Drug-induced permeabilization of parasite’s digestive vacuole is a key trigger of programmed cell death in *Plasmodium falciparum*

J-H Ch’ng1, K Liew2, AS-P Goh1, E Sidhartha1 and KS-W Tan*1,2,3

Having previously characterized chloroquine (CQ)-induced programmed cell death (PCD) hallmarks in the malaria parasite *Plasmodium falciparum* and delineating a pathway linking these features, the roles of non-classical mediators were investigated in this paper. It was shown that the later stages of this pathway are Ca2⁺-dependent and transcriptionally regulated. Moreover, it was demonstrated for the first time that micromolar concentrations of CQ partially permeabilized the parasite’s digestive vacuole (DV) membrane and that this important upstream event appears to precede mitochondrial dysfunction. This permeabilization of the DV occurred without rupture of the DV membrane and was reminiscent of lysosome-mediated cell death in mammalian cells. As such micromolar concentrations of CQ are found in the patient’s plasma after initial CQ loading, this alludes to a clinically relevant antimalarial mechanism of the drug which has yet to be recognized. Furthermore, other ‘non-antimalarial’ lysosomotropic compounds were also shown to cause DV permeabilization, triggering PCD in both CQ-sensitive and -resistant parasites. These findings present new avenues for antimalarial developments, which induce DV destabilization to kill parasites.

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Although much progress has been made towards controlling and eliminating malaria in some countries, half the world’s population continues to be at risk of infection with an estimated one million deaths occurring annually.1 Although artesunate-based antimalarials are still effective in treating the disease, resistance to these last-line antimalarials has already been observed near the Thai–Cambodian border, where parasite clearance times have been delayed.2 It is therefore essential for new antimalarial leads to be discovered such that therapeutic options stay ahead of resistance development.3

We have previously suggested that a novel programmed cell death (PCD) mechanism may be useful in aiding antimalarial design.4 Though there have been reports of PCD features occurring in unicellular organisms like *Blastocystis*, *Leishmania*, *Trypanosoma*, *Saccharomyces*, and *Trichomonas*,5 the existence of such a pathway in *Plasmodium* remains controversial. PCD features in *Plasmodium falciparum*6–10 and the rodent malaria parasite *P. berghei*11,12 have been documented, characterized by the fragmentation of DNA and a loss of mitochondria membrane potential. However, other reports suggested that *Plasmodium* does not undergo PCD.6,13,14

Accounting in part the discrepancies between these studies, our previous work demonstrated that PCD features were indeed observable in parasites exposed to high micromolar concentrations of chloroquine (CQ) but not at lower nanomolar levels.4 Moreover, these features were shown to be linked in a linear pathway: low-levels of cysteine protease activity triggered mitochondrial dysregulation, which preceded the amplification of cysteine protease activity and ultimately led to DNA fragmentation and the parasite’s death. Through the use of specific cysteine protease inhibitors, clan CA cysteine proteases were implicated as the key mediators of this pathway, contrary to the hypothesis that clan CD proteases are responsible.8 Additionally, the extent of PCD features was shown to vary between different strains, with CQ-resistant parasites showing reduced cell death features than CQ-sensitive parasites when exposed to the same dose of CQ.

This study began with the identification of other ‘non-classical’ mediators involved in this CQ-induced PCD pathway through the use of various inhibitors of cell death, and later stages were shown to be Ca2⁺-dependent and transcriptionally regulated. The role of the parasite’s digestive vacuole (DV) in facilitating this cell-death pathway was also

1Department of Microbiology, National University of Singapore, Singapore, Singapore; 2Singapore-MIT Alliance for Research and Technology, National University of Singapore, Singapore, Singapore; 3Infectious Disease Program, National University of Singapore, Singapore, Singapore

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Abbreviations: PCD, programmed cell death; CQ, chloroquine; DV, digestive vacuole; PPy-KO, pre-procathepsin K knockout; ROS, reactive oxygen species; RNS, reactive nitrogen species; CM-CQ, coumarin-labelled chloroquine; TEM, transmission electron microscopy; ΔΨm, mitochondria transmembrane potential; LMP, lysosomal membrane permeabilization; PBN, phenyl-tert-butylnitronone; DSP, desipramine; CPZ, chlorpromazine; PMZ, promethazine; 4HT, 4-hydroxytamoxifen; LL-OMe, Leucine–Leucine-O-Methyl Ester

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investigated and it was demonstrated for the first time that the destabilization of this organelle with lysosomotropic compounds led to PCD in the parasite. This is reminiscent of the lysosomal membrane permeabilization (LMP)-associated PCD pathway well studied in mammalian cell lines. Unlike CQ, treatment with these lysosomotropic compounds led to PCD features, which were consistent between CQ-resistant and -sensitive parasite strains, suggesting that such an antimalarial strategy would likely be effective even against CQ-resistant parasites.

**Results**

**Inhibitors of CQ-induced PCD.** To further characterize mediators in the CQ-induced PCD pathway, 3D7 parasites were pre-treated with a panel of inhibitors at various concentrations for 30 min before being induced with CQ. This panel of inhibitors consisted of calcium chelator BAPTA (1–100 μM), clan CD cysteine protease-specific inhibitor Q-VD-OPh (10–50 μM), V-ATPase inhibitor bafilomycin A1 that inhibits the acidification of the DV (10 nM–1 μM), calpain inhibitor calpastatin (10 nM–10 μM), reactive nitrogen species (RNS) scavenger phenyl-tert-butylnitrone (PBN, 10 μM–10 mM), reactive oxygen species (ROS) scavengers glutathione and mannitol (10 μM–10 mM for both), translation inhibitor cycloheximide (1–50 μM) and transcription inhibitor actinomycin D (1–50 μM). Cells were then costained with JC-1 and Hoechst, and assayed for loss of mitochondria transmembrane potential (Δψm) and DNA loss, respectively.

None of the inhibitors tested reduced the proportion of parasites showing loss of Δψm (data not shown). It is therefore unlikely that any of the suspected mediators are involved in causing mitochondrial dysregulation in CQ-induced PCD.

Most of the compounds tested did not affect DNA degradation (data not shown), except for BAPTA, cycloheximide and actinomycin D, which significantly reduced the proportion of cells showing sub-G1 DNA content (Figure 1). This suggested that DNA degradation, a late-stage feature of the pathway, is dependent on Ca2⁺, and is also transcriptionally regulated. Furthermore, the involvement of calpains and clan CD cysteine proteases like metacaspase appears unlikely as calpastatin and Q-VD-OPh were ineffective inhibitors.

**Microarray data and validation.** Microarray analyses of these parasites were performed to elucidate differentially transcribed genes. Of the genes that were dysregulated (Supplementary Data, Supplementary Table S1), a cathepsin C precursor (PFL2290w) was found to be more highly transcribed (6.54-fold increase) at 8 h after treatment. Real-time RT-PCR was performed to verify this (Supplementary Data, Supplementary Figure S2).

**Pre-procathepsin C knockout and corresponding PCD profile.** A 3D7-pre-procathepsin C knockout (PPC-KO) strain was generated (Supplementary Data, Supplementary Figure S3), but JC1 and Hoechst-loss assays on the KO strain showed no difference with the wild type (data not shown). This may have been a result of redundancy of mediators as previously suggested, which complicated further characterization of this protease’s role in cell death.

**Coumarin-labelled CQ and Fluo-4-AM redistribution.** Confocal imaging showed that coumarin-labelled CQ (CM-CQ) and Fluo-4-AM colocalize inside the hemazoin-containing DV, when CM-CQ was administered at 300 nM (Figures 2a and c). However, at 30 μM of CM-CQ, the labelled drug and Fluo-4-AM both dispersed throughout the parasite’s cytoplasm (Figures 2b and c). Coupled with the involvement of clan CA proteases previously demonstrated, the simultaneous redistribution of both CM-CQ and Ca2⁺ led us to hypothesize that a permeabilization of the DV membrane may be triggering the PCD pathway.

**Abrupt Ca2⁺ redistribution.** Time-course studies showed a dramatic redistribution of Fluo-4-AM from the DV, the primary calcium store, to the cytoplasm between the 2–4 h time-points (Figure 3a). This 2–4 h after treatment window was further narrowed using real-time single-cell tracking, which showed that this redistribution of Fluo-4-AM from the DV to the parasite’s cytoplasm occurred very rapidly (within 1 min) in cells at about 3–3.5 h after treatment with CQ (Figure 3b and Supplementary Data, Supplementary Video S4). This observation supported our hypothesis of a DV-breach as an increasing accumulation of lysosomotropic CQ is likely to have reached a threshold at around 3–3.5 h, leading to the DV membrane being compromised. Control cells showed no redistribution of Fluo-4-AM throughout this
period, and the fluorescence signal was not compromised by photobleaching (data not shown).

To validate this model further, CQ-resistant strains bearing the CQ-resistance transporter (PfCRT) mutation were also investigated. Interestingly, 7G8 parasites showed a similar staining profile to that of 3D7 after 4 h exposure to 3 \( \mu \text{M} \) of CQ, while K1 showed a markedly reduced proportion of cells with Fluo-4-AM redistributed to the cytoplasm (Figure 3c). This is likely due to differences in the PfCRT mutations and will be discussed later.

Figure 2  Colocalization studies of CM-CQ and calcium-sensing dye, Fluo-4-AM. (a) 3D7 parasites (white arrows) were treated with 300 nM of CM-CQ for 10 h, stained in Fluo-4-AM and observed by confocal microscopy. Colocalization of CM-CQ and Fluo-4-AM to the hemazoin-containing DV (black arrows) is clearly seen at such nanomolar concentrations of the drug. (b) Parasites exposed to 30 \( \mu \text{M} \) of CM-CQ show localization of both CM-CQ and Fluo-4-AM to the cytoplasm of the parasite but not the cytoplasm of the host erythrocyte. (c) At least 50 parasites were counted and the results were tabulated to show the association in the localization of Fluo-4-AM and CM-CQ. (d) Fluorescence intensity traces of the cross-section (yellow) of three representative uninduced parasitized erythrocytes stained with Fluo-4-AM shows that the hemazoin-containing DV is intensely stained. (e) The fluorescent trace of three representative induced parasites (3 \( \mu \text{M} \) of CQ for 4 h) shows that Fluo-4-AM fluorescence is distributed throughout the length of the parasite cross-section.
Transmission electron microscopy. To determine whether the DV membrane had been affected as a result of CQ-treatment, ultrastructures of saponin-enriched 3D7 parasites were analyzed. In healthy control parasites, the DV membrane was clearly stained and formed a continuous boundary enveloping the hemazoin crystals (Figure 4ai and ii). In CQ-treated cells, a variety of cell morphologies were observed after 4 h incubation. A majority of cells appeared similar to the healthy control cells with visible and continuous membrane surrounding the DV (Figure 4bi and ii). However, their hemazoin crystals were less well defined and membranous whorls or vacuoles appeared within the DV. Occasionally, cells showed DV with electron-lucent membrane with some having minute breaks showing DV material being extruded into the cytoplasm (Figure 4ci–iii). In a few cells, a massive rupture of the DV membrane was observed, leading to DV contents being emptied into the parasite’s cytoplasm (Figure 4d) – these are likely to be necrotic cells.

Mitochondria dysregulation and DNA degradation induced by lysosomotropic compounds. The JC-1 and Hoechst assay were used to screen a variety of potential DV-permeabilizing compounds through a range of concentrations: Leu–Leu-OMe (LL-OMe, 100 nM–10 mM), norfloxacin (100 nM–100 μM), ciprofloxacin (250 μM–10 mM), miltefosine (100 nM–200 μM), vincristine (50 nM–10 μM), verapamil (VP, 1–200 μM), diltiazem (1–200 μM),...
desipramine (DSP, 1–200 μM), chlorpromazine (CPZ, 10 nM–200 μM), promethazine (PMZ, 10 nM–200 μM), 4-hydroxytamoxifen (4HT, 10 nM–200 μM), propranolol (1 nM–200 μM) and chlorpheniramine (1 nM–200 μM; data not shown).

Of the five compounds, which showed significant PCD features in 3D7 after 10 h incubation, the lowest concentration of the compound at which the features were detected was used on 7G8 and K1 as well (Figures 5a and b). Their concentrations are as follows: 200 μM DSP, 100 μM CPZ, 200 μM PMZ, 150 μM 4HT and 10 mM LL-OMe (due to its limited solubility, the concentration of Leu–Leu could not be increased beyond this even though the PCD features were not extensive).

As previously accounted for,\(^4\) CQ caused 3D7 and 7G8 parasites to undergo loss of ΔΨ\(_m\) but to a lesser extent in 7G8 parasites, while the proportion of K1 parasites exhibiting loss of ΔΨ\(_m\) was unaffected (Figure 5a). In contrast, DSP, CPZ, PMZ, 4HT and LL-OMe were all shown to be effective in causing loss of ΔΨ\(_m\) in 3D7, 7G8 and K1 (Figure 5a). Notably, their effects were different between strains, but CQ-resistant 7G8 and K1 parasites were either equally or more sensitive to the effects of these lysosomotropic compounds than CQ-sensitive 3D7.

Similarly, CQ caused DNA degradation in all three strains of parasites, but to a lesser degree in K1 and 7G8 strains (Figure 5b). In comparison, DSP, CPZ, PMZ and 4HT were all effective in causing DNA degradation in all three strains (Figure 5b).

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**Figure 4** TEM of 3D7 parasites. (ai and ii) Untreated 3D7 parasites show clearly defined hemazoin crystals located within an intact DV membrane. (bi and ii) Parasites exposed to 3 μM of CQ for 4 h also show intact DV membrane but with less discrete hemazoin crystals and membrane whorls forming inside the DV. (ci–iii) Some of the treated parasites also show compromised DV membrane with DV material extruding into the cytoplasm. (d) Other CQ-treated parasites show DV rupture leading to hemazoin crystals being dispersed into the cytoplasm. Scale bar shows 0.5 μm for ×11 000 and ×18 000 magnifications, 0.2 μm for ×28 000 and ×36 000 magnifications, and 0.1 μm for ×56 000 magnification.
IC50 values for the other lysosomotropic compounds were 170 nM and 355 nM, respectively (Figures 7a–c), while the CQ for 3D7, 7G8 and K1 were determined to be 48 nM, determined to range from 5.024–27.563 these compounds than the CQ-sensitive strains. At this point, CQ-resistant strains appeared equally or more susceptible to vehicle control (NA), 30 CQ-sensitive 3D7 and CQ-resistant 7G8 and K1 parasites were exposed to a 150 and analysis by flow cytometry. (Figure 5 PCD features in parasites exposed to lysosomotropic compounds. CQ-sensitive 3D7 and CQ-resistant 7G8 and K1 parasites were exposed to a vehicle control (NA), 30 μM of CQ, 200 μM of DSP, 100 μM of CPZ, 200 μM PMZ, 150 μM of 4HT or 10 μM LL-OMe for 10 h before staining with JC-1 and Hoechst, and analysis by flow cytometry. (a) JC-1 staining shows mitochondrial outer-membrane permeabilization in treated parasites. Data represent means ± S.E. (A1 versus A2–A7, P < 0.001; B1 versus B2–B7, P < 0.001; C1 versus C3–C7, P < 0.001; N = 3). (b) Weak Hoechst staining shows parasites with sub-G1 DNA content after exposure to lysosomotropic compounds. Data represent means ± S.E. (A1 versus A2–A6, P < 0.001; B1 versus B2, P < 0.05; B1 versus B3–B6, P < 0.001, B1 versus B7, P = 0.053; C1 versus C2, P < 0.01; C1 versus C3–C6, P < 0.001; C1 versus C7, P = 0.072; N = 3).

