Central Role of Hemoglobin Degradation in Mechanisms of Action of 4-Aminoquinolines, Quinoline Methanols, and Phenanthrene Methanols

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We have used a specific inhibitor of the malarial aspartic proteinase plasmspin I and a nonspecific cysteine proteinase inhibitor to investigate the importance of hemoglobin degradation in the mechanism of action of chloroquine, amodiaquine, quinine, mefloquine (MQ), halofantrine, and primaquine. Both proteinase inhibitors antagonized the antiparasitic activity of all drugs tested with the exception of primaquine. An inhibitor of plasmspin I, Ro40-4388, reduced the incorporation of radiolabelled chloroquine and quinine into malarial pigment by 95%, while causing a 70% reduction in the incorporation of radiolabelled MQ. Cysteine proteinase inhibitor E64 reduced the incorporation of chloroquine and quinine into malarial pigment by 60 and 40%, respectively. This study provides definitive support for the central role of hemoglobin degradation in the mechanism of action of the 4-aminoquinolines and the quinoline and phenanthrene methanol antimalarials.

The 4-aminoquinolines chloroquine (CQ) and amodiaquine (AQ), the quinoline methanols quinine (QN) and mefloquine (MQ), and the phenanthrene methanol halofantrine (HF) all exert selective toxicity towards the erythrocytic stages of malaria parasites and were developed based on a knowledge of quinine structure and activity (29, 31, 41). Although there are structural similarities, QN, MQ, and HF are generally considered to constitute a group distinct from CQ and AQ. This classification is based on a number of reported differences. The 4-aminoquinolines are diprotonated and less lipid soluble at physiological pH, whereas the others, most notably QN and MQ, are much weaker bases (26, 28, 50). Recent reports suggest an inverse relationship between parasite sensitivity to CQ and sensitivity to MQ, HF, and AQ (22, 40, 44, 49). CQ and AQ induce pigment clumping in Plasmspin berghel (24, 46). The quinoline methanols do not induce pigment clumping but can inhibit 4-aminoquinoline-induced clumping (20, 27, 30). Based on these observations and spectrophotometric studies it has been suggested that the interactions between the two drug classes and hematin are fundamentally different (48, 49).

Morphological effects following treatment with MQ, QN, and HF are similar to that observed following treatment with CQ, i.e., an initial swelling of the acid food vacuole (20, 27, 30). It is generally accepted that CQ (and AQ) exerts its antimalarial action by an initial swelling of the acid food vacuole (20, 27, 30). The inhibition of hematin polymerization (12) shows similar 50% inhibitory concentration (IC50) values in vitro. This evidence has been the basis for questioning whether these drugs interact at different points within the hemoglobin degradation process or if MQ has an additional or an independent mechanism of action distinct from that of AQ and CQ (10, 14).

Hemoglobin degradation within the parasite is an ordered process (16, 18) involving at least three proteinases. Aspartic proteinase plasmspin I is responsible for the initial cleavage of the hemoglobin tetramer at the hinge position, the Phe33-Leu34 bond in the α-globin chain (17). A second aspartic proteinase, plasmspin II, has also been identified and may have a role in the cleavage of denatured hemoglobin (16). Falcipain, a cysteine proteinase, is also implicated in the cleavage of peptides from the denatured hemoglobin (15, 16). The amino acids resulting from this process are presumably used by the parasite (37, 43). It is generally agreed that the remaining hematin residue, which is potentially toxic, is removed via a polymerization process (11, 38), degradation, or export. Much

<table>
<thead>
<tr>
<th>Drug</th>
<th>Mean IC50 (nM) ± SD for isolate:</th>
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<tr>
<td></td>
<td>K1</td>
</tr>
<tr>
<td>Ro40-4388</td>
<td>256 ± 26</td>
</tr>
<tr>
<td>E64</td>
<td>8,982 ± 744</td>
</tr>
<tr>
<td>CQ</td>
<td>183 ± 1.5</td>
</tr>
<tr>
<td>HF</td>
<td>110 ± 10</td>
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<tr>
<td>MQ</td>
<td>2.6 ± 0.9</td>
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<tr>
<td>QN</td>
<td>14.8 ± 3.2</td>
</tr>
<tr>
<td>PQ</td>
<td>176 ± 25</td>
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<td>2,072 ± 635</td>
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** Data were derived from at least three assays performed at a hematocrit of 1% and a parasitemia of 1%.
evidence has accumulated to support the hypothesis that quinoline type blood schizontocides exert their antimalarial activity through interacting with hematin (3, 8, 12, 42). We have recently extended these observations and have provided strong evidence, in the case of CQ, that both the mechanism of action and resistance in the parasite are based on drug access to hematin (5). Further, it has been reported that a specific inhibitor of malarial plasmepsin I, Ro40-4388, antagonizes the actions of CQ (25).

We have used Ro40-4388 and a nonselective inhibitor of cysteine proteinase, E64, as probes to determine if the antimalarial activities of QN, HF, MQ, AQ, and CQ all depend on the efficient degradation of hemoglobin. Primaquine (PQ), an aminoquinoline antimalarial which does not inhibit hematin polymerization (12, 19) and which probably exerts its antimalarial action via a heme-independent mechanism (12), was used as a control.

MATERIALS AND METHODS

Drugs used in the study. CQ, AQ, QN, PQ, and trans-epoxysuccinyl-l-leucylamido-(4-guanidino)-butane (E64) were purchased from Sigma, Dorset, United Kingdom. MQ and Ro40-4388 were obtained from Hoffmann-La Roche, Basel, Switzerland, and HF was obtained from SmithKline Beecham.

Parasite isolates and cultivation. A CQ-resistant isolate of Plasmodium falciparum K1 and CQ-sensitive isolate HB3, obtained from D. C. Warhurst, London School of Hygiene and Tropical Medicine, London, United Kingdom, were used throughout this study. Parasite cultures were maintained by an adaptation of the method of Jensen and Trager (21). Cultures were synchronized by the method of Lambros and Vandenburg (23) before use.

In vitro sensitivity assays. Drug susceptibilities were assessed by the measurement of [3H]hypoxanthine incorporation into parasite nucleic acid as previously described by Desjardins et al. (9). Drug IC_{50}s were calculated from the log of the dose/response relationship, as fitted with Grafit software (Erithacus Software, Kent, United Kingdom). Results are given as the means of at least three separate experiments.

Drug combination assays. To analyze the combined effect of the antimalarials and proteinase inhibitors (plasmepsin I inhibitor Ro40-4388 and cysteine proteinase inhibitor E64) the IC_{50} for each drug alone was obtained as described above. From these values, a stock solution of each drug was prepared such that the IC_{50} of each drug would fall around the fourth serial dilution. Combinations of the stock solutions were prepared in constant ratios of 0:10, 1:9, 3:7, 5:5, 7:3, 9:1, and 10:0. Each combination was serially diluted across a microtiter plate and processed as for the standard sensitivity assay. The fractional inhibitory concentration (FIC; FIC = IC_{50} of the drug in the combination/IC_{50} of the drug when tested alone) of each drug was calculated and plotted as an isobologram (4).

Hemozoin purification. Ring stage parasites were incubated for 24 h in the presence of radiolabelled drug and in the presence or absence of a fixed concentration of proteinase inhibitors (plasmepsin I inhibitor Ro40-4388 and cysteine proteinase inhibitor E64) the IC_{50} for each drug alone was obtained as described above. From these values, a stock solution of each drug was prepared such that the IC_{50} of each drug would fall around the fourth serial dilution. Combinations of the stock solutions were prepared in constant ratios of 0:10, 1:9, 3:7, 5:5, 7:3, 9:1, and 10:0. Each combination was serially diluted across a microtiter plate and processed as for the standard sensitivity assay. The fractional inhibitory concentration (FIC; FIC = IC_{50} of the drug in the combination/IC_{50} of the drug when tested alone) of each drug was calculated and plotted as an isobologram (4).

FIG. 1. Isobolograms showing the relationship between the FICs of Ro40-4388 and AQ (A), CQ (B), HF (C), MQ (D), and QN (E) in the K1 isolate.
RESULTS

In vitro sensitivity of the parasites to antimalarial drugs and proteinase inhibitors. The IC$_{50}$ data for all antimalarial drugs tested and for proteinase inhibitors Ro40-4388 and E64, indicative of their activities against CQ-resistant isolate K1 and CQ-sensitive isolate HB3, are shown in Table 1. The ability of specific plasmepsin I inhibitor Ro40-4388 to inhibit parasite growth is shown to be more potent than that of cysteine proteinase inhibitor E64. The CQ-resistant and CQ-sensitive parasite isolates showed no differential susceptibilities to these proteinase inhibitors. PQ displayed antimalarial activity weaker than those of the other quinoline-containing drugs used in this study.

The interaction between quinolines and proteinase inhibitors. Representative isobolograms for antimalarial drug-proteinase inhibitor combinations are shown in Fig. 1, 2, and 3. The interactions between Ro40-4388 and CQ, AQ, MQ, and HF against the K1 isolate were antagonistic (Fig. 1). Similar antagonism was observed between E64 and these five drugs (Fig. 2). In contrast, the interaction between PQ and Ro40-4388 was additive (Fig. 3). Similar data were obtained with the HB3 isolate (data not shown).

The effect of proteinase inhibitors on the incorporation of quinolines into hemoglobin. Incubation of ring stage parasites with either radiolabelled CQ, QN, or MQ over 24 h resulted in radiolabelled drug incorporation within the malarial pigment. Ro40-4388 at its IC$_{50}$ reduced CQ and QN incorporation by more than 95% (Fig. 4A and B) and produced a 70% reduction in MQ incorporation (Fig. 4C). E64 was less efficient in reducing the incorporation of radiolabelled drugs (Fig. 4A and B). The reductions produced by E64 at its IC$_{50}$ were approximately 60% for CQ and 40% for QN. This effect of E64 is consistent with the observations of Asawamahasakda et al. (1).
The malaria parasite needs to degrade hemoglobin for successful growth and development. We believe that this is highlighted by the ability of the two proteinase inhibitors used in this study to inhibit parasite growth, as measured by the incorporation of hypoxanthine. The IC_{50} of proteinase inhibitors Ro40-4388 and E64 in P. falciparum presented here are comparable to those reported earlier (2, 25). Cysteine proteinase inhibitor E64 has previously been shown to inhibit parasite growth at the trophozoite stage, causing the accumulation of undegraded hemoglobin within the food vacuole (2, 35, 36). E64 was shown to reduce the formation of hemozoin via an inhibition of hemoglobin degradation (1, 35), and this effect was irreversible (35). In contrast, Ro40-4388 has been shown to inhibit the growth of P. falciparum parasites in vitro at nanomolar concentrations (25). Interestingly, the inhibitory effects of Ro40-4388 on hemozoin formation and parasite growth were reversible (unpublished observations). The removal of inhibitor after 24 h of incubation by minimal washing in complete medium was followed by pigment production and parasite growth. This apparent parasitostatic effect may have important implications for the use of these inhibitors as antimalarials.

Moon et al. (25) have shown that Ro40-4388 and CQ interact antagonistically against P. falciparum. We have found similar antagonism between Ro40-4388 and AQ, QN, MQ, and HF. These data suggest that all of the drugs tested have a common mechanism of action based on some component of the hemoglobin degradation process. The fact that [3H]QON and [3H]MQ are incorporated into hemozoin in a manner similar to that of [3H]CQ (42) lends support to this argument but does not provide conclusive proof. Since we have previously shown that Ro40-4388 produces a marked decrease in the number of binding sites, specifically for heme-binding drugs (5), we believe that the antagonism between Ro40-4388 and the quinolines and phenanthrenes is due to a reduction in the amount of heme available for drug binding. Therefore, the observation that the incorporation of a radiolabelled drug (CQ, QN, or MQ) into the growing hemozoin polymers is almost completely arrested in the presence of Ro40-4388 would suggest that it is the interaction of the drug with the heme monomer or polