had disappeared in most parasites imaged at 49 hpi when merozoites had formed (Fig. 3C). This demonstrates that accumulations of parasite ER are indeed a natural feature of P. berghei development. We also found these ER accumulations in other phases of parasite replication (Supporting Information Figs S1 and S2). During blood stage schizogony, we observed ER accumulations from the ring stage onwards until they disappear in maturing schizonts. Blood stage merozoites seem to contain only the perinuclear ER, similar to liver stage merozoites. Interestingly, we also observed ER accumulations in gametocytes (Supporting Information Fig. S1). ER accumulations were also found during sporogony of the parasite in the mosquito midgut, but are absent in mature sporoblasts. Midgut and salivary gland sporozoites possess a perinuclear ER and a peripheral ER network, which spans through the entire sporozoite, but are devoid of ER accumulations (Supporting Information Figs S2 and S4). Together, it appears that ER accumulations are formed in proliferative parasite stages and are absent in non-proliferative stages.

High resolution microscopy offers insight into the nature of ER accumulations

When we examined ER accumulations using standard widefield fluorescence microscopy (FM), they appeared as
oval to round ER bodies, potentially formed through expansion of ER sheets or tubules (Fig. 4A). To unravel the architecture of ER accumulations in detail, we used stimulated emission depletion (STED) microscopy, which typically gives a resolution of 30–80 nm (Hell and Wichman, 1994). STED imaging of WT parasites fixed at 48 hpi and stained with anti-PbBiP antiserum, revealed that ER accumulations have a dense membranous tubular organization (Fig. 4B). In order to resolve the structure of ER accumulations even further, we used electron microscopy (EM). Recently, there have been major advances in 3D EM using SBFSEM, which has been already used to look at morphological and volumetric changes of parasites in different blood stages of *P. falciparum* (Sakaguchi et al., 2016). We used SBFSEM for imaging of FACS-sorted PbRmCherry parasites that were fixed at 48 hpi. By vertically cutting thin slices through our sample and subsequent imaging we
were able to identify parasites containing ER accumulations. SBFSEM imaging confirmed that the ER forms an intricate dense and seemingly interconnected tubular network, which most often has a round to oval shape (Fig. 4C-a and 4C-a', Supporting Information Fig. S5 and Movie S2 and S3). Using transmission EM (TEM) we were able to obtain an even higher resolution of the tubular network and to resolve ER membranes (Fig. 4C-b, b', b''). We found that ER tubules within the network have a mean diameter of $68 \pm 6$ nm (95% confidence interval), which is consistent with the typical ER diameter of 50–100 nm found in most eukaryotic organisms. (Voeltz et al., 2002).

We measured the area of ER accumulations at their greatest width in SBFSEM images and compared this with standard FM of parasites, for which infections had been performed identically up to and including parasite fixation. We found that ER accumulations imaged with either SBFSEM or FM have a similar mean area of about 6.3 $\mu m^2$ and we, therefore, conclude that structures found by SBFSEM and FM are the same (Fig. 4D).

**FLIP analysis of the parasite ER**

Having established that the ER consists of vast accumulations in addition to a tubular network and the perinuclear ER, we sought to investigate whether or not these are connected. To get a better understanding of the ER architecture, we performed fluorescence loss in photobleaching (FLIP) experiments in which we repeatedly photobleached a 3.75 $\mu m^2$ area of PbCgfGFP-Sec61$\beta$ parasites and measured fluorescence intensity in the remainder of the parasite. If the GFP-tagged protein can diffuse freely throughout the entire parasite ER, we would expect that the fluorescence intensity of the entire ER would rapidly decrease (Teixeira and Huston, 2008). To control for
fluorescence loss by repeated imaging we imaged a separate set of parasites where the bleaching area was set outside the parasite. When we analysed parasites at 40 hpi using FLIP we observed an almost complete loss of fluorescence in the entire parasite ER after 20 cycles of photobleaching. This loss is specific as control parasites show only a minor decrease in fluorescence intensity (Fig. 5A and Supporting Information Movies S4 and S5). These results imply that the GFP-Sec61β fusion protein can diffuse freely throughout what must be a single interconnected ER compartment despite its various forms. We had noticed that the ER seems to be divided shortly before single merozoites are formed and we assumed that the ER has lost its connection through the whole parasite at this point (Fig. 2C). When we photobleached a 3.75 μm² area of the parasite ER at 56 hpi, at which point only perinuclear ER is seen, only minor levels of fluorescence intensity are lost in the remaining parasite, meaning that at this stage, the ER has mostly lost its interconnectivity (Fig. 5B and Supporting Information Movies S6 and S7) confirming our observation made by fluorescence microscopy (Fig. 2C).

