A programmed cell death pathway in the malaria parasite Plasmodium falciparum has general features of mammalian apoptosis but is mediated by clan CA cysteine proteases

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Several recent discoveries of the hallmark features of programmed cell death (PCD) in Plasmodium falciparum have presented the possibility of revealing novel targets for antimalarial therapy. Using a combination of cell-based assays, flow cytometry and fluorescence microscopy, we detected features including mitochondrial dysregulation, activation of cysteine proteases and in situ DNA fragmentation in parasites induced with chloroquine (CQ) and staurosporine (ST). The use of the pan-caspase inhibitor, z-Val-Ala-Asp-fmk (zVAD), and the mitochondria outer membrane permeabilization (MOMP) inhibitor, 4-hydroxycoumarin, enabled the characterization of a novel CQ-induced pathway linking cysteine protease activation to downstream mitochondrial dysregulation, amplified protease activity and DNA fragmentation. The PCD features were observed only at high (μM) concentrations of CQ. The use of a new synthetic coumarin-labeled chloroquine (CM-CQ) showed that these features may be associated with concentration-dependent differences in drug localization. By further using cysteine protease inhibitors z-Asp-Glu-Val-Asp-fmk (zDEVD), z-Phe-Ala-fmk (zFA), z-Phe-Phe-fmk (zFF), z-Leu-Leu-Leu-fmk (zLLL), E64d and CA-074, we were able to implicate clan CA cysteine proteases in CQ-mediated PCD. Finally, CQ induction of two CQ-resistant parasite strains, 7G8 and K1, reveals the existence of PCD features in these parasites, the extent of which was less than 3D7. The use of the chemoreversal agent verapamil implicates the parasite digestive vacuole in mediating CQ-induced PCD.

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The most deadly of the four clinically relevant malaria parasites, Plasmodium falciparum, continues to kill an estimated one million people annually.¹ Although current drugs can combat the disease, the increase in resistance to these medicines makes it crucial for new leads to be uncovered, such that new therapeutic initiatives will be one step ahead of disease progression.² One pathway yet to be thoroughly explored for antimalarial development is that of programmed cell death (PCD).

Recent reports highlight the existence of PCD features in unicellular organisms including Blastocystis, Leishmania, trypanosomes, Saccharomyces cerevisiae and Trichomonas vaginalis.³ Indeed, accounts of PCD in P. falciparum⁴ and the rodent malaria parasite P. berghei⁵ have been documented, as featured by the fragmentation of DNA and a loss of mitochondria membrane potential. However, other reports suggest that P. falciparum does not undergo PCD.⁶⁻⁷

Besides inconclusive cell death features in the malaria pathogen, there is a lack of evidence for a clear pathway linking such features. In addition, PCD pathway mediators have yet to be identified. Some studies have made use of caspase inhibitors or substrates to suggest the involvement of clan CA cysteine protease mediators such as metacaspase-orthologs⁸ or caspase 3-like proteases.⁹ These assays, however, have been shown to be nonspecific and do not reliably discriminate between the clan CA and CD cysteine proteases.¹⁰⁻¹¹

Our study began by characterizing the PCD pathway of P. falciparum. First, we conducted three cell-based assays and detected mitochondrial dysregulation, cysteine protease activation and DNA fragmentation in both CQ- and
ST-induced parasites. Second, the mitochondria outer membrane permeabilization (MOMP) inhibitor, 4-hydroxytamoxifen, and pan-caspase inhibitor, z-Val-Ala-Asp-fmk (zVAD), were used to determine the order in which these PCD features occurred. Interestingly, the sequence of events in CQ-induced parasites showed a striking resemblance to the classical mammalian apoptotic pathway even in the absence of classical apoptotic machinery. Third, owing to the cross-reactive nature of fluoromethylketone (fmk) inhibitors, clan CA-specific protease inhibitors E64d and CA-074 were used and showed that clan CA cysteine proteases are crucial mediators in the CQ-induced PCD pathway. Clan CA proteases, in particular the cathepsins and calpains, have already been implicated in protozoan PCD.12–15 As such, the discovery of clan CA proteases as PCD mediators in P. falciparum presents a clear new target of therapeutic intervention for future antimalarial design. Finally, the existence of CQ-induced PCD was shown in drug-resistant strains 7G8 and K1, and the importance of CQ’s lysosomotropic nature in triggering these features is discussed.

Results

Mitochondrial membrane potential assay. The accumulation of JC-1 J-aggregates in uninduced cells showed that about 90% of the parasites contained functional mitochondrion with transmembrane potential ($\Delta V_{m}$) (Figure 1a and c). Confocal microscopy confirmed the specificity of the aggregated mitochondrial stain (Figure 1b). Treatment with CQ or ST caused a significant decline in this proportion to approximately 69 and 75%, respectively (Figure 1a).

Pretreatment of parasite culture with 4-hydroxy-tamoxifen (4HT) inhibited CQ-induced MOMP almost entirely, returning the proportion of JC1-positive cells to ~90% (Figure 1a). However, 4HT was unable to exert a similar protective effect for ST-induced parasites, suggesting that ST-induced decrease in JC-1 values was independent of MOMP.

Pan-caspase inhibitor, zVAD, was similarly able to rescue CQ-induced cells from loss of $\Delta V_{m}$ but was unable to protect cells from ST-induced JC-1 loss (Figure 1a). Almost 90% of CQ-induced cells pretreated with zVAD were JC1-positive compared with 77% of ST-induced cells. This suggests that the activity of cysteine proteases is upstream of MOMP in CQ-induced cells.

The use of 4HT and zVAD in combination did not rescue ST-induced cells from JC-1 loss, whereas CQ-induced cells were completely rescued (Figure 1a).

Cysteine protease activity. In parasitized erythrocytes, high background staining within the erythrocyte (data not shown) deemed it necessary for parasites to be enriched by saponin-lysis of the host erythrocyte such that fluorescence of the parasite itself could be clearly recorded. In uninduced parasites, a basal level of CaspaTag activity was detected in 10% of the cells (Figure 2a and c). This proportion increased significantly on CQ and ST induction to 34 and 32%, respectively. The retention of CaspaTag in a greater proportion of CQ- and ST-induced parasites suggested that the pan-caspase stain was bound to cysteine proteases that had been activated by the drug treatments.

Preincubation of the parasite culture with zVAD reduced the basal proportion of uninduced cells that are CaspaTag-positive to 5% and suppressed CQ-induced CaspaTag binding to below 10% (Figure 2a). Interestingly, zVAD did not reduce the proportion of CaspaTag-positive cells after ST treatment.

Even in the absence of CQ and ST, 4HT increased the proportion of CaspaTag-positive cells slightly to 17% (Figure 2a). Moreover, 4HT increased the proportion of CaspaTag-positive cells in ST-treated cultures from 32 to 80%, suggesting synergism between 4HT and ST on the parasite. This synergistic effect was not preventable by zVAD treatment. Despite this inherent toxicity, 4HT reduced CaspaTag binding in CQ-induced cells from 34 to 15%, showing clearly that MOMP is essential for cysteine protease activity in CQ-induced parasites.

In situ DNA fragmentation. As with the CaspaTag assay, parasitized cultures treated with various conditions were saponin-enriched just before TUNEL staining and gated in an identical manner. There was an increase in the proportion of TUNEL-positive cells from 10 to 27% and 56% in parasites induced by CQ and ST, respectively, indicating that these two drugs induced in situ DNA fragmentation in parasites (Figure 3a and c).

Pretreatment of parasite cultures with 4HT alone resulted in an increase in TUNEL-positive cells to 22.5% (Figure 3a). Moreover, 4HT was ineffective in reducing the percentage of TUNEL-positive cells in both CQ- and ST-treated cultures, possibly owing to its inherent toxicity.

Pretreatment with zVAD, however, significantly reduced the TUNEL-positive population in CQ-induced but not in ST-induced cultures (Figure 3a). This suggested that zVAD was able to effectively inhibit the pathway leading to in situ DNA fragmentation in CQ-induced parasites but not in ST-induced populations.

In double inhibition studies, the proportion of CQ-induced TUNEL-positive cells decreased but the toxic effects of 4HT were still observable in the non-induced cultures (Figure 3a). ST-induced cultures remained highly TUNEL-positive even when inhibitors were used in combination.
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CQ + 4HT
CQ + zVAD

NA

Counts

R1  9.68%

Counts

R2  10.05%

Counts

R1  3.50%

Counts

R2  5.56%

Counts

R1  5.71%

Counts

R2  7.05%

Counts

R1  5.96%

Counts

R2  6.43%

Counts

Field 1

Counts

Field 2

Counts

Field 3

Counts

Field 4

Percentage of JC-1 Positive Parasites (%)

Veh  4HT  zVAD  4HT + zVAD

Counts

Phase  JC1 (Red)

Counts

JC1 (Green) Overlay

A  B  C  D  E  F

R1  R2

JC1 (Red) Log

JC1 (Green) Log

JC1 (Red) Log

JC1 (Green) Log

JC1 (Red) Log

JC1 (Green) Log

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As CQ-induced parasites displayed clearer PCD induction and inhibition in MOMP, cysteine protease activity and in situ DNA fragmentation assays, we focused on the characterization of the CQ-induced pathway and its mediators.

**Time course experiments with CaspaTag and JC-1.** Time course experiments on CQ-induced parasites showed that a significant decrease in JC-1-positive cells occurred after 4 h of drug administration in contrast to the increase in CaspaTag-positive cells that was significant only after 6 h (Figure 4). This suggests that mitochondrial dysregulation is an early-onset feature of PCD, which precedes the activation of cysteine proteases. As it was also shown that zVAD is able to prevent MOMP, the activity of cysteine proteases must be essential for the loss of ΔΨm to take place. We therefore posit that a small increase in protease activity leads to mitochondrial permeabilization, which then amplifies the protease cascade in a positive feedback loop.

**Preparation and validation of fluorescent-labeled chloroquine.** To determine the localization of chloroquine (CQ), we synthesized a CQ analog conjugated to biologically benign coumarin (CM) fluorophore. HPLC and NMR analyses indicated that coumarin-labeled chloroquine (CM-CQ) was synthesized with excellent purity of 98% (Supplementary data, Figure S1–13). To ascertain the biological efficacy of CM-CQ, IC50 curves were measured and the inhibitory efficiencies of the CQ compounds determined (Supplementary data, Figure S1A). In addition, the effect of varying CM-CQ concentrations on JC-1 assay was also measured (Supplementary data, Figure S2). These assays show that CM-CQ behaves similar to CQ in parasite killing and inducing MOMP, but with about 10-fold less efficacy.

**Dose-dependent effect of CQ.** JC-1 and CaspaTag assays were used to investigate PCD features at different
concentrations of CQ (Figure 5a). At low nanomolar concentrations, there were basal levels of PCD features but at micromolar concentrations of CQ, there was a large increase in the proportion of cells displaying MOMP and activation of cysteine proteases. Confocal microscopy revealed a dose-dependent relocalization of labeled CQ from the DV at nanomolar concentrations to the cytoplasm at micromolar concentrations (Figure 5b and Supplementary Data, Figure S3).

**Necrosis control experiments.** To exclude the possibility that the CQ-induced PCD is an unregulated necrotic death, parasitized cultures were subjected to sodium azide and 50°C treatments in the presence or absence of zVAD
To verify the increase in the proportion of CQ-induced parasites mediating by clan CA cysteine proteases, the pathway. Cathepsin B-specific CA-074 was unable to inhibit these features. Heat treatment altered JC-1 background staining of the erythrocytes and therefore data were excluded.

Effect of different inhibitors on CaspaTag binding. Owing to the well-documented nonspecific binding of zVAD to both clan CD and CA proteases, coupled with the absence of any caspase-orthologs in *Plasmodium*, the range of inhibitors was extended to elucidate the preferred substrate specificity of the target protease. Besides CA-074, all the tested inhibitors prevented the increase of CaspaTag-positive cells in CQ-induced samples (Figure 6a). Inhibition by clan CA-specific inhibitor E64d showed that target proteases were clan CA cysteine proteases. Inhibition by z-Phe-Ala-fmk (zFA), z-Phe-Phe-fmk (zFF) and z-Leu-Leu-Leu-fmk (zLLL) showed that target proteases lack the P1-aspartate specificity characteristic of caspases, and suggests the involvement of clan CA proteases such as cathepsins and calpain-like proteases in the pathway. Cathepsin B-specific CA-074 was unable to fully prevent CaspaTag binding, suggesting that cathepsin B-like proteases are partly involved in mediating the pathway.

Hoechst staining of DNA content. To verify the effectiveness of the extended panel of inhibitors, parasites were assayed for a loss of DNA content 10 h after CQ treatment. Healthy ring-stage parasites were used to distinguish the G1 population (Supplementary data, Figure S7A) and a control experiment excluded the possibility that CQ competed with Hoechst DNA-binding (Supplementary data, Figure S7C). Finally, flow cytometry histograms and fluorescence microscopy images (Supplementary data, Figure S7B and D) were compared to characterize parasite cultures that had been subject to various experimental conditions.

CQ-treated parasites showed an increased proportion of cells with sub-G1 DNA content from 10 to over 20% (Figure 6b). The effect of different inhibitors corroborated findings of the previous section, suggesting that clan CA proteases are essential in mediators for DNA degradation in CQ-induced PCD and that cathepsin B-like proteases act in concert with other clan CA mediators to give rise to DNA breakdown.

Intracellular calcium studies. In healthy parasites, calcium indicator fluo-4-AM localized to the digestive vacuole (DV) of healthy parasites (Supplementary data, Figure S8). However, CQ treatment resulted in a redistribution of Ca$^{2+}$ to the cytoplasm in a dose-dependent manner (Supplementary data, Figure S8 and S9). To explore the possibility that MOMP is Ca$^{2+}$-mediated, a cell-permeable calcium chelator, BAPTA-AM, was used but this did not significantly reduce the proportion of CQ-induced parasites undergoing MOMP (Supplementary data, Figure S10).

Inter-strain comparisons for JC1 and Hoechst loss. We subsequently assayed for PCD features in CQ-resistant 7G8 and K1 strains (Figure 7). The IC$_{50}$ values of CQ in 7G8 and K1 parasites were determined to be 76 and 202 nM, respectively (Supplementary Data, Figure S1B and S1C). Pretreatment with VP lowered the IC$_{50}$ values of 7G8 and K1 to 61 and 120 nM, respectively. Interestingly, 7G8 cultures mirrored 3D7 and showed a significant drop in the JC1-positive population at concentrations of CQ exceeding 3 μM of CQ (Figure 7a). In comparison, the drop in JC1-positive K1 cells was markedly less at this same concentration, and it was only at a higher concentration of the drug that there was a significant decrease in the proportion of JC1-positive cells. However, pretreatment of K1 with 10 μM of chemoreversal agent VP resulted in a significant decrease in JC1-positive cells even at 3 μM of CQ.

As with 3D7, both 7G8 and K1 showed a significant increase in the proportion of parasites with sub-G1 DNA content after CQ treatment (Figure 7b). However, these proportions were significantly lower than those observed in 3D7. Although VP pretreatment was able to significantly increase the proportion of K1 parasites with sub-G1 DNA content, this reversion was incomplete. It should be noted that BD LSRII showed better resolution of Hoechst-positive parasites from the uninfected RBC population, and consequently the proportion of cells showing DNA loss as measured by this cytometer was slightly greater than that measured in the previous set of experiments using DAKO cytometry.

