**Plasmodium Secretion Induces Hepatocyte Lysosome Exocytosis and Promotes Parasite Entry**

**HIGHLIGHTS**

- *Plasmodium* sporozoites induce host lysosome exocytosis during invasion.
- Hepatocyte lysosome exocytosis occurs in a SPECT2-independent manner.
- Inhibition of lysosome-plasma membrane fusion inhibits sporozoite invasion.
- Secreted parasite factors are sufficient to induce lysosome exocytosis.

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**SUMMARY**

The invasion of a suitable host hepatocyte by *Plasmodium* sporozoites is an essential step in malaria infection. We demonstrate that in infected hepatocytes, lysosomes are redistributed away from the nucleus, and surface exposure of lysosome-associated membrane protein 1 (LAMP1) is increased. Lysosome exocytosis in infected cells occurs independently of sporozoite traversal. Instead, a sporozoite-secreted factor is sufficient for the process. Knockdown of SNARE proteins involved in lysosome-plasma membrane fusion reduces lysosome exocytosis and *Plasmodium* infection. In contrast, promoting fusion between the lysosome and plasma membrane dramatically increases infection. Our work demonstrates parallels between *Plasmodium* sporozoite entry of hepatocytes and infection by the excavate pathogen *Trypanosoma cruzi* and raises the question of whether convergent evolution has shaped host cell invasion by divergent pathogens.

**INTRODUCTION**

*Plasmodium* parasites, the causative agents of malaria, are transmitted to humans by the bite of infected female *Anopheles* mosquitoes. The sporozoite form of the parasite is deposited into human skin during a blood meal. Sporozoites are motile and rapidly migrate through the skin to enter a capillary, which allows the parasite to travel to the liver. *Plasmodium* sporozoites have the capacity to transmigrate through cells using a process termed cell traversal (CT) (Mota and Rodriguez, 2001). Recent studies have demonstrated that sporozoites can also enter hepatocytes within a transient vacuole, independently of CT, and that parasites that are CT deficient within a transient vacuole associate with lysosomes and are eliminated (Risco-Castillo et al., 2015). In contrast, productive invasion occurs when sporozoites invade a hepatocyte, form a parasitophorous vacuole (PV), and develop into a liver stage (LS) schizont, from which merozoites are released into the bloodstream and invade erythrocytes. The secretion of a multitude of sporozoite factors are released during motility, CT, and invasion, yet a precise role for most secreted factors remains undefined.

Membrane vesicle trafficking is fundamental to eukaryotic life and plays a regulatory role in nearly all cellular activities. Many intracellular pathogens target and subvert these trafficking events for their own benefit (Alix et al., 2011; Asrat et al., 2014). Previous work has demonstrated that the *Plasmodium* liver stage PV membrane co-localizes with late endosomes (Petersen et al., 2017), lysosomes (Lopes da Silva et al., 2012; Niklaus et al., 2019; Prado et al., 2015; Risco-Castillo et al., 2015), and autophagic vesicles (Prado et al., 2015; Real et al., 2018; Wacker et al., 2017). Although these studies have suggested that the vesicle association is gradually lost over the course of liver stage infection (Niklaus et al., 2019; Prado et al., 2015; Risco-Castillo et al., 2015), when this association is initiated during *Plasmodium* life cycle progression remains unknown, and none of these studies have evaluated infection prior to 3 h post-entry. Moreover, the role of host vesicular trafficking processes in sporozoite entry of hepatocytes has not been explored.

**RESULTS AND DISCUSSION**

*Plasmodium* Sporozoites Co-localize with LAMP1-Positive Vesicles

We quantitatively surveyed the extent of co-localization between the parasite and five markers of endocytic compartments. Freshly isolated *Plasmodium yoelii* sporozoites were added to Hepa1-6 cells. After 90 min, cells were fixed, stained, and visualized by 3D fluorescence deconvolution microscopy. We used antibodies against early endosome antigen 1 (EEA1) and Ras-related protein 5 (Rab5) to mark early endosomes, Rab7a to mark late endosomes (LE), Rab11a to mark recycling endosomes, and lysosome-associated membrane protein 1 (LAMP1) to mark late endosomes (LE).
protein 1 (LAMP1) to mark LE/lysosomes. Sporozoites were labeled with an antibody against circumsporozoite protein (CSP) (Figure 1A). Intensity-based co-localization was used (Bolte and Cordelieres, 2006) to evaluate the extent of overlap between CSP and staining for each vesicular compartment (Figures 1A and 1B). The Pearson’s correlation coefficient between CSP and LAMP1 was 0.6, but staining did not significantly overlap between CSP and EEA1, Rab5, Rab7a, or Rab11a (Figures 1A and 1B). These data are consistent with earlier observations (Lopes da Silva et al., 2012; Petersen et al., 2017).