**Effect of ER stress on ER accumulations in *P. berghei* liver stage parasites**

*Plasmodium* parasites appear to be fairly limited in the machinery they harbour to react to and cope with ER stress (Gosline *et al.*, 2011; Harbut *et al.*, 2012) (Fig. 1). We reasoned that the parasite might, therefore, be much more sensitive to inhibitors that induce ER stress and that inhibitor treatment might have an influence on
abundance and size of ER accumulations, as ER extension is known to alleviate ER stress (Schuck et al., 2009). Infected cells were treated with inhibitor concentrations that did not affect host cell development, and their effect on parasite development was investigated (Fig. 6 and Supporting Information Fig. S6). ER stress was induced using the well-described proteasome inhibitor MG132 or using two different signal peptide peptidase (SPP) inhibitors, namely (ZLL)2 and Ly411575. MG132 selectively blocks the proteolytic activity of the 26S proteasome, which is known to be present in Plasmodium parasites, leading to the accumulation of ubiquitinated proteins and thus ER stress (Aminake et al., 2012; Prasad et al., 2013). (ZLL)2 and Ly411575 have been described to specifically inhibit the activity of Plasmodium SPP (Parvanova et al., 2009; Harbut et al., 2012). SPP is an ER-resident protein, which is involved in the removal of signal peptides. In mammalian cells, an additional role has been found in transport of proteins destined for proteasomal degradation from the ER lumen to the cytoplasm (Loureiro et al., 2006). We infected HeLa cells with PbGFP parasites (PbGFP<sub>CON</sub>, constitutive cytoplasmic GFP expression (Franke-Fayard et al., 2004)) and initiated inhibitor treatment at 24 hpi (MG132 at 50 nM, (ZLL)<sub>2</sub> at 2 μM and Ly411575 at 20 nM). Applied inhibitors did not interfere with host cell growth and proliferation at concentrations used (Supporting Information Fig. S6). At 48 hpi, we visualized the ER with ER-Tracker<sup>TM</sup> Red, which gives an ER staining that correlates to the previously observed ER localization (Supporting Information Fig. S7A). A confocal spinning disc microscope was used to assess the size and number of ER accumulations in relation to the parasite size of inhibitor-treated parasites compared to control parasites (Fig. 6A). Measured parasites had a comparable thickness along the z-dimension (Supporting Information Fig. S7B). Parasites treated with Ly411575 were not assessed for size and number of ER accumulations, since we observed that parasite development was dramatically impaired; parasites stayed very small and did not form any ER accumulations. For untreated parasites and those treated with inhibitors other than Ly411575, we first analysed the mean area of ER accumulations to investigate whether inhibitor treatment influenced the morphology of single ER accumulations independent of the parasite size (Fig. 6B). Next, we investigated if inhibitor-treated parasites harboured

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**Fig. 5.** FLIP imaging of the *P. berghei* ER shows that the parasite ER is a single continuous organelle, which is divided once merozoites are formed. HeLa cells were infected with PbGFP-Sec61β parasites and imaged live at either 40 hpi A. or 56 hpi B. Parasites were photobleached 20 times at 10 s intervals at the indicated area. Fluorescence intensity was measured at the ROI. In control parasites, the bleaching area was set outside of the parasite to account for loss of fluorescence intensity by imaging. Images of representative parasites are shown. Fluorescence intensity values of the prebleached parasites were set as 100%. Depicted values are means, error is depicted as 95% confidence intervals (grey area above and under the curve) of the relative fluorescence intensity at each time point, n = 3, with each eight imaged parasites per settings.
different numbers of ER accumulations. As parasites can vary considerably in size, we analysed the total parasite area per ER accumulation (Fig. 6C). Finally, we analysed the total extent of ER accumulations, represented by the percentage of parasite cytoplasmic area occupied by ER accumulations (Fig. 6D). When we treated parasites with MG132, we observed that they form significantly larger ER accumulations, but seem to have slightly (nonsignificantly) reduced numbers of ER accumulations per parasite relative to the parasite area. Nevertheless, the total parasite area that is occupied by ER accumulations seems to be similar in control parasites (Fig. 6B–D). Parasites treated with (ZLL)2 form ER accumulations that are relatively similar in size and number compared to those in control parasites (Fig. 6B–D). In addition to the abundance and size of ER accumulations, we also analysed the effect of inhibitor treatment on parasite size and on completion of liver stage development. Interestingly, we found that parasites treated with ER stress-inducing inhibitors were generally smaller at 48 hpi. This was most significant upon treatment with Ly411575, where parasite size was only half that of control parasites (Fig. 6E). Furthermore, all inhibitor-treated parasites were severely impaired in successfully completing liver stage development, illustrated by a more than 50% reduction in the generation of detached cells compared to control parasites (Fig. 6F).
Markers of the tER, cis- and trans-Golgi are localized at ER accumulations

To further characterize the role of parasite ER accumulations, we were interested in understanding whether these accumulations are sites of protein synthesis and subsequent transport. For initial characterization of the *P. berghei* ER we used transgenic parasites expressing a (sf)GFP-PbSec61β fusion protein (Fig. 2A–C). Sec61β, together with Sec61α and Sec61γ, builds the Sec61 complex, which forms a channel through the ER membrane and is central to co- and post-translational protein translocation. Membrane-bound ribosomes are known to be tightly associated with the Sec61 complex to facilitate co-translational protein transport (Kelkar and Dobberstein, 2009).

Successfully folded proteins exit the ER at ER subdomains, so-called tER sites, to then be transported to the cis-Golgi for further modification and sorting. Sec13p is a component of COPII-coated vesicles that has been used previously in *P. falciparum* to visualize the tER (Struck et al., 2008b). Here, we generated transgenic parasites expressing the *P. berghei* homologue of Sec13p (PBANKA_144540) as a GFP fusion protein. PbSec13p-GFP-infected HeLa cells were fixed between 24 and 56 hpi and stained with anti-PbBiP antiserum to allow visualization of the ER with tER sites at mid to late liver stage development (Fig. 7). PbSec13p is localized in distinct foci, which we found to be distributed along the parasite ER during the *Plasmodium* liver stage. Interestingly, tER foci are also present at ER accumulations, seemingly concentrated at their periphery. Generally, tER foci appear to be more abundant at ER accumulations than at other sites of the ER. Merozoites of mature schizonts each harbour a single tER site, localized at a distinct region of the parasite perinuclear ER (Fig. 7).

We were next interested in how the Golgi is positioned in relation to the ER. In the cells of most organisms, the cis-Golgi is found juxtaposed to the tER, to allow effective COPII-mediated protein transport between the ER and Golgi apparatus. As *Plasmodium* parasites possess a non-classical unstacked Golgi, the trans-face of the Golgi is found in close proximity to the cis-Golgi. The Golgi reassembling stacking protein (GRASP) and Rab6 are well-established marker proteins of the cis-Golgi and trans-
Golgi, respectively, and have been used in *P. falciparum* for visualization of both Golgi faces in the parasite’s blood stage (Struck et al., 2008a,b). To visualize the cis-Golgi and trans-Golgi, we generated PbLSGRASP-GFP and PbLSGFP-Rab6 parasites (Supporting Information Figs S8, S9 and S12). PbLSGRASP-GFP parasites were fixed at different time points of liver stage development and the ER was stained with anti-PbBIP antiserum. cis-Golgi faces are found at distinct ER sites, with the cis-Golgi being mostly found at ER accumulations (Supporting Information Fig. S8). To verify these observations, we additionally generated double fluorescent PbCsfGFP-Sec61β-PbLSGRASP-mCherry parasites. Confocal live cell imaging of these double-fluorescent parasites confirmed that cis-Golgi faces are indeed mainly found at ER accumulations (Fig. 8 and Supporting Information Movies S8 and S9). Interestingly, when the PM starts to invaginate at the cytomere stage, cis-Golgi structures are found underlying the PM and become absent from ER accumulations. Following this, each forming daughter merozoite was observed to inherit one cis-Golgi face, which is then localized at a distinct site of the perinuclear ER (Fig. 8). For analysis of the trans-Golgi localization, PbLSGFP-Rab6 parasites were stained with ER-Tracker™ Red and imaged live. Interestingly, we found that trans-Golgi faces are also localized at ER accumulations but are also frequently located at other regions of the ER (Supporting Information Fig. S9). Finally, when single merozoites are formed, they contain a single cis- and trans-Golgi face (Supporting Information Figs S8 and S9). In summary, we find that ER and cis- and trans-Golgi compartments are localized at ER accumulations and hence we suggest that ER accumulations might be hubs of protein export from the ER to the Golgi apparatus.

**The ER forms extensions towards the parasite plasma membrane and parasitophorous vacuole membrane**

Visualizing the parasite ER by confocal imaging of sfGFP-tagged Sec61β and ER-Tracker™ Red and by EM, we noticed that the ER not only forms a huge network throughout the parasite but also seemingly extends towards the parasite PM and even to the PVM (Fig. 9A). To further investigate this observation, HeLa cells were infected with PbLSExp1-GFP parasites, which express the PVM-resident protein PbExp1 (*P. berghei* exported protein 1), fused to GFP. PbLSExp1-GFP parasites were found to develop similarly to PbLSExp1-mCherry parasites (Graewe et al., 2011). At 48 hpi parasites were stained with ER-Tracker™ Red and imaged using a confocal spinning disc microscope. We observed that the parasite ER forms long tubular extensions towards the PVM, often originating from node-like ER structures close to the PM (Fig. 9A, red arrows). Tubular extensions are different ER structures than the previously described ER accumulations. Interestingly, we found that the PVM is also stained by ER-Tracker™, which is a drug conjugate glibenclamide BODIPY® TR (Fig. 9A). Glibenclamides are known to bind selectively to sulphonylurea receptors of ATP-sensitive K⁺ channels, prominent on the ER (Hambrock et al., 2002). Due to the staining of the PVM with ER-Tracker™, we hypothesized that the ER potentially feeds lipids into the PVM. Interestingly, we found that staining of the PVM by ER-Tracker™ is most obvious in regions where the PbExp1-GFP signal is weak or absent (Fig. 9A, white arrows). Seeking to characterize the spatiotemporal presence of ER extensions directed towards the parasite periphery we imaged PbLSExp1-GFP, PbWT and PbLGFP parasites stained with ER-Tracker™ Red, live at different time points throughout liver stage development. We found ER extensions in all imaged parasites at each time point investigated between 24 hpi and 56 hpi (*n* > 3, with more than 15 parasites imaged per replicate and time point). Data of PbLSExp1-GFP and PbWT are not shown, examples of imaged PbLGFP parasites are provided in Supporting Information Fig. S10A. Live imaging of transgenic double fluorescent PbCsfGFP-Sec61β-PbLSExp1-mCherry parasites allowed detailed analysis of parasite ER and PVM dynamics without potential artefacts through live stains or fixation. We imaged parasites during liver stage development (40, 48 and 56 hpi) using a confocal microscope and were able to confirm the presence of ER extensions towards the PVM in all imaged parasites (*n* = 3 per time point, with each 20 imaged parasites per replicate) (Fig. 9B upper panel, arrows and Supporting Information Fig. S10B). PbCsfGFP-Sec61β-PbLSExp1-mCherry parasites, fixed at 48 hpi were imaged at a higher resolution using STED microscopy, which indicates that tubules of the parasite ER seemingly interface with the PVM and/or PM to establish potential ER-PM/PVM contact sites (Fig. 9B lower panel, arrows). In addition to our observations made using confocal microscopy, we also detected potential connections between the ER and the parasite PM and/or PVM by SBFSEM, which again seem to originate from a node-like ER structure close to the PM. Thus, EM data strengthens our observation of potential contact sites between the ER and the parasite PM and/or PVM (Fig. 9C).

**Discussion**

Here, we report for the first time a detailed description of highly concentrated ER accumulations that are evenly distributed within and formed during development of *Plasmodium* parasites, additionally to an interconnected perinuclear and network-like parasite ER. A combination of high-resolution STED microscopy and SBFSEM allowed us to define ER
Fig. 8. The cis-Golgi (red), defined by PbGRASP-mCherry, is localized at distinct regions of the parasite ER (green). HeLa cells were infected with Pb^3sfGFP-Sec61β-Pb^3GRASP-mCherry. Cells were imaged live with a confocal microscope at different time points of parasite liver stage development between 36 and 56 hpi. Between the trophozoite and cytomere stage, cis-Golgi structures are mainly present at ER accumulations. Upon parasite PM invagination around groups of nuclei the cis-Golgi is found at the parasite periphery and absent from ER accumulations. Merozoites each contain one cis-Golgi face. Red boxes are shown at a higher magnification. Scale bars correspond to 10 μm.
accumulations as an intricate network of densely packed ER tubules.

Even though the *Plasmodium* ER has been the focus of numerous studies, ER accumulations described in the present study have never been mentioned. Couffin *et al.* showed that *P. falciparum* Sec61 is localized at the parasite ER in blood stage parasites using antibody staining. Although FM images cannot be interpreted conclusively and blood stage parasites are much smaller than liver stage parasites, one can already identify ER accumulations in their staining (Couffin *et al.*, 1998). Another study characterized ER morphology in *P. falciparum* blood stage parasites expressing GFP with an ER retention signal -SDEL. They describe the parasite ER,
similarly to our observations, as a ring around the nucleus that forms extensions into the cytoplasm, which become considerably branched in late stages. However, ER accumulations, clearly present in their images of transgenic trophozoites and schizont parasites are not mentioned in the text (van Dooren et al., 2005). Similarly, other studies investigating vesicle-mediated trafficking, PfSec22, PfSec24 and PfSec12 of the P. falciparum blood stage also show images of the parasite ER with visible ER accumulations, but these were not made reference to (Adisa et al., 2007; Lee et al., 2008; Ayong et al., 2009). Additionally, a recent study using P. yoelii liver stage parasites found the lysophosphatidic acid acyltransferase (LPAAT) PY01678 to be localized at the parasite ER. Staining of the P. yoelii ER with anti-BiP in this study clearly shows the presence of numerous ER accumulations in P. yoelii liver stage parasites, which were again not discussed (Lindner et al., 2014).