Discussion

Our results show that both CQ and ST treatment gives rise to features of PCD in *Plasmodium* including MOMP, the activation of cysteine proteases and *in situ* DNA fragmentation. These findings corroborate growing evidence supporting the existence of PCD hallmarks in parasites induced by a variety of agents including drugs, febrile temperatures, exposure to bilirubin resulting in reactive oxygen species increase and binding of platelets to infected erythrocytes.

Observation of apoptotic features such as chromatin condensation, DNA fragmentation, phosphatidylserine exposure, apoptotic body formation and caspase activity have been shown by Al-Olayan et al. in the mosquito midgut stage.
of *P. berghei*. However, this could not be reproduced by Le Chat *et al*.22 In apoptotic *P. falciparum*, as induced by drugs and heat-shock treatments, Nyakeriga *et al.*6 and Totino *et al.*7 were unable to detect MOMP, DNA fragmentation and laddering, or chromatin cleavage. In contrast, Meslin *et al.*8 and Kumar *et al.*9 showed MOMP, DNA fragmentation, chromatin condensation and DEVDase activity in other strains of *P. falciparum* induced by similar stimuli. Still other authors observed evidence of autophagy or secondary necrosis including cytoplasmic vacuolation in drug-resistant strains such as PSS1 or a swelling of the food vacuole in CSC-1.7,19 These discrepancies may be due to differing drug concentrations, parasite strains, developmental stages, inducers and assaying methods, complicating the conclusions that can be reliably discerned.

Another limitation of the existing studies has been the absence of definitive evidence outlining any pathway of PCD. As unicellular eukaryotes may display more than one PCD pathway,23 the focus of this study was primarily to delineate the pathway induced by CQ.
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Several established MOMP inhibitors were investigated (data not shown) and 4HT was shown to inhibit MOMP in CQ-induced cells without significantly reducing the proportion of healthy cells in the control group. Current evidence suggests that 4HT inhibits Ca$$^{2+}$$-induced MOMP in ischemic myocardiocytes by binding to Ca$$^{2+}$$-activated K$$^+$$ channels of the mitochondria.24–25 However, this is not likely to be the mechanism of action in the parasite as preincubation of parasites with the Ca$$^{2+}$$-chelator BAPTA-AM did not prevent MOMP. In addition to 4HT, the effectiveness of zVAD in inhibiting MOMP in CQ-induced cells points clearly toward the activity of cysteine proteases being upstream of MOMP.

Despite some toxicity, 4HT was also able to partially reduce the major proportion of CaspaTag-positive cells in CQ-treated cultures, suggesting that MOMP occurs upstream of the bulk of cysteine protease activation in CQ-treated cells. This conclusion was supported by time course experiments that showed MOMP occurring in a significant proportion of cells at 4 h after CQ induction, whereas the increase in CaspaTag-positive cells was only significant after 6 h.

Pretreatment with 4HT was unable to prevent CQ-induced DNA fragmentation. Instead, 4HT toxicity made it difficult to interpret findings. In contrast, zVAD pretreatment clearly reduced TUNEL staining in CQ-induced cells, placing the activation of cysteine proteases upstream of *in situ* DNA fragmentation.

The effectiveness of zVAD's inhibition of cysteine proteases, leading to a concomitant change in the proportion of CQ-induced parasites showing MOMP, cysteine protease activity and DNA fragmentation, is strongly indicative of a regulated pathway linking these PCD features. In addition, the effect of 4HT in reducing CaspaTag-positive cells and time point experiments provides further insight into the CQ-induced pathway outlined below (Diagram 1b). Our model of CQ-induced PCD suggests that the activity of cysteine proteases is necessary for the loss of mitochondrial membrane potential. This preceding event is likely to be involved in...
the amplification of protease activity, which results in DNA fragmentation. As experiments with 4HT were complicated by its toxicity, we cannot exclude the possibility that this amplification of the cysteine proteases can occur independently of mitochondrial permeabilization.

Our CQ-induced PCD model resembles the classical model of receptor-mediated apoptosis, which was recently reviewed by Degterev and Yuan. Although caspase 8 is able to directly activate executioner caspase 3 to initiate apoptosis, this process is greatly amplified through the mitochondrial route (Diagram 1a).

Although these models display overall similarities in the assayed characteristics, the absence of caspase homologs and classical mediators of apoptosis, such as Bid, Bax, AIF,
Apaf-1 and Bcl-2 proteins, alludes to different mediators being involved in the process of CQ-induced PCD in *P. falciparum*. In ST-induced cells, the ineffectiveness of zVAD and 4HT in reducing PCD features suggests that a different phenomenon is being observed. The high concentrations of ST may have resulted in unregulated necrosis that is unlikely to be prevented by any number of inhibitors.

An apparent limitation of this study is the high concentrations of CQ used. In an attempt to consolidate competing findings by different studies using different drug concentrations (35 nM–100 μM), parasites in this study were initially exposed to a range of drug concentrations (30, 300 nM, 3 and 30 μM) of CQ for 8 h and observed for features of PCD. To ensure that parasites were all still in a comparable trophozoite stage of development, an incubation time exceeding 10 h was not considered. As a significant loss of ΔΨm and an increase in CaspaTag-positive cells were observed only with 3 and 30 μM of CQ treatment, the latter dose was used. These parameters worked optimally with our protocols and allowed the investigation of multiple PCD phenotypes at a single time point. Moreover, effective zVAD-mediated inhibition of PCD features even at such high doses of CQ suggests that key modulators of a well-regulated pathway are indeed being targeted.

Conclusions on studies using micromolar concentrations of CQ must not be projected as the primary mechanism of CQ’s antimalarial activity. At physiological nanomolar concentrations, lysosomotropic CQ accumulates in the DV and interferes with hemazoin formation. However, our findings showed the drug being present in the cytoplasm of the parasite at micromolar concentrations where it could bind nonselectively to unknown cytosolic PCD mediators. Alternatively, relocation of the drug at a high dose could be due to the excessive accumulation of the lysosomotropic drug in the DV leading to its compromise, the acidification of the cytoplasm and the accumulation of protonated CQ inside the entire parasite.

To exclude the possibility that the proposed CQ-induced PCD pathway is an unregulated necrotic event, two necrosis controls were included. Both sodium azide and heat treatments resulted in distinctly different cell death profiles compared with CQ-induced cultures. Moreover, it was clearly shown that zVAD was unable to modulate the extent of cell death features, strongly suggesting that these necrosis controls were genuinely undergoing an unregulated and disordered cell death. The ineffectiveness of zVAD in limiting necrotic cell death has been previously shown and contrasts with the well-ordered ‘programmed’ pathway characterized in CQ-induced cells where downstream features of cell death can clearly be inhibited by 4HT and zVAD.

Although the concentration of the CQ used limits this study’s usefulness in understanding the therapeutic mechanisms of CQ’s antimalarial activity, the objective of discovering novel mediators of *P. falciparum* PCD is not undermined. As further research characterizes what takes place in the parasite at such a dose of CQ, novel compounds specifically targeting the mediators of this PCD pathway may be optimally designed for antimalarial use.

Unlike the mammalian model, the identities of participating cysteine proteases in CQ-induced *P. falciparum* remains unknown, given the conspicuous absence of classical apoptotic mediators. Meslin et al. have suggested that a clan CD protease PfMCA1 (a metacaspase homolog) was a likely mediator of cell death in *P. falciparum*. However, Le Chat et al. showed that *P. berghei* metacaspases did not have a role in PCD. There is growing evidence that caspase inhibitors with fmk or aldehyde reactive groups (including CaspaTag) are not as specific as once believed, but can efficiently inhibit non-caspase cysteine proteases including metacaspases, calpains and cathepsins. This nonspecific nature of zVAD binding in our study is confirmed by the fact that other fmk moiety-containing inhibitors with different amino-acid sequences (z-Asp-Glu-Val-Asp-fmk (zDEVD), zFA, zFF and zLLL) seemed equally effective in reducing CaspaTag binding. Moreover, zFA, zFF and zLLL do not have the aspartate residue at the P1 position that is essential for classical caspase binding. Although the nonspecific nature of zVAD binding has allowed us to characterize the aforementioned pathway, we cannot ascertain the identity of its target protease on the basis of its binding to zVAD alone. Thus, clan CA-specific inhibitors such as E64d and CA-074 were included as well.

Despite being a broad-acting inhibitor of clan CA cysteine proteases, E64d does not cross-react with clan CD proteases. As E64d was effective at inhibiting CaspaTag binding and DNA loss in CQ-induced cells, we concluded that clan CA proteases are essential mediators of CQ-induced PCD. In addition, as CaspaTag binds both Clan CA and CD proteases, and Clan CA-specific E64d completely prevented the increase in CaspaTag-positive cells, this suggests that clan CA mediators are likely to be the proteases accounting for the cysteine protease activity being detected. The partial inhibition of PCD features by specific cathepsin B inhibitor, CA-074, shows that there may be multiple pathways involving clan CA proteases. Nevertheless, it would be premature to exclude the involvement of clan CD proteases such as PfMCA1 as regulators or even key downstream mediators of CQ-induced PCD.

Conventionally, Clan CD caspases mediate apoptosis. However, recent work highlights the importance of clan CA papain-like proteases such as calpains and cathepsins in PCD regulation through truncation of Bid or activation of Bax, leading to MOMP. Even though Bid and Bax homologs do not exist in the parasite, it is conceivable that one or more parasite clan CA proteases may function likewise to cleave a novel parasite-specific mediator of MOMP. Clan CA proteases are well associated with parasites and include cathepsins and falcipains from the C1 family and calpains from the C2 family. *P. falciparum* 3D7 possesses a preprocathepsin C homolog, a calpain homolog, four falcipains, SERA (serine-repeat antigens) and other uncharacterized C1 proteases.

Interestingly, a study by Gazarini et al. showed that micromolar quantities of CQ triggered calcium release from acidic compartments of *P. chabaudi*. Moreover, Farias et al. observed that calcium release was accompanied by a detection of cysteine-protease activity in the cytoplasm of *P. chabaudi* and *P. falciparum*. Although the authors suggest calpain involvement, this is unlikely as putative calpain...
homologs in *Plasmodium* do not have a calcium-binding domain. Our own studies on 3D7 also showed a relocalization of Ca\(^{2+}\) from the DV of the parasite to the cytoplasm. However, the use of the calcium chelator, BAPTA-AM, did not inhibit MOMP in CQ-induced parasites. However, the role of Ca\(^{2+}\) in limiting other features of CQ-induced PCD has yet to be fully investigated and it is still possible that in CQ-induced PCD, cysteine protease are activated in a Ca\(^{2+}\)-dependent mechanism as these authors have suggested.

This CQ-induced PCD pathway would only be of interest to drug developers if its existence were also established in CQ-resistant strains of *P. falciparum*. As such, drug-resistant 7G8 and K1 were also investigated for features of PCD. CQ resistance was initially believed to arise from mutations in the DV transmembrane protein CQ-resistance transporter (PICRT). Two distinct sets of mutations were found, one originating from South America and the other from Southeast Asia and Africa. These mutations in PICRT allow the efflux of CQ from its primary site of action, the DV, thereby conferring resistance to the drug. Interestingly, this resistance, originating from the Asia-Africa mutations, was shown to be reversible by VP and similarly structured compounds.

Despite the differences between 7G8 and K1 in PICRT mutation sets, both strains showed a significant proportion of cells with JC1-1 loss and DNA degradation, suggesting that PCD pathways exist even in these resistant strains. Interestingly, 7G8 showed a similar extent of cells exhibiting MOMP as 3D7 but a lower proportion of cells with sub-G1 content, even at higher CQ concentrations. In comparison, K1 showed a VP-reversible reduction in the proportion of cells with MOMP and DNA degradation that was only detectable at higher CQ levels. These findings show the importance of PICRT and CQ’s localization in the DV for PCD to occur (Diagram 1b).

In addition, the lesser extent of PCD features even at high CQ concentrations suggests that these CQ-resistant strains may have a reduced propensity to undergo PCD. This model for drug resistance has also been established in cancer cells where treatment pressure leads to a selection of clones with altered apoptotic machinery or regulatory components. Such cells are more likely to survive and may be cross-resistant to other drugs using the apoptotic pathway. Although further work on field isolates and other PCD inducers are needed, the potential of finding a link between drug resistance and resistance to cell death in *P. falciparum* would be very illuminating.

In summary, our present study not only shows that CQ and ST treatments give rise to characteristic features of PCD, but also a linear pathway implicating mitochondrial dysregulation, clan CA cysteine protease activity and DNA fragmentation is evident after CQ-induction. Although no specific mediators have been identified at this juncture, we have determined clan CA proteases as key pathway modulators. In addition, we have shown that PCD features exist in drug-resistant 7G8 and K1 parasites, and are suitable for therapeutic exploits. Besides providing promising targets for therapeutic drugs, breakthroughs in this area of PCD modulators should also enable a greater understanding of drug resistance and the physiological role of PCD pathways in unicellular organisms.

### 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 with minor modifications to the methods previously described. Parasite cultures were maintained in 75 or 125 cm\(^2\) flasks using malaria culture media (MCM) consisting of RPMI 1640 supplemented with 0.5% (w/v) albumin (Intralipid, Auckland, New Zealand), 0.005% (w/v) hygroxyline, 0.03% (w/v) L-glutamine, 0.22% (w/v) gentamycin and 2.5% human erythrocytes. Flasks were gassed with 3% CO\(_2\), 4% O\(_2\) and 93% N\(_2\) and incubated at 37°C in a dark incubator. Culture medium was replenished during subculturing on alternate days. Thun Giemsa smears were made to determine parasitemia before subculture and at the start of each experiment, Sorbitol synchronization was carried out weekly to select for ring-stage parasites.

#### Drug preparation and treatments

CQ diphosphate (Sigma-Aldrich, Dorset, UK) was dissolved in PBS and filter-sterilized to obtain working solutions of 300 μM, and stored in the dark at 4°C. A new batch of CQ working solution was made before each experiment. Unless otherwise stated, parasitized erythrocytes were incubated for 8 or 10 h in 30 μM of CQ and washed twice with MCM before assaying for apoptotic features. Staurosporine (ST, Sigma-Aldrich) stock solution (1 mM) was prepared by dissolving the drug in 1% DMSO and stored at −20 °C. Working concentrations of ST at 100 μM were prepared before each experiment by diluting the stock solution with MCM. Infected erythrocytes were treated for 8 or 10 h with 10 μM ST, and washed twice with MCM before cell death assays. For necrosis controls, cultures were incubated with either 0.1% (w/v) of sodium azide for 8 or 10 h, or heat treated at 50°C for 1 h, and then incubated at 37°C for 8 or 10 h. Verapamil (VP, Sigma-Aldrich) was dissolved in PBS with 0.5% acetic acid, to a working concentration of 1 mM, and parasite cultures were pretreated with 10 μM of VP for 30 min before adding CQ. Vehicle controls consisted of equivalent amounts of DMSO or acetic acid.