Figure 1. Plasmodium Sporozoites Co-localize with LAMP1-Positive Vesicles

(A) Hepa1-6 cells were infected with P. yoelii sporozoites for 90 min and processed for fluorescence microscopy using antibodies to EEA1, Rab5a, Rab11, Rab7a, and LAMP1 (red) and PyCSP (green). Scale bar represents 5 μm.

(B) Pearson’s correlation coefficients were calculated for each endocytic vesicle channel with the sporozoite CSP channel of 25 different microscopic fields from three independent experiments. Hepa1-6 cells were infected with P. yoelii sporozoites for 5 min (C) and 30 min (D) and processed for fluorescence microscopy using DAPI (blue) for DNA, phalloidin (white) for actin visualization, antibodies to LAMP1 (red) for LE/lysosomes, and CSP (green) for parasites. Isospot rendering for LE/lysosomes and isosurface rendering for LE/lysosomes, parasites, host cell nucleus, and plasma membrane are shown. Red isospots represent LAMP1-positive structures co-localized with CSP. Magnified inset is 15 μm x 15 μm.

(E) Hepa1-6 cells were infected with wild type or SPECT2- P. yoelii sporozoites and fixed after 5, 30, 60, and 90 min. Intensity-based colocalization was performed on at least 25 parasites per time point and Pearson’s correlation coefficients were calculated. Box and whiskers plot depict mean ± SD of three independent experiments.
We next assessed the kinetics of co-localization between sporozoites and LAMP1. Hepa1-6 cells were infected with *P. yoelii* sporozoites and fixed after 5 (Figure 1C), 30 (1D), 60, or 90 min (1E). LAMP1 structures that co-localized with CSP were observed as early as 5 min and were elongated in the infected cells (Figure 1C). These elongated LAMP1 structures were not observed in bystander or unexposed cells. Thus, the association between LAMP1-positive LE/lysosomes and sporozoites occurs during or very soon after infection and is maintained. We observed a very similar association between LAMP1 and CSP in the CT-deficient parasite, *PySPECT2* (Ishino et al., 2004) (Figures 1E, S1A, and S1B). Our data are consistent with the hypothesis that lysosomes interact with the sporozoite during or very soon after entry, independently of CT. Since *PySPECT2* parasites cannot egress from the transient vacuole, these data do not distinguish between productive and non-productive entry.

**Sporozoite Entry Is Associated with LE/Lysosome Redistribution**

Lysosomes are typically located in juxtanuclear regions of the cell under basal conditions but can be redistributed under times of stress or during infection (Yokota et al., 1989). To evaluate lysosome localization during *Plasmodium* infection, we infected Hepa1-6 cells with *P. yoelii* sporozoites and fixed after 30 or 90 min (Figure 2A). To assess the quantity and localization of lysosomes within infected and uninfected cells we defined LE/lysosomes as LAMP1-positive structures between 0.25 and 1 µm in diameter, corresponding to the typical size of LE/lysosomes within the mammalian cell. Hepa1-6 cells contained an average of ~450 LAMP1-positive structures, similar to measurements obtained by other groups (Cortez et al., 2016). We defined perinuclear lysosomes as LAMP1-positive structures that were within a region surrounding the nucleus, delineated by extrapolating the DAPI signal (Cortez et al., 2016). In unexposed or mock-treated samples containing material from the salivary glands of uninfected mosquitoes, ~85% of LE/lysosomes were perinuclear (Figure 2A). In infected cells, lysosomes were slightly higher in number (Figure 2B) and significantly less perinuclear (Figure 2C). Interestingly, bystander cells, which were defined as being immediately proximate to the infected cell, also exhibited an increase in lysosome numbers and redistribution (Figures 2B and 2C).

To assess the fate of redistributed lysosomes, we asked if there was evidence of LAMP1 positive-vesicle fusion with the hepatocyte plasma membrane in infected cells. We infected Hepa1-6 cells with *P. yoelii* sporozoites and evaluated total and surface-exposed LAMP1 (sLAMP1) by flow cytometry (Figure 2D) and immunofluorescence microscopy (Figure S2A). Both LAMP1 and sLAMP1 were elevated in infected cells compared with exposed uninfected and unexposed control cells (Figures 2D and 2E). Interestingly, little impact on sLAMP1, if any, was observed in uninfected cells, despite our earlier observation that lysosomes redistribute in these cells. These data suggest that lysosomes traffic away from the nucleus in infected and neighboring cells but undergo exocytosis only in infected cells. Although we observed an increase in the total number of lysosomes in bystander cells using microscopy-based methodology, we did not observe an increase in total LAMP1 when it was evaluated by flow cytometry on uninfected cells within infected cultures. This difference might originate from the lack of spatial information that can be resolved in flow cytometry experiments. Alternatively, the ability to quantify LAMP1-positive vesicles may be greater using microscopy-based methodology, which could result in our inability to distinguish the modest difference in total LAMP1 levels between bystander and uninfected cells. We observed a similar pattern of lysosome redistribution (Figures S2B–S2D) and elevated levels of sLAMP1 (Figures S3A and S3B) when we infected with *PySPECT2* . Together, these data suggest that lysosome trafficking and exocytosis are altered in infected hepatocytes, independently of CT.