These results demonstrated that the extent of CQ-induced PCD is dependent on PICRT mutations, whereas PCD induced by DSP, CPZ, PMZ and 4HT is unaffected by PICRT mutations and consistently triggers a significant degree of loss of Δψm and DNA degradation in all three strains.

Ca2+ redistribution in 3D7, 7G8 and K1 induced by CQ and other lysosomotropic compounds. Treatments with DSP, CPZ, PMZ or 4HT resulted in Fluo-4-AM redistribution from the hemazoin-containing parasite DV to the parasite’s cytoplasm in all three strains (Figures 6a–c), similar to what had been observed in Figures 2 and 3. This simultaneous occurrence of DV permeabilization and PCD features suggests that these are coupled events, and supports the hypothesis that leakage of clan CA cysteine proteases from the DV triggered the hallmark PCD features. There is insufficient data to address the causes or implications of these differences.

Discussion
We have previously reported evidence that micromolar levels of CQ initiate a unique PCD pathway that involves the activity of clan CA cysteine proteases, loss of Δψm, DNA fragmentation and cell death. This paper begins by further characterizing this pathway through the use of inhibitors to non-classical PCD mediators.

Even though some studies have shown that ROS and RNS exposure resulted in parasites with cell death characteristics, we were unable to detect their involvement in the CQ-induced PCD pathway. Although Ca2+ -chelating BAPTA-AM was able to reduce DNA degradation, experiments involving calpastatin were unable to show calpain involvement as had been proposed by others. Clan CD-specific cysteine protease inhibitor, Q-VD-OPh, was also ineffective in reducing PCD features in CQ-induced parasites even though the role of a parasite metacaspase (PIMCA1 – a Ca2+ -sensitive clan CD cysteine protease) had been proposed by others.

Notably, it was demonstrated for the first time that CQ-induced cell death had a transcriptionally regulated component – cycloheximide and actinomycin pre-treatment reduced the proportion of parasites showing sub-G1 DNA content. This suggests that the late stage of cell death may, at least in part, rely on transcriptionally dependent mechanisms.

Microarray data of CQ-treated parasites showed that a number of cysteine protease genes were indeed upregulated, including a putative clan CA pre-procathepsin c gene. However, knockout parasites were not more resistant to PCD, and these inconclusive results may suggest that redundancy factors are involved.

Recognizing that clan CA cysteine proteases are typically localized to the parasite’s DV and that they are associated with haemoglobin digestion, it was possible their redistribution to the cytoplasm may trigger/mediate CQ-induced PCD. This would require that the DV membrane becomes permeabilized and that DV-localizing moieties (such as Ca2+) to be released together with these proteases. Indeed, a simultaneous redistribution of both CQ and Ca2+ out of the DV was observed, but this only occurred at micromolar concentrations of CM-CQ. This was not surprising as CQ is known to be lysosomotropic, and its excessive accumulation in the parasite DV may have led to a destabilization of the DV membrane. Such permeabilization of lysosome membrane has also been demonstrated in mammalian cells exposed to micromolar concentrations of CQ.

The physiological relevance of Ca2+ redistribution in the regulation of P. falciparum and P. chabaudi development has been demonstrated previously by the Garcia group. It was shown that high levels of melatonin, a hormone regulated by the circadian cycle, resulted in phospholipases C activation, the generation of IP3, and a release of Ca2+ from the intracellular ER-stores, eventually leading to the loss of parasite synchronization. Moreover, the redistribution of Ca2+ was accompanied by an increase in mitochondria Ca2+ levels and an increase in the activity of cysteine proteases within the parasite’s cytoplasm. It would be interesting to...
study the possible role of Ca\(^{2+}\)-associated PCD in the in vivo synchronization of parasites that results in the characteristic cycle of fever and chills in afflicted patients.