#### Synthesis of CM-CQ

Under a standard synthetic procedure, using flame-dried glass apparatus under nitrogen atmosphere, des(N-ethyl)-CQ (25.0 mg, 0.09 mmol), that is, N-(7-chloro-4-quinolinyl)-N’-ethyl-1,4-diaminobutane, was dissolved in anhydrous acetonitrile (5.0 ml). Finely powdered and predried potassium carbonate (24.8 mg, 0.2 mmol) was then added, followed by the addition of an anhydrous acetonitrile solution (1.0 ml) of the known CM (44.2 mg, 0.1 mmol), that is, 2-bromo-5-(2-[(7-dimethylamino)-2-oxo-2H-chromen-4-yl]acetamido)-ethyl)-acetamide. The mixture was stirred for 12 h at room temperature, then filtered and the filtrate evaporated to give a yellow residue. The residue was then purified by flash column chromatography (230–400 mesh silica gel) to give CM-CQ (45.0 mg, 0.074 mmol) in 82% yield.

NMR measurements of CM-CQ were recorded on a 300 MHz (for 1H) and 500 MHz (for 13C) Bruker spectrometer (Fällanden, Switzerland) fitted with a pulse-field gradient probe. The residual resonance of deuterated solvents was used as an internal reference.

1H NMR (300 MHz, MeOD): δ 0.87 (t, J = 6.4 Hz, 3H), 1.52–1.57 (m, 2H), 1.62–1.74 (m, 2H), 2.41 (t, J = 7.0 Hz, 2H), 2.49 (q, J = 7.0 Hz, 2H), 2.97 (bs, 8H), 3.21 (m, 2H), 3.28–3.40 (m, 4H), 3.63 (s, 2H), 5.98 (s, 1H), 6.34 (d, J = 2.3 Hz, 1H), 6.43 (d, J = 5.8 Hz, 1H), 6.55 (dd, J = 2.6 Hz, 9.1 Hz, 1H), 7.25 (dd, J = 2.0 Hz, 8.8 Hz, 1H), 7.42 (d, J = 8.8 Hz, 1H), 7.71 (d, J = 2.0 Hz, 1H), 7.99 (d, J = 8.1 Hz, 1H), 8.29 (d, J = 5.8 Hz, 1H). 13C NMR (125 MHz, MeOD): δ 12.1, 26.2, 27.1, 40.0, 40.1, 40.2, 44.7, 50.5, 55.7, 58.7, 98.6, 99.6, 109.7, 110.3, 110.5, 118.7, 124.5, 125.8, 126.8, 127.3, 136.2, 149.4, 152.2, 152.7, 154.6, 157.1, 164.2, 171.5, 175.2.

Analytical HPLC and mass spectrometry of CM-CQ was carried out on a Shimadzu LC-IT-TOF system (Kyoto, Japan) equipped with a autosampler, using a reverse-phase Phenomenex Luna (Torrance, CA, USA) 5 μm C18, 50 × 3.0 mm column. Solvent mixtures of 0.1% TFA/H\(_2\)O (solvent-A) and 0.1% TFA/CH\(_3\)CN (solvent-B) were used as coeluents under a linear gradient of 1–100% of solvent B (over 45 min; flow rate of 0.5 ml/min). This gave a purity of CM-CQ of 98% with a found m/z [M + 1\(^+\)] of 607.277 (calculated for C\(_32\)H\(_39\)ClN\(_6\)O\(_4\): 607.280).

#### Inhibitor preparation and treatments

Inhibitors were all purchased from Sigma-Aldrich, dissolved in manufacturer’s recommended solvents to stock concentrations and stored at −20°C. 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 were as follows: 4HT, 20 μM; zVAD, zDEVD and CA-074, 50 μM; zFA, zFF and zLLL, 5 μM; E64d, 12.5 μM; BAPTA-AM, 1–50 μM. Lower concentrations of zFA, zFF and zLLL were used because of their poor solubility. Vehicle control consisted of equivalent amounts of DMSO and/or ethanol in accordance with solvent amounts used in various inhibitor treatments.

Mitochondria transmembrane potential assay. Cell-permeable lipophilic cation probe JC-1 (Molecular Probes, Eugene, OR, USA), also known as 5,5′,6,6′-tetrachloro-1,3,3′,3′-tetraethyl-benzimidazolylcarbocyanine iodine, normally has green fluorescence (525 nm) but aggregates at high transmembrane potential of functional mitochondria to emit an orange-red fluorescence (exitation maximum at 590 nm). A final concentration of 6 μM of JC-1 was added to 0.5 ml of treated parasitized erythrocytes and incubated at 37 °C in a dark humidified candle jar for 30 min. Cells were washed twice with PBS and resuspended to the original volume before being analyzed by flow cytometry. Percentage of JC1(Red)/JC1(Green) ratios were obtained to determine the extent of mitochondrial dysregulation as per the manufacturer’s recommendations.

Cysteine protease activity assay. CaspaseTag pancaspase in situ assay kit, sulforhodamine (Chemicon, Temecula, CA, USA), consists of a membrane-permeable fluorometthy ketone inhibitor of caspases conjugated to sulforhodamine (SR-VAD-fmk). Saponin enrichment was performed on 2 ml of treated cells, washed twice with 2 ml PBS and resuspended in 300 μl MCM. In accordance with the manufacturer’s instructions, each vial of CaspaseTag was dissolved in 50 μl DMSO to produce the CaspaseTag stock solution (150 μg) and stored in the dark at −20°C. Before staining, further dilutions were made using MCM till a working solution of 30 μl CaspaseTag was obtained. To each sample, 7 μl of the freshly prepared CaspaseTag reagent was added and the culture incubated at 37 °C in the dark for 1 h. Thereafter, two washes were carried out with 1 ml of the wash buffer provided. The samples were resuspended in 1 ml of wash buffer before flow cytometry analysis.

In situ DNA fragmentation assay. ApoDirect DNA Fragmentation Assay kit (Cintech, San Diego, CA, USA) is a TUNEL (terminal deoxynucleotidyltransferase dUTP nick-end labeling) assay that makes use of FITC-conjugated dUTP incorporation into nicked DNA by means of terminal deoxynucleotidyltransferase. Saponin enrichment of 2 ml of the treated cells also permeabilized the parasite’s plasma membrane to the contents of the TUNEL stain. Parasites were then spun down and washed twice with 1 ml of wash buffer provided in the kit. Samples were then resuspended in 50 μl of the premixed staining solution, as directed by the manufacturer, and incubated in the dark for 90 min at 37 °C. Samples were then washed twice in 1 ml of rinse buffer and resuspended in 1 ml of PBS before flow cytometry.

Hoechst staining of parasite DNA content. Hoechst 33342 stain (provided together with CaspaseTag kit, Chemicon) is a lipophilic DNA-binding fluorescent stain that is excited by ultraviolet light (350 nm), and emits blue fluorescence light (461 nm). After the parasite cultures were incubated in the absence/presence of drugs/inhibitors for 10 h, cells were washed twice with PBS and 2 μl of the 200 μg/ml stain was added per 1 ml of culture for 30 min. The cells were then washed twice and resuspended in 1 ml of PBS before flow cytometry.

Intracellular calcium detection. Cell-permeable Fluor-4-AM (Molecular Probes) fluoresces green when bound with Ca2+ and was used to detect intracellular calcium stores. A final concentration of 2 μl of Fluor-4-AM was added to 0.5 ml of treated parasitized erythrocytes and incubated at 37 °C in a dark humidified candle jar for 30 min. Cells were washed twice with PBS and resuspended to the original volume before viewing under a confocal microscope. At least 30 parasitized cells were counted for each condition.

Inhibitory concentration (IC50) determination. Parasitized cultures (ring stage) diluted with fresh erythrocytes and MCM to 2% parasitemia and 1.25% hematocrit were incubated with various concentrations of commercial CQ (Sigma, Dorset, UK), CM (i.e., 7-dimethylamino-4-coumarinacetic acid) and CM-QC for 48 h. To determine parasitemia, cultures were stained with Hoechst 33342, as described above, and analyzed by flow cytometry.

Flow cytometric analyses. Cell numbers and fluorescence intensity after staining with JC-1, CaspaseTag, FITC-upt or Hoechst were assayed by a flow cytometer (DAKO Cytomation Cyte ADP, Fort Collins, CO, USA), equipped with an argon-ion laser tuned to 488 nm and a violet laser tuned to 405 nm. Blue, green and red fluorescence was detected using 450, 525 and 595 nm band-pass filters, respectively. For experiments involving 7G8 and K1, a high-throughput flow cytometer (BD LSRII Special Order System, San Jose, CA, USA) was used to manage the large number of samples involved. Using BD LSRII, JC-1 stain was excited with 488 nm blue laser with green fluorescence being detected using the 505LP/525/570BP filter, whereas red fluorescence was detected using the 570LP/585/620BP filter. For Hoechst stain, excitation was by the 355 nm UV laser, whereas detection used the 450LP/505BP. Both flow cytometers were adjusted for forward and side scattering to accommodate either infected erythrocytes or liberated parasites. Between 100 000 and 200 000 erythrocytes or 10 000–20 000 enriched parasites were analyzed from each sample and its respective duplicate to ensure reliability. At least three independent experiments were performed unless otherwise stated.

Fluorescence and confocal microscopy. For fluorescence microscopy, wet mounts of stained infected erythrocytes or enriched parasites were observed using an image analysis setup consisting of a light and fluorescence microscope (Olympus BX60, Japan) coupled to a microscope digital camera (Olympus DP71, Tokyo, Japan) and images were captured using computer software DP Manager version 3.1.1.208 (Olympus). Confocal imaging was performed using Olympus Fluoview FV1000 (Japan) equipped with solid state and argon ion lasers tuned to 405 and 488 nm, respectively. Images were captured using Olympus Fluoview version 1.6b. For enumeration purposes, images of at least 50 parasitized erythrocytes were screened per concentration of drug used.

Statistical analyses. All data shown are means ± S.E.M. Statistical difference between percentages of stained cells was measured using univariate t-test for means and comparisons between multiple groups were analyzed using ANOVA and post hoc comparison using Tukey’s test for paired comparisons. Significantly different results (P < 0.05) were highlighted.

Conflict of interest

The authors declare no conflict of interest.

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