The PV membrane (PVM) is critical for liver-stage development. Soon after productive infection, parasite factors, including upregulated in infectious sporozoites 4 (UIS4), are translated and trafficked to the PVM (Matuschewski et al., 2002). We infected Hepa1-6 cells with wild-type *P. yoelii* sporozoites and fixed samples 3 h after infection. LS parasites with an intact PVM were distinguished by positive CSP and UIS4 staining, and LS parasites positive for CSP but negative for UIS4 were defined as unsuccessful invasion events. We observed co-localization of LAMP1 with parasite markers in both cases (Figure S3C), suggesting that lysosomal contents are associated with intracellular parasites, independently of the status of their PVM.

**The Role of Lysosome Exocytosis Varies across Species of Intracellular Parasites**

Lysosomes have been previously demonstrated to play a role in *Trypanosoma cruzi* host cell entry (Tardieux et al., 1992). Specifically, a portion of *T. cruzi* parasites utilize a lysosome-mediated event to enter the host cell (Hissa et al., 2012; Tardieux et al., 1992). In contrast, *Toxoplasma gondii*, an apicomplexan parasite...
closely related to \textit{Plasmodium}, sequesters host lysosomes to the vacuolar space (Coppens et al., 2006) but is not thought to use lysosomes to facilitate host cell entry. To elucidate how the invasion of \textit{Plasmodium} parasites is compared with these two disparate systems, we infected Hepa1-6 cells with \textit{P. yoelii} sporozoites, \textit{T. gondii} tachyzoites, or \textit{T. cruzi} trypomastigotes and assessed infection and sLAMP1 by flow cytometry after 90 min. Cells infected with \textit{P. yoelii} or \textit{T. cruzi}, but not \textit{T. gondii}, exhibited increased sLAMP1 (Figures 3A and 3B).
Figure 3. The Role of Lysosome Exocytosis Varies across Species Of Intracellular Parasites

(A and B) Hepa1-6 cells were infected with *P. yoelii* sporozoites, *T. gondii* tachyzoites or *T. cruzi* trypomastigotes for 90 min and analyzed by flow cytometry. The histogram shows surface LAMP1 from infected, uninfected, and unexposed control cells. Surface LAMP1 is expressed as fold change over uninfected cells. The bar graph displays the mean ± SD of three independent experiments. *p < 0.001.

(C) Hepa1-6 cells were transduced with shRNA lentiviruses against SNAP23, SYN4, SYT7, VAMP7, or a scrambled control and challenged with *P. yoelii* sporozoites, *T. gondii* tachyzoites, or *T. cruzi* trypomastigotes for 90 min. The bar graph displays the infection rate after knockdown of each transcript of interest normalized to scramble shRNA cells, indicated by dashed line (n = 5 for *Plasmodium* and *Trypanosoma*; n = 7 for *Toxoplasma* infection; mean ± SD).

(D) Hepa1-6 cells were incubated with or without the indicated compound for 15 min, washed, and then infected with the indicated parasite for 90 min. The bar graph represents mean ± SD of three independent experiments.

(E) Pearson’s correlation coefficients were calculated from the data in (D) for each pairwise combination of infections.
Fusion between lysosomes and the plasma membrane is mediated by the SNARE complex, which includes synaptotagmin VII (SYT7), syntaxin 4 (SYN4), vesicle-associated membrane protein 7 (VAMP7), and synaptoosomal-associated protein 23 (SNAP23) (Rao et al., 2004). We knocked down each factor in Hepa1-6 cells using lentivirus-encoded shRNAs and observed decreased levels of transcript (Figure S4A) and reduced sLAMP1 (Figure S4B) (Rao et al., 2004). We then infected each knockdown line with *T. gondii*, *P. yoelii*, or *T. cruzi* parasites. Knockdown of each member of the SNARE complex significantly reduced *P. yoelii* and *T. cruzi* infections, but not *T. gondii* infection (Figure 3C), and did not reduce the viability of infected cells (Figure S4C).

Genetic knockdowns can sometimes lead to compensatory changes that produce off-target effects. To partially circumvent this, we evaluated the impact of a range of small molecules that modulate lysosome exocytosis (Table S1). We treated Hepa1-6 cells with each compound for 15 min, washed the cells, and then infected with *P. yoelii* sporozoites, *T. gondii* tachyzoites, or *T. cruzi* trypomastigotes for 90 min. Molecules that increase lysosome exocytosis or redistribution (ionomycin, thapsigargin, brefeldin A; Table S1 and Figure S4C) significantly increased *P. yoelii* and *T. cruzi* infection (Figure 3D). In contrast, pretreatment with MJCD, which has been previously shown to reduce surface LAMP1 levels by disrupting lipid rafts, reduced our readout of sLAMP1 (Figure S4D) and diminished *P. yoelii* and *T. cruzi* infections (Figure 3D). Levels of sLAMP1 and infection returned to baseline when cholesterol was added, restoring lipid raft formation. No inhibitors substantially altered *T. gondii* infection (Figure 3D). Overall, changes in *P. yoelii* and *T. cruzi* infections were tightly correlated (Pearson Correlation Coefficient, 0.8297) (Figure 3E), whereas other pairwise comparisons were less correlated. Our data suggest that *T. cruzi* and *P. yoelii*, but not *T. gondii*, rely on a lysosome-mediated mechanism to enter the host cell. The extent of these parallels, and ways in which the entry strategies diverge, remains an important area for further investigation.