To better understand the mechanics of the permeabilization, time-course experiments were performed and the study was extended to include CQ-resistant 7G8 and K1 parasite strains. Interestingly, DV destabilization was identified to occur in a majority of 3D7 and 7G8 parasites between 2 and 4 h after exposure to 3 \(\mu\)M of CQ, whereas only a minority of K1 parasites exhibited this. This is likely due to the effects of the PICRT in reducing CQ levels within the DV of K1 and thereby stabilizing it. As can be seen from the lower IC\(_{50}\) value, the PICRT of 7G8 is less efficient that in K1 and may not have adequately effluxed CQ to stabilize the organelle at such micromolar doses.\(^{27}\)

The early onset of DV destabilization in the majority of 3D7 parasites after only 2–4 h suggests that this feature may present before the loss of \(\Delta F_{\text{m}}\) which was previously shown to occur in only a small proportion of induced parasites 4–6 h after CQ administration.\(^{4}\)

Through the use of live cell imaging, the redistribution of Ca\(^{2+}\) in 3D7 was observed to occur in an abrupt and rapid manner. This supported a model where the accumulation of lysosomotropic CQ within the DV reached a threshold beyond which the DV membrane was permeabilized and its contents rapidly leaked into the parasite cytosol. However, this observation does raise concerns about the ‘programmed’ nature of cell death. Although an unregulated rupture of lysosomes has been shown to result in necrosis in mammalian cell, a regulated LMP has been demonstrated to result in cathepsin-mediated PCD.\(^{28–31}\) It was therefore important to closely examine the DV membrane to observe for membrane integrity.

Transmission electron microscopy (TEM) imaging of CQ-treated 3D7 did indeed show a small proportion of parasites with ruptured DVs. This has also been observed in \textit{P. falciparum} subjected to heat shock, and such cells are likely to be necrotic parasites.\(^{14}\) Some other cells had minute pores through which DV enzymes may have leaked into the cytoplasm and it cannot be ascertained whether these cells are undergoing regulated or unregulated cell death. A majority of cells, however, showed intact and clearly defined DV membrane. This is in line with other reports showing the cells undergoing LMP often have no observable ultrastructural differences in lysosome membrane morphology when compared with healthy cells.\(^{30}\) Given that the majority of CQ-treated parasites also show Fluo-4-AM redistribution to the cytosol following by PCD hallmarks, it is possible that these parasites have ‘regulated’ membrane permeability leading to Ca\(^{2+}\) release without visible membrane compromise and leading to a ‘regulated’ PCD pathway.

Although the process of lysosome-mediated cell death has been well studied, the exact mechanisms remain unclear.\(^{31}\) One model suggests apoptosis-associated LMP is size selective such that only smaller molecules are able to diffuse through the partially permeable lysosomal membrane while larger molecules remain trapped.\(^{32}\) If this model is applicable to \textit{P. falciparum}, it might allow specific clan CA proteases to exit the parasite DV (together with CM-CQ, and Ca\(^{2+}\)) and trigger PCD without jeopardizing the DV membrane.

**Figure 6** Effect of lysosomotropic compounds on Ca\(^{2+}\)-localization. Parasites were exposed to a vehicle control (NA), 3 \(\mu\)M of CQ, 200 \(\mu\)M of DSP, 100 \(\mu\)M of CPZ, 200 \(\mu\)M PMZ, 150 \(\mu\)M of 4HT for 4 h, or 10 mM LL-OMe for 10 h before staining with Ca\(^{2+}\)-sensing Fluo-4-AM and observation under a confocal microscope. (a) CQ-sensitive 3D7 and (b) CQ-resistant 7G8 show similar redistribution of Fluo-4-AM from the hemazoin-containing DV to the parasite’s cytoplasm after exposure to CQ, DSP, CPZ, PMZ and 4HT. (c) Multi drug-resistant K1 parasites, however, shows decreased redistribution after exposure to CQ but similar redistribution when exposed to the other lysosomotropic compounds. At least 30 infected erythrocytes were counted per condition.
Notably, this novel discovery of DV permeabilization in response to micromolar concentrations CQ is clinically relevant as serum levels of CQ actually do reach such concentrations.33–35 Although the main antimalarial mechanism of CQ lies in its inhibition of hemazoin formation, DV destabilization may account for the rapid clearance of parasites (especially those at the late-trophozoite and schizont stage) during the initial drug-loading phase of treatment.

To investigate whether PCD is CQ specific or truly a result of DV destabilization, other lysosomotropic compounds were assayed. Of the panel of compounds tested, DSP, CPZ, PMZ and 4HT all produced both early (loss of Dc_m) and late-stage (DNA degradation) hallmarks of PCD. Moreover, confocal microscopy confirmed that these compounds induced DV permeabilization after only 4 h treatment. Interestingly, these compounds have been shown to be strong lysosomotropic compounds that reduced LysoTracker Red staining and showed cytotoxicity in H9c2 rat cardiomyocyte-derived cells as well.20 The manner in which DV destabilization triggers cell death is also very similar to that described for metazoan cells which have been shown to involve LMP, loss of Dc_m (shown to be an indispensable component of the pathway) and caspase-independent or -dependent cell death execution.21–22

Notably, these compounds function equally well in inducing parasite PCD regardless of CQ sensitivity. This might be expected as the presence of PfCRT and other multi-drug resistant transporters are not expected to affect their accumulation inside the DV. Nevertheless, it should be cautioned that resistance might develop if compound-specific transporters evolve to efflux such drugs out of the DV.

Unlike CQ which primarily exerts its antimalarial activity through inhibiting hemazoin polymerization,36 these DV-destabilizing compounds are not known to bind with any specific target and, resultantly, their IC_{50} values are relatively high (6–30 μM) compared with that of CQ (48–355 nM). Though this limits their usefulness in antimalarial therapy, the high IC_{50} values of these compounds suggest that these compounds inhibit parasite growth through their lysosomotropic nature – these compounds exert their cytotoxic effects only when they have accumulated to sufficient levels in the DV to alter its biochemistry. That these compounds affect mammalian lysosomes at similar concentrations (3–40 μM) is also suggestive of this.20 As micromolar concentrations of these compounds are only sufficient to alter the pH of the DV and inhibit parasite development, a significantly higher dose is likely required to permeabilize the DV membrane. This explains why an approximately tenfold higher concentration (100–200 μM) is needed to induce the PCD features earlier observed.

Taken together, these studies on lysosomotropic compounds present for the first time, clear evidence that drug-induced DV destabilization can trigger PCD to occur similarly in both CQ-resistant and -sensitive P. falciparum. Although

![Figure 7](image_url)
these compounds cause lysosomal damage to mammalian cells at the same concentration that they induce DV destabilization in *P. falciparum*, analogues of these compounds or some other parasite DV-specific compounds may be developed to eventually become functional antimalarials.

In summary, we show that the later stages of the CQ-induced PCD pathway are both Ca$^{2+}$-dependent and transcriptionally regulated in *P. falciparum*. Additionally, it is demonstrated for the first time that selective permeabilization of the DV membrane occurs at high micromolar concentrations of CQ and is an important upstream event in the PCD pathway. This permeabilization is shown to be rapid, abrupt and can occur without rupture of the DV membrane, reminiscent of lysosome-mediated cell death in mammalian cells. As micromolar concentrations of CQ are only found in the plasma immediately after the initial loading dose is administered, this may be a clinically relevant yet uncharacterized antimalarial mechanism of the drug that will rapidly clear the mature stages of the parasite. Evidence that DV destabilization can be induced by other ‘non-antimalarial’ compounds was also presented, and DV permeabilization was associated with PCD features. Finally, CQ-sensitive and -resistant parasites were also shown to succumb to these lysosomotropic compounds with hallmark PCD features. This opens up new possibilities for the development of a novel class of drugs, which exerts their antimalarial effect through compromising the DV membrane’s integrity.

Materials and Methods

**Parasite culture.** A CQ-sensitive laboratory strain of *P. falciparum, 3D7* (MRA-102, MR4; ATCC, Manassas, VA, USA), and CQ-resistant strains 7G8 (MRA-154, MR4; ATCC) and K1 (MRA-159, MR4; ATCC), were cultured continuously as previously described.₄ Cultures of ~7–10% parasitemia were used, with parasites being in the late-ring stage (22–26 h after invasion) during the administration of inhibitor and drugs unless otherwise stated.

**Preparation of cell death inducers and treatments.** Chloroquine, diprophosphate, VP, diltiazem, DSP, CPZ, PMZ, propanolol and chlorpromazine (all from Sigma-Aldrich, Dorset, UK) were prepared fresh for each experiment by dissolving in PBS and filter sterilizing. Stock solutions of LL-OMe, norfloxacin, ciprofloxacin, mitofusine and vincristine (all from Sigma-Aldrich) were dissolved in DMSO and stored at ~20 °C. Stock solution of 4HT (Sigma-Aldrich) was dissolved in ethanol and stored at ~20 °C. CM-CQ was synthesized as previously described,₄ dissolved in DMSO and stored at ~20 °C. Unless otherwise stated, parasitized erythrocytes were incubated for 4 or 10 h in these inducers before being washed twice with MCM and assaying for apoptotic features. Vehicle controls consisted of equivalent amounts of DMSO or PBS.

**RNA preparation and reference pool.** Microarray oligonucleotide design, spotting, parasite RNA purification, cDNA synthesis, Cy Dye labelling and hybridizations were performed as previously described.₃ Samples per time-point were subjected to Cy Dye switching to eliminate any bias in labelling. Samples for individual time-points were hybridized against a reference pool that contained RNA samples representing all the developmental erythrocytic stages of the parasite. A total of 4 µg of sample RNA per time-point was hybridized against 4 µg of reference RNA were used for each microarray, and hybridizations were incubated at 60 °C for 16 h.

**Microarray acquisition and quality control.** Data from each time-point were acquired and analyzed on a GenePix Pro 4.0 (Axon, Sunnyvale, CA, USA), and the ratios normalized for the swapped Cy Dye replicates. The data were then reformatted using MEV (http://www.tm4.org/mev) and normalized using MIDAS (http://www.tm4.org/midas.html). Briefly, a scalar normalization factor was calculated for each array using unflagged features with median intensities >0 for each channel and a pixel resolution correlation coefficient >0.75. Quality spots were retained based on the log2 ratio for array features that were unflagged and had a sum of median intensities greater than the local background plus two standard deviations of the background were extracted from the database for further analysis. The ratios were then averaged for each time-point replicate.

**Real-time RT-PCR.** Genomic DNA contamination from purified total RNA was removed with the TURBO DNA-free Kit (Ambion, Austin, TX, USA) according to the manufacturer’s protocol. Amplification was performed using the Maxima SyBr Green 2X Master mix (Fermentas, Glen Burnie, MD, USA) and analyzed on an ABI 7500 thermocycler (Applied Biosystems, Carlsbad, CA, USA). Primer sequences for the candidate genes can be found in Supplementary Data (Supplementary Table S5). The absolute amount of transcripts were quantified relative to the standard curve and normalized to actin as a normalization housekeeping gene. The unpaired T-test was used to determine the significance of the differences between biological triplicates and a P-value of <0.05 was reported.

**Generation of PPC knockout.** The pARL-STEVORΔ3 vector was linearized at the BglII and AvrII sites to excise the existing STEVOR sequence. A fragment encoding the 5’-UTR together with the first exon of PPC gene (PFL2290w) was amplified using the PFL2290w_BglII (5’-AGTCAGATGGACTCATACATTTACGACCTGC-3’) and PFL2290w_AvrII (5’-AGTCCTACGTCGGTTAACTTTACAATTTCT-3’) primers to create the pARL-PPC targeting vector. Parasites were transfected with 100 µg plasmid DNA and drug resistant transfectants selected with 5 µM WR99210.₅

**Southern blotting.** The PPC probe was generated via PCR amplification using the PFL2290w_BglII and PFL2290w_AvrII primers (Supplementary Data, Supplementary Table S5) with 10 nM of dUTP-DIG (Roche, Indianapolis, IN, USA). In all, 3 µg of genomic DNA was digested with both EcoRV and AvrII restriction enzymes (New England Biolabs, Ipswich, MA, USA) before loading onto a 0.8% agarose gel. The size-fractionated samples were transferred onto a Zeta-Probe GT membrane (Bio-Rad, Hercules, CA, USA) via capillary transfer using 0.4 M NaOH, and overnight hybridization was performed at 42 °C. All prehybridization, stringency washes and colorimetric development were performed using the NBT/BCIP DIG-High Primer DNA Labelling and Detection Starter Kit (Roche) as per the manufacturer’s protocol.

**Inhibitor preparation and treatments.** Calpstatin was purchased from Calbiochem (La Jolla, CA, USA) and dissolved in PBS. Q-VD-OPh was purchased from SM-Biochemicals (Anaheim, CA, USA) and dissolved in DMSO and stored at ~20 °C. All other assayed cell death inhibitors were all purchased from Sigma-Aldrich and include BAPTA-AM, PBN, cycloheximide, actinomycin D and bafilomycin A1, all dissolved in DMSO and stored at ~20 °C, as well as L-glutathione and Mannitol that were prepared fresh each time and dissolved in PBS. Working solutions were obtained by diluting the stock concentration with MCM just before administration. Parasite cultures were preincubated with inhibitors for 30 min before drug treatment. The final inhibitor concentrations are as indicated in each study. Vehicle control consisted of equivalent amounts of DMSO and/or ethanol in accordance with solvent amounts used in various inhibitor treatments.

**Costaining with JC-1 and Hoechst.** The PPC probe was generated via PCR amplification using the PFL2290w_BglII and PFL2290w_AvrII primers (Supplementary Data, Supplementary Table S5) with 10 nM of dUTP-DIG (Roche, Indianapolis, IN, USA). In all, 3 µg of genomic DNA was digested with both EcoRV and AvrII restriction enzymes (New England Biolabs, Ipswich, MA, USA) before loading onto a 0.8% agarose gel. The size-fractionated samples were transferred onto a Zeta-Probe GT membrane (Bio-Rad, Hercules, CA, USA) via capillary transfer using 0.4 M NaOH, and overnight hybridization was performed at 42 °C. All prehybridization, stringency washes and colorimetric development were performed using the NBT/BCIP DIG-High Primer DNA Labelling and Detection Starter Kit (Roche) as per the manufacturer’s protocol.

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Inhibitory concentration (IC_{50}) determination. Parasitized cultures (ring stage) diluted with fresh erythrocytes and MCM to 2% parasitemia and 1.25% hematocrit were incubated with various concentrations of test substances for 48 h. To determine parasitemia, cultures were stained with Hoechst 33342 and analyzed by flow cytometry. Plotting of the dose–response curve as well as the computation of the IC_{50} was done with GraphPad Prism 5 demo version using a four-parameter logistic curve (variable slope).