The ER size and shape is highly dependent on the function and requirements of the cell. For example, the ER of B lymphocytes triples its size during differentiation into highly secreting plasma cells, which are clearly in need of a greater protein folding capacity (Van Anken et al., 2003). The UPR has been shown to play an essential role in the expansion of ER membranes. Under ER stress, the transcription factor XBP1 is activated by IRE1, leading to the elevation of the chaperone level in the ER. Additionally, UPR activates the associated ERAD and lipid biosynthesis pathway (Cox et al., 1997). Interestingly, XBP1-deficient B lymphocytes are not able to differentiate into plasma cells as they fail to expand their ER (Reimold et al., 2001). Interestingly, yeast cells deficient in either IRE1 or Hac1 (XBP1 in metazoans) were not able to normally expand their ER membrane by generation of ER sheets but instead expanded their ER through formation of ER patches under ER stress (Schuck et al., 2009). Using EM, they were identified as abnormally shaped, intertwined ER tubules. Surprisingly, even though ER expansion in UPR-deficient yeast cells was based on tubular expansion and was uncoupled from increasing amounts of chaperones, ER expansion alone, independent of its shape, was sufficient to alleviate ER stress (Schuck et al., 2009). ER patches identified in yeast are quite similar in their organization to ER accumulations we observed during Plasmodium asexual development.

Bearing in mind that Plasmodium parasites harbour only a dramatically reduced UPR machinery, deficient in IRE1, AFT6 and their downstream components and are thus limited in their response to ER stress (Fig. 1), ER accumulations might indeed be generated to alleviate the ER stress caused by the enormous growth rate of the parasite (Gosline et al., 2011). Upon inhibitor treatment, we observed only a tendency for the total parasite area occupied by ER accumulations to be slightly larger in inhibitor-treated parasites than in control parasites. A possible reason we do not see more significant effects on the size and number of ER accumulations, when ER stress is induced with inhibitors, is that parasites might already have an intrinsically increased level of ER stress, defined by ER expansion, as a result of the tremendous parasite growth rate. As a consequence, inhibitor-induced ER stress might shift the ER stress response from seeking to restore ER homeostasis towards a cell death response. This hypothesis is supported by the fact that we observed a clear reduction in the number of successfully developing parasites and in parasite size, upon treatment with ER stress-inducing inhibitors. The fact that Plasmodium parasites harbour a vastly reduced UPR machinery and the elevated sensitivity of parasites to ER stress, illustrate the potential of interfering with parasite development by targeting components of the parasite UPR.

Using classical marker proteins of the tER, cis- and trans-Golgi, we have examined the relationship between ER accumulations and the immediate secretory system. We found that foci of ER exit sites and Golgi cisternae are concentrated at ER accumulations. Accordingly, we assume that ER accumulations are functional divisions of the parasite ER, involved in protein and lipid dispatch.

Once parasites have finished nuclear division and start to build merozoites, we noticed by in vitro and intravitral live cell imaging that ER accumulations disappear. It is believed that the parasite PM needed to encase forming merozoites originates from invaginations of the surrounding PM, primarily around spheres of nuclei, and finally around single merozoites. Invagination of the PM to form single merozoites is restricted to a short period of time spanning only a few hours at the end of parasite development (Sturm et al., 2009; Stanyo et al., 2011). Little is known about how the parasite is able to provide sufficient amounts of membrane to surround thousands of merozoites in such a short time frame. In Plasmodium parasites, membrane reservoirs are not found directly at the PM but rather at the surrounding PVM, which forms extensive protrusions into the parasite host cell during liver and blood stage development (Deschermeier et al., 2012; Grützke et al., 2014; Prado et al., 2015; Matz et al., 2015). The PM and PVM exhibit regions of close alignment, which would theoretically allow for a direct exchange ofmembranous material between the two membranes. However, as the parasite matures and especially when the PM starts to invaginate, the distance between PM and PVM increases (Graewe et al., 2011), hindering potential membrane transfer. We have observed by live cell imaging and EM that the parasite ER extends tubules from node-like ER patches towards the parasite PVM.
and/or PM, appearing to build up membrane contact sites. Based on these findings, we hypothesize that lipids might be transferred directly via ER-PM contact sites for allow for PM expansion and invagination. In other eukaryotes, ER-PM contact sites are involved in many cellular processes, such as nonvesicular lipid transport, store-operated Ca$^{2+}$ entry and cell signalling (Henne et al., 2015). Since we could never observe direct transport of proteins, such as sfGFP-PbSec61β, through ER-PM contact sites, we propose that membrane lipids might be transported via non-vesicular lipid transport from the ER to the PM rather than through membrane fusion or vesicular transport.

Here, we provide first observations of potential contact sites between the parasite ER and PM, yet further investigations are needed to analyse whether lipids are transferred via MCSs and if potential transfer leads to the depletion of ER accumulations and helps membrane expansion.

In summary, the use of high-end microscopy techniques has enabled us to analyse the morphology of the P. berghei ER in detail and to provide explanations regarding the origin and function of ER accumulations identified.

**Experimental procedures**

**Animal work statement**

Experiments were conducted with strict accordance to the guidelines of the Swiss Tierschutzgesetz (TSchG; Animal Rights Laws) and approved by the ethical committee of the University of Bern (Permit Number: BE109/13).

C57BL/6 and BALB/c mice used in experiments were between 6 and 10 weeks of age and were either bred in the central animal facility of the University of Bern, or were supplied from Harlan Laboratories or Charles River.

**Culture and in vitro infection of HeLa cells**

In this study, we used HeLa cells, which in contrast to HepG2 cells, grow only in a single layer. On the one hand, this allows sporozoites to infect host cells more easily, which results in a higher infection rate and on the other hand facilitates imaging of subcellular structures of P. berghei liver stages in vitro. Development of parasites in HeLa cells and HepG2 cells to infectious merozoites is comparable. It has been previously demonstrated that P. berghei parasites are able to successfully develop in HeLa cells (Calvo-Calle et al., 1994). HeLa cells (generous gift from Robert Menard, Pasteur Institute, Paris) were grown in MEM (minimum essential medium) with Earle’s salts, supplemented with 10% heat inactivated FCS (foetal calf serum), 1% penicillin/streptomycin and 1% L-glutamine (PAA Laboratories) in a humid incubator at 37°C with 5% CO$_2$. Cells were passaged twice a week using accutase. For infection, either 3 x 10$^5$ cells were seeded onto glass cover slips contained separately in wells of a 24-well cell culture plate, or 1 x 10$^5$ cells seeded into glass-bottom dishes. About 24 h post seeding, HeLa cells were infected with sporozoites prepared from salivary glands of female Anopheles stephensi mosquitoes infected with the aforementioned P. berghei parasite lines. Sporozoites contained in infection medium [MEM with 2.5 μg ml$^{-1}$ Amphotericin B (PAA Laboratories)] were incubated with HeLa cells for 2 h and following this, the sporozoite-containing medium was removed and new infection medium was added. Afterwards medium was changed daily.

**Inhibitor treatment and ER-Tracker™ staining**

Inhibitors were dissolved in DMSO for working stocks (MG-132 (Sigma-Aldrich) at 250 μM and ZLL$_2$ (Calbiochem) at 10 mM and Ly411575 (Sigma-Aldrich) at 100 μM). HeLa cells were infected with sporozoites of PbC6GFP parasites, inhibitors (MG132 at 50 nM, ZLL$_2$ at 2 μM and Ly411575 at 20 nM) were added initially at 24 hpi and exchanged at 48 hpi. For microscopic evaluation of ER accumulations at 48 hpi, the ER was stained using ER-Tracker™ Red (glibenclamide BODIPY® TR). Infected HeLa cells were incubated with 500 nM ER-Tracker™ Red for 1 h at 37°C and 5% CO$_2$. As a next step, the staining solution was replaced with probe-free infection medium and parasites were imaged no later than 45 min after staining.

**Detached cell assay and size measurement**

For quantification of parasites that are able to successfully complete liver stage development, HeLa cells were infected as described above. Parasite numbers were counted at 48 hpi and compared to the number of successfully developed parasites (detached cells) that were counted in the supernatant at 65 hpi. The parasite area in μm$^2$ was measured at 48 hpi by density slicing using Fiji software. Counting and imaging for the size measurement was performed using a Leica DMi6000B widefield epifluorescence microscope.

**Live cell imaging and FLIP analysis**

For live cell imaging, 1 x 10$^5$ HeLa cells were seeded into glass bottom dishes (MatTek) and infected as described, DNA was stained with a final concentration of 1 μg ml$^{-1}$ Hoechst 33342 (Sigma) and cells were kept in 5% CO$_2$ at 37°C. Imaging and time-lapse microscopy were performed using the Leica DMi6000B widefield epifluorescence microscope (Wetzlar) with a Leica HCX PL APO 100 x 1.4 oil objective and the Leica LAS AF software. Confocal live cell images of parasites were acquired using the Leica TCS SP8 confocal microscope with the HC PL APO 63 x/1.4 oil objective and the Leica Application Suite X software. Imaging of PbC6GFP parasites stained with ER-Tracker™ Red and FLIP analysis of PbC6GFP-Sec61β parasites was performed using a confocal spinning disc microscope (Till Photonics) and a 60 x oil objective (numerical aperture 1.35). For FLIP analysis, a bleaching area of 3.75 μm$^2$ was selected within the parasite and for the control analysis, to account for bleaching by imaging, outside of the parasite. Fluorescence intensity (ROI) was measured for the whole parasite apart from the bleaching area plus a 1.5 μm wide
threshold region. A region outside of the cell was selected for background correction. After acquiring three pre-bleach images, bleaching was performed using 100% power of a 488 nm solid-state laser with five iterations per bleaching. Parasites were bleached 20 times at 10 s intervals and fluorescence intensity was normalized to the mean intensity of the corresponding prebleached images.

In vivo analysis of PbSec61β parasites

PbCsGFP-Sec61β (250,000) sporozoites were injected intravenously into C57BL/6 mice. Intravital microscopy was performed as previously published (Thiberge et al., 2007) at either 42 or 49 hpi using the LSM 510 Zeiss microscope in the LSM 5 live mode. Images were acquired by confocal line scanning microscopy with the Zeiss Plan-Apochromat 63×/1.40 oil DIC M27 objective and the Zeiss LSM 5 Duo Release software.

Serial block-face scanning electron microscopy and transmission electron microscopy

HeLa cells (5 × 10^4) were seeded per well into a 96-well plate and infected with PbCsCherry parasites, as described above. Infected cells were FACS-sorted at 6 hpi for PbCsCherry infected cells and re-seeded at a density of 10,000 sorted cells per well into a 96-well optical plate (Greiner bio one). Following this, parasites were fixed in a glutaraldehyde buffer at 48 hpi and processed according to a previously published protocol (Deerick et al., 2010). SBFSEM images were acquired with a Quanta FEG 250 (FEI Company) equipped with a Gatan 3View2XP ultramicrotome (accelerating voltage = 3.5 kV; low vacuum). TEM images were acquired using a Philips/FEI CM12 Transmission Electron Microscope. Images were processed using Fiji.

Statistical analysis

Statistical analysis was conducted with GraphPad Prism version 6.0. An unpaired, two-tailed t-test was used for analysing data of the detached cell assay and for analysing the nature of ER accumulations of inhibitor-treated parasites. The Mann–Whitney test was used to analyse the size of parasites treated with inhibitors and to compare accumulation sizes measured either with the Leica DMi6000B widefield epifluorescence microscope or SBFSEM. Experiments were conducted in biological triplicates. The error is depicted as 95% confidence intervals. *P* values < 0.05 were considered significant.

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Author contributions

Conception or design of the study: GK, RRS, VTH. Acquisition, analysis or interpretation of the data: GK, RRS, BZ, BK, MDN, PCB, VTH. Writing of the manuscript: GK, VTH, RRS. All authors read and approved the final manuscript.

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Supporting information
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