**Secreted Sporozoite Factor(s) Contribute to Lysosome Redistribution during Invasion**

Lysosome exocytosis is induced by the soluble *T. cruzi* factor Tcgp82 (Cortez et al., 2016). To assess if a parallel process occurs during *Plasmodium* entry, we treated sporozoites with FBS to induce secretion and then collected supernatants. We exposed Hepa1-6 cells to this sporozoite-derived, secretion-enriched supernatant at different sporozoite:hepatocyte ratios for 90 min. Cells were monitored for lysosome redistribution by 3D fluorescence microscopy (Figures 4A, 4B, 5A) and sLAMP1 by flow cytometry (Figure 4C). Treating cells with even low quantities of secretion-enriched supernatant, but not heat-inactivated supernatant, promoted lysosome redistribution (Figure 4B), and sLAMP1 was induced in a dose-dependent manner (Figures 4C and 5B). Therefore, sporozoite-induced lysosome redistribution is impacted by different factors or the same factors at different levels than lysosome exocytosis. These results are consistent with a model where two separate secretion-mediated events induce hepatocyte lysosome redistribution and lysosome exocytosis (Figure 4D). Taken together, our data suggest a role for lysosome exocytosis in hepatocyte entry of sporozoites, independently of CT or the presence of the PVM.

A growing collection of evidence suggests that parasites that differ only slightly in genetic makeup can exhibit drastically altered host cell tropism. For example, *Plasmodium* species rely differentially on host proteins CD81 and SRB1 for entry (Manzoni et al., 2017; Risco-Castillo et al., 2014; Silvie et al., 2003), and this relationship cannot be predicted by evolutionary similarity alone (Frech and Chen, 2011). Here, we demonstrate that lysosome-related alterations impact *P. yoelii* and *T. cruzi* infections similarly but have no effect on the apicomplexan parasite, *T. gondii*. These observations raise the question of how quickly pathogens can evolve host cell tropism and whether the similarities we observe are sculpted by convergent evolution. Systematic investigation into mechanisms of host cell invasion across many pathogens with well-defined evolutionary relationships, will allow us to obtain a better understanding of the major influences that shape host cell engagement over evolutionary time.

**Limitations of the Study**

In this study, we addressed the role of lysosome exocytosis during *Plasmodium* liver stage infection. Although we systematically explored this process using temporally resolved immunofluorescence imaging, our work does not definitively demonstrate that lysosome exocytosis occurs at the moment of infection. Subsequent work that utilizes live cell imaging could add temporal detail to our findings.
The functional studies contained within this manuscript are performed using pharmacological inhibitors and shRNA-mediated knockdown, each which have off-target effects. Moreover, we have performed these experiments using cell lines in vitro. These technical limitations represent potential caveats of our study.

**Figure 4. Secreted Sporozoite Factor(s) Contribute to Lysosome Redistribution during Invasion**

(A) Hepa1-6 cells were infected with *P. yoelii* sporozoites or treated with sporozoite secretion-enriched supernatants at different sporozoite-hepatocyte (S:H) ratios. After 90 min, cells were processed using DAPI (blue) for DNA, phalloidin (white) for actin visualization, antibodies to LAMP1 (red) for LE/lysosomes, and CSP (green) for parasites and displayed as maximum intensity projections. Scale bar represents 5 μm. The isospots corresponding to lysosomes away from the nucleus and perinuclear lysosomes were depicted in cyan and magenta, respectively.

(B) Values represented in bar graphs correspond to percentage of perinuclear lysosomes identified in (A). Data represent the mean ± SD of at least 10 different microscopic fields per condition from three independent experiments.

(C) Hepa1-6 cells were infected with *P. yoelii* sporozoites or exposed to sporozoite secretion-enriched supernatants. Surface LAMP1 was analyzed by flow cytometry. Values represent the mean ± SD of three independent experiments.

*D* *p* < 0.001.

(D) Model of *Plasmodium* sporozoites promoting lysosome exocytosis. Sporozoites are depicted in yellow, hepatocytes in gray, lysosomes in red, and a *Plasmodium*-derived secreted factor in green.
Finally, we demonstrate that secreted parasite factor(s) are sufficient to induce hepatocyte lysosome exocytosis. Subsequent work will elucidate the specific effectors of this process.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

DATA AND CODE AVAILABILITY
The data supporting the findings of this study are available within the paper and its Supplemental Information.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2019.10.054.

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AUTHOR CONTRIBUTIONS
K.V., E.K.K.G., I.C., S.M., H.S.K., and A.M.B. performed experiments. K.V., I.C., and F.D.M. analyzed data. J.D.A., K.S., and A.K. supervised the research. K.V. and A.K. wrote the paper with input from all other authors.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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Supplemental Information

*Plasmodium* Secretion Induces Hepatocyte Lysosome Exocytosis and Promotes Parasite Entry

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**Transparent Methods**

**Cell lines and culture**

Hepa1-6 cells were obtained from American Type Culture Collection. Cells were maintained in DMEM-Complete Medium (Dulbecco's modified eagle medium (Cellgro, Manassas, VA), supplemented with 10% FBS (Sigma-Aldrich, St. Louis, MO), 10000 IU/ml penicillin, 100 mg/ml streptomycin (Cellgro), 2.5 mg/ml fungizone (HyClone/Thermo Fisher, Waltham, MA) and 4 mM L-Glutamine (Cellgro). Cells were split 2-3 times weekly. All experiments were performed using Hepa1-6 cells that were passaged between 4 and 20 times after purchase from ATCC.

**Mosquito rearing and sporozoite production**

For *P. yoelii* sporozoite production, female 6–8-week-old Swiss Webster mice (Harlan, Indianapolis, IN) were injected with blood stage *P. yoelii* (17XNL) parasites to begin the growth cycle. Animal handling was conducted according to the Institutional Animal Care and Use Committee-approved protocols. *Anopheles stephensi* mosquitoes were allowed to feed on infected mice after gametocyte exflagellation was observed. Salivary gland sporozoites were isolated using a standard protocol at day 14 or 15 post-blood meal. The sporozoites were activated with 20% FBS and spun at 1000 x g to remove debris from salivary gland. Spin sporozoites at 15,000 x g for 4 min at 4°C to pellet and resuspend in the desired volume of complete medium.

**T. gondii production**

*T. gondii* strain RHΔHXGPR T Gra2:GFP, Tub:βgal, a kind gift from Marilyn Parsons (CIDR, Seattle), was maintained by continual cycling through human foreskin fibroblasts (HFF). For
infections, parasites were lysed from HFFs by passing 2× through a 27-gauge needle and then counted on a hemocytometer.

**T. cruzi production and labeling**

Tissue culture-derived trypomastigotes from the *T. cruzi* Cl Brener strain were obtained by weekly passage in confluent monolayers of Hepa1-6 cells at 37 °C and 5% CO₂, in DMEM medium supplemented with 10% FBS. Motile trypomastigotes were obtained from the supernatant and purified as previously described (Schenkman et al., 1991) and stained with 5(6)-Carboxyfluorescein diacetate N-succinimidy ester (Sigma Aldrich) as suggested by manufacturer’s protocol.

**shRNA-mediated gene knockdown**

MISSION shRNA vectors for SNAP23, VAMP7, SYT7 and SYN4 were obtained from Sigma Aldrich (St. Louis, MO). Non-replicating lentiviral stocks were generated by transfection of HEK293-FT cells. 4 × 10⁶ HEK293-FT cells were plated on poly-L-lysine coated dishes to achieve 70-80% confluency at time of transfection. Approximately 24 h after plating, transfection mixtures were prepared by mixing 20 µL Polyethylenimine MAX (Polysciences Inc, Warrington, PA) prepared at 1 mg/ml, together with 4.75 µg of shRNA construct or a scramble shRNA control, 1.5 µg viral envelope plasmid (pCMV-VSV-G), and 3.75 µg viral packaging plasmid (psPax2). After incubating for 10 min at room temp in DMEM, transfection complexes were added drop-wise to cells. After overnight incubation, cells were washed to remove transfection mixtures and were fed with 10 mL fresh media. Lentivirus-containing supernatant was harvested 36 hours later, passed through 0.45 µm syringe filters, and either used immediately for transduction or stored at -80 °C.
To induce knockdown of candidate SNARE proteins, Hepa1-6 cells were transduced with lentiviral supernatants in 6-well plates at a cell density of $1 \times 10^6$ per well. At time of plating, cells were transduced with 1 mL of supernatant in the presence of 0.5 µg/mL polybrene (Sigma Aldrich St. Louis, MO). In order to select for cells with stable integration of shRNA transgenes, supernatant was replaced with complete media with the addition of 2 µg/mL puromycin 24 h post-transduction, and cells were selected for at least 5 days prior to experiments.

**Validation and quantification of shRNA mediated knockdown**

Total RNA was extracted using TRIzol reagent according to the manufacturer’s procedure (Invitrogen). cDNA synthesis was performed using the Thermo Scientific RevertAid RT Kit according to the manufacturer’s instructions (Thermo Scientific). For quantitative PCR (qPCR) a standard curve was generated using 1:4 dilutions of a reference cDNA sample for PCR amplification of all target PCR products using target specific primers (Table S1). The values of each transcript were normalized to mouse GAPDH. Experimental samples were compared to this standard curve to give a relative abundance of transcript.

**Infection assays**

$5 \times 10^5$ Hepa1-6 wild type cells or knockdowns cells were seeded in each well of a 24-well plate (Corning) and infected with *P. yoelii* sporozoites at a multiplicity of infection (MOI) = 0.25, *Toxoplasma gondii* tachyzoites (MOI = 0.5) or *Trypanosoma cruzi* trypomastigotes (MOI = 5) for 90 min. For small molecule treatment experiments, Hepa1-6 cells were treated with or without 10 µM ionomycin, 400 nM thapsigargin, 10 µM brefeldin A, 10 µM nocodazole and 5mM methyl-β-cyclodextrin (MβCD) for 15 min, washed and infected for 90 min. For cholesterol replenishment, MβCD treated cells were washed and incubated with 1 mM cholesterol for 15 min before parasite
addition. Cells were stained with Live/Dead marker after 60 mins of infection as per manufacturer’s instruction (ThermoFischer). After 90 min of infection, cells were harvested with accutase (Life technologies) and fixed with Cytoperm/Cytofix (BD Biosciences). Cells were blocked with Perm/Wash (BD Biosciences) + 2% BSA for one hour at room temperature then stained overnight at 4 °C with primary antibody. Cells were washed three times with PBS then stained for one hour at room temperature with secondary antibodies. The cells were then washed and resuspended in PBS + 5 mM EDTA. Infection rate was measured by flow cytometry on an LSRII (Becton-Dickinson) and analyzed by FlowJo (Tree Star). Surface LAMP-1 levels were calculated using mean fluorescent intensity of specific population in FlowJo and represented as fold change between infected and control population.

For the evaluation of surface expression of LAMP-1, cells were detached using Accutase (Sigma) and were incubated with a polyclonal antibody to LAMP1 (DSHB) in medium + 2% BSA for 30 minutes in ice, fixed with 3.7% paraformaldehyde, permeabilized with 0.01% Triton X-100. Cells were then stained with monoclonal antibody to *P. yoelii* Circumsporozoite protein (CSP) conjugated to AlexaFluor 488 (Life Technologies) at 1:500, *T. gondii* P30 mouse monoclonal antibody at 1:1000 (Novus Biologicals). Cells were washed three times with PBS then stained for one hour at room temperature with secondary antibodies. The cells were washed and suspended in PBS+5 mM EDTA. Infection rate and surface expression of LAMP1 was measured by flow cytometry on an LSRII (Becton-Dickinson) and analyzed by FlowJo (Tree Star).

**3D Fluorescence Microscopy**

For imaging experiments, Hepa1-6 cells were plated in 8 well chamber slides (Labtek) and infected with *P. yoelii* wild type or SPECT2⁻ sporozoites. Cells were fixed with 10% formalin (Sigma) at
defined timepoints after infection (5, 30, 60, or 90 min), permeabilized with Triton X-100, and
stained with rabbit anti-EEA1 (CST), rabbit anti-Rab5 (CST), rabbit anti-Rab7 (CST), rabbit anti-
Rab11 (CST), rat anti-LAMP1 (DSHB), goat anti-UIS4 (SCI-GEN) and mouse anti-CSP
antibodies. Nuclei were stained with DAPI (Vectashield) and AlexaFluor 647-phalloidin (Life
technologies) was used for actin visualization. Images were acquired with a 100× 1.4 NA objective
(Olympus) on a DeltaVision Elite High Resolution Microscope (GE Healthcare Life Sciences).
The sides of each pixel represent 64.5 × 64.5 nm and z-stacks were acquired at 300 nm intervals.
Approximately 20-30 slices were acquired per image stack. For deconvolution, the 3D data sets
were processed to remove noise and reassign blur by an iterative Classic Maximum Likelihood
Estimation widefield algorithm provided by Huygens Professional Software (Scientific Volume
Imaging BV, The Netherlands).

**Image analysis and quantification**

Imaris software (Bitplane) was used to obtain 3D reconstructions of the fluorescence microscopy
image stacks and quantification of lysosomes in x-y-z coordinates. Deconvolved images of
immunostained cells stained with anti-LAMP1 (lysosomes) anti-CSP (sporozoites) and phalloidin
(actin) with DAPI (nucleus) were processed, thresholded and segmented by Imaris software, to
render isospots and isosurfaces from the fluorescence signal (Real and Mortara, 2012). Phalloidin
channel was used for the 3D reconstruction of the cells and only the sporozoites encased inside the
3D phalloidin structure are considered as invaded sporozoites and proceeded further. Isosurfaces
were constructed by extrapolating the DAPI signal to the local minima (Cortez et al., 2016) in
order to define a perinuclear region where lysosomes can be differentially counted. Using a mask
tool, all LAMP1 signal outside this nuclear/perinuclear isosurface was suppressed, allowing
addition of a new LAMP1 fluorescence channel corresponding exclusively to LAMP1 localized
in perinuclear area. After image processing, we obtained an unmasked LAMP1 signal corresponding to total lysosomes and a masked LAMP1 signal localized to the perinuclear region, corresponding to perinuclear lysosomes. Isospots were constructed based on these two classes of LAMP1 signal, which allowed the quantification of total and perinuclear lysosomes per cell for each image-stack. The surface segmentation function of Imaris was used to identify the cell boundary using phalloidin-signal. Intensity based co-localization was performed by creating region of interest (ROI) specific to the sporozoite structure in the CSP channel using the Imaris isosurface module. Pearson’s correlation coefficient for co-localization analysis of endocytic vesicles and CSP was performed in the ROI using the Imaris co-localization module.

**Generation of sporozoite supernatant**

Salivary gland sporozoites were isolated using standard protocols. Sporozoites incubated with 20% FBS for 20 min at RT and spun at 13,000 x g for 4 min at 4 °C. The supernatant is collected and again spun at 13,000 x g for 4 min at 4 °C to ensure the preparation is free of intact sporozoites. Hepal-6 cells were exposed to supernatants at different sporozoite to hepatocyte ratio for 90 min. Cells were washed and subjected to 3D immunofluorescence microscopy or flow cytometry as described previously.

**Statistical analysis**

p-values were determined in GraphPad Prism 8 software using two tailed end t-test for samples with unequal variance.
**Fig. S1.** CT deficient or wild type *Plasmodium* sporozoites interacts with lysosomes in similar fashion. Immunofluorescent microscopy of Hepa1-6 cells infected with SPECT2<sup>−</sup> *P. yoelii* sporozoites. Infection was assessed after (a) 5 or (b) 30 min and processed for fluorescence microscopy using DAPI (blue) for DNA, Phalloidin (white) for actin visualization, antibodies to LAMP1 (red) for LE/lysosomes and CSP (green) for parasites. Isospots for LE/lysosomes and isosurfaces for LE/lysosomes, parasites, host cell nucleus and plasma membrane were created using Imaris software and the LE/lysosomes interacting with parasites were identified by detecting overlap between isospots and the isosurface. Red spots represent LAMP1-positive structures colocalized with CSP. Magnified inset is 15 µm x 15 µm. Related to Figure 1.
Supplementary Figure 2

(a) DAPI CSP LAMP1 PHALLOIDIN MERGE

3D segmented reconstruction

90 min post-infection with SPECT2

(b) DAPI CSP LAMP1 PHALLOIDIN MERGE 3D segmented reconstruction

30 min post-infection with SPECT2

90 min post-infection with SPECT2

(c) 30 minutes

Perinuclear lysosomes (%)

Control Infected Bystander

p = 0.03

Wildtype SPECT2 NS

(d) 90 minutes

Perinuclear lysosomes (%)

Control Infected Bystander

Wildtype SPECT2 NS NS NS
**Fig. S2. Sporozoite induced lysosome-plasma membrane fusion is independent of cell traversal.** (a) Hepa1-6 cells were infected with *P. yoelii* sporozoites and fixed after 90 min. Cells were stained with antibodies to LAMP1 prior to permeabilization and stained with DAPI (blue) for DNA, phalloidin (white) for actin visualization, antibodies to CSP (green) for parasites and displayed as maximum intensity projections. Bar = 5 µm. (b) Hepa1-6 cells were infected with SPECT2− sporozoites and fixed after 30 and 90 min. Cells were processed for 3D fluorescence microscopy using DAPI (blue) for DNA, phalloidin (white) for actin visualization, antibodies to LAMP1 (red) for LE/lysosomes and CSP (green) for parasites and displayed as maximum intensity projections. Scale bar = 5 µm. Images were obtained on a Deltavision fluorescence microscope and processed by Imaris software to construct isosurfaces (nuclei and parasite) and LAMP1-positive isospots (LE and lysosomes) by predefined algorithms for identification of surfaces and spots. Perinuclear isosurfaces were created by extrapolating the DAPI signal to define a perinuclear region where lysosomes could be differentially quantified. The isospots corresponding to total and perinuclear lysosomes were depicted in cyan and red, respectively. Bar = 5 µm. (c and d) Values represented in box and whiskers plot correspond to lysosomes from total and perinuclear area represented as mean ± SD of 25 different microscopic fields from three independent experiments. Related to Figure 2.
Supplementary Figure 3

(a) Flow cytometry histogram showing the surface expression of LAMP1- Alexa Fluor 594 compared to SPECT2.

(b) Bar graph showing the MFI fold change in LAMP1 (over control cells) with infected and uninfected samples. A p-value of 0.0006 indicates statistical significance.

(c) Confocal microscopy images of CSP Lamp1 and Lamp1 UIS4 merge.
**Fig. S3. Sporozoite induced lysosome-plasma membrane fusion is independent of productive invasion.** (a) Hepa1-6 cells were infected with SPECT2⁻ *P. yoelii* sporozoites for 90 min and analyzed by flow cytometry using antibodies specific to LAMP1 and CSP. Surface LAMP1 was evaluated by staining cells prior to permeabilization, while total LAMP1 was evaluated by staining for LAMP1 after permeabilization. The histogram shows the distribution of surface LAMP1 from SPECT2⁻ infected, uninfected and unexposed control cells from one of three independent experiments. (b) Surface LAMP1 levels were compared between uninfected and SPECT2⁻ infected cells as a fold change over control cells. The bar graph depicts the mean ± the SD of three independent experiments. (c) Hepa1-6 cells were infected with *P. yoelii* sporozoites and fixed after 90 min. Cells were stained with DAPI (blue) for nuclei visualization, antibodies to LAMP1 for lysosomes, UIS4 (white) and CSP (green) for parasite detection and displayed as maximum intensity projections. Bar = 5 µm. Red spots represent LAMP1-positive structures co-localized with CSP. Magnified inset is 15 µm × 15 µm. Related to Figure 2.
Fig. S4. Selective knockdown of SNARE proteins can be achieved using lentivirus-mediated shRNA. (a) Bar graph depicting relative gene knockdown compared to non-targeting control shRNA. Hepa 1-6 cells were transduced with lentivirus expressing shRNA constructs selectively targeting SNARE proteins, or a non-targeting control. Mean knockdown level was determined using qPCR. Values are normalized to non-targeting control which is indicated by solid black line. Data is mean ± the SD of 3 independent experiments. (b) Hepa1-6 cells were transduced with shRNA lentiviruses against SNAP23, SYN4, SYT7, or VAMP7 or a scrambled control and assessed for surface LAMP1 by flow cytometry. Surface LAMP1 levels were compared between scramble and specific knockdowns and expressed as a fold change in mean fluorescent intensity (MFI). The bar graph depicts the mean ± the SD of 5 independent experiments. (c) Hepa1-6 cells were transduced with shRNA lentiviruses against SNAP23, SYN4, SYT7, VAMP7 or a scrambled control and challenged with \( P. \text{yoelii} \) sporozoites for 90 min and stained with Live/Dead stain for 30 mins before fixation. The bar graph displays the dead cell percentage in knockdowns normalized to scramble shRNA cells, indicated by dashed line following infection. (d) Hepa1-6 cells were incubated with or without 10 \( \mu \text{M} \) ionomycin, 400 \( \text{nM} \) thapsigargin, 10 \( \mu \text{M} \) brefeldin A, 10 \( \mu \text{M} \) nocodazole, 5 \( \text{mM} \) methyl-\( \beta \)-cyclodextrin (M\( \beta \)CD) for 15 min, fixed and assessed for surface LAMP1 by flow cytometry. For cholesterol replenishment, methyl-\( \beta \)-cyclodextrin (M\( \beta \)CD) treated cells were incubated with 1 \( \text{mM} \) cholesterol for 15 min prior to fixation. Surface LAMP1 levels were compared between DMSO vehicle control and specific treatments and expressed as a fold change in mean fluorescent intensity (MFI). The bar graph depicts the mean ± the SD of three independent experiments. Related to Figure 3.
Supplementary Figure 5

a

![Graph showing normalized count vs. PE Texas Red - Lamp1 for different conditions: Control, Uninfected, Infected, S:H ratio 1:5, S:H ratio 1.0, S:H ratio 0:5, S:H ratio 1:5 (HI).]
Fig. S5. Sporozoite secreted factor(s) contribute to relocalisation of lysosomes. Hepa1-6 cells were infected with *P. yoelii* sporozoites or exposed to sporozoite secretion-enriched supernatants. Surface LAMP1 was analyzed by flow cytometry. Histogram is representative of three independent experiments. Related to Figure 4.
Supplementary Table 1

Lysosomal trafficking modulators selected for the study. Related to Figure 3

| Inhibitor            | Function                                                                 | Reference                                      |
|----------------------|----------------------------------------------------------------------------|------------------------------------------------|
| Ionomycin            | ionophore, increases intracellular Ca++ and induces lysosome exocytosis.   | (Xu et al., 2012)                              |
| Brefeldin A          | redistributes LE/lysosomes towards periphery                              | (Lippincott-Schwartz et al., 1991; Tardieux et al., 1992) |
| Nocodazole           | a microtubule-depolymerizing agent, prevents lysosome redistribution.     | (Tardieux et al., 1992)                       |
| Thapsigargin         | inhibitor of the sarco/endoplasmic reticulum Ca++ ATPase, elevates cytosolic Ca++ and promotes lysosome exocytosis. | (Sivaramakrishnan et al., 2012)                |
| Methyl-β-cyclodextrin| depletes membrane cholesterol and reduces LAMP1 levels on surface.        | (Hissa et al., 2012)                          |
| Wortmannin           | PI3K inhibitor, inhibits various stages of endocytic network.             | (Sinnberg et al., 2009)                       |
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