Flow cytometric analyses. Cell numbers and fluorescence intensity after staining with JC-1 and Hoechst were assayed by a high-throughput flow cytometer (BD LSRII Special Order System; BD, San Jose, CA, USA). JC-1 stained was excited with 488 nm blue laser with green fluorescence being detected using 525/50BP filter (396 V), while the red fluorescence was detected using 570LP 585/42BP filter (350 V). For Hoechst stain, excitation was by 355 nm UV laser while detection utilized 450LP 450/50BP filter (380 V). Forward and side scatter parameters were adjusted (200 V and 154 V, respectively) to accommodate erythrocytes and to exclude any debris. Unstained controls as well as stained healthy 3D7 cultures were used to determine gates at the start of each experiment to ensure consistency. In single-cell tracking experiments, chambered glass slides were pre-transmssivity of 3.0%. These parameters were used in all experiments to ensure flow cytometry. Briefly, cells on a wet-mount were visualized under 30–50 parasitized erythrocytes were screened per condition, as stated in each experiment. Argon ion lasers tuned to 405 nm and 488 nm, respectively. Images were captured Confocal imaging was the original volume before viewing under a confocal microscope. At least 30–50 parasitized cells were counted for each condition as stated in each experiment.

Statistical analyses. All data shown are means ± S.E.M. Statistical difference was analyzed using ANOVA and post-hoc comparison using Tukey's test. Significantly different results (P < 0.05) were highlighted.

Intra-cellular calcium detection. Cell-permeable calcium-sensing dye, Fluo-4-AM (Molecular Probes), fluoresces green when bound with Ca^{2+} and was used to detect calcium storage in the DV or its release into the parasite cytoplasm. Subsequently, this stain also acted as a surrogate marker for DV permeabilization. A final concentration of 2 μM of Fluo-4-AM was added to parasitized erythrocytes and incubated at 37 °C for 30 min. Cells were washed twice with PBS and resuspended to the original volume before viewing under a confocal microscope. At least 30–50 parasitized cells were counted for each condition as stated in each experiment.

Confocal microscopy and single-cell tracking. Confocal imaging was done using Olympus Fluoview FV1000 (Tokyo, Japan) equipped with solid state and Argon ion lasers tuned to 405 nm and 488 nm, respectively. Images were captured using Olympus Fluoview version 1.6b. For enumeration purposes, images of at least 30–50 parasitized erythrocytes were screened per condition, as stated in each experiment. Briefly, cells on a wet-mount were visualized under ×100 magnification and × 5 zoom with the following calibrations: 2.0 μm/Pixel sampling speed, line Kalman integration, 488 nm laser (for Fluo-4-AM) at 713 V and transmissivity of 10.0%, 405 nm laser (for CM-CQ and Hoechst) at 655 V and transmissivity of 3.0%. These parameters were used in all experiments to ensure consistency. In single-cell tracking experiments, chambered glass slides were pre-treated for 5 min with 1: 8 dilution of poly-l-lysine (Sigma-Aldrich) and left to dry for 1 h at 60 °C. Cultures that were prestained with Fluo-4-AM were diluted to a final hematocrit of 0.1% and 1 ml was added per chamber. A vehicle control or 3 μM of CQ was then administered while the cells were given time to settle to the base of the chamber. The viewing stage was perfused with standard parasite culture gas mixture and kept at 37 °C for the entire period of viewing. At 2 h after CQ was added, micrographs were captured every 1 or 3 min for the next 2 h, using a reduced laser transmissivity of 1.0% to reduce photobleaching. Fluorescence intensity tracing of parasite cross-sections were carried out using ImageJ 1.44p software (National Institutes of Health, Bethesda, MD, USA).

Transmission electron microscopy. Erythrocyte-free parasites were first isolated by saponin enrichment as previously described and washed twice in cold PBS. Cells were then fixed 24 h in 2% glutaraldehyde (w/v PBS) at 4 °C. After washing three times in cold PBS, cells were post-fixed for 1 h with 1% osmium tetroxide (w/v PBS) with 1% potassium ferricyanide followed by dehydration with a graded ethanol series. Thereafter, cells were infiltrated and embedded with LR-White resin (London Resin Company, London, UK) at 60 °C for 4 days and ultrathin sections (70–90 nm thickness) were prepared using UltraCut E microtome (Reichert-Jung, Wein, Austria). These sections were then collected on copper mesh grids, stained a second time with 1% osmium tetroxide for 5 min, rinsed, treated with uranyl acetate for 6 min, rinsed, and in lead citrate for 8 min before final rinsing. Sections were then examined using an EM208S transmission electron microscope (Phillips, Eindhoven, The Netherlands) and DigitalMicrograph software v1.82.365 (Pleasanton, CA, USA).

Conflict of interest

The authors declare no conflict of interest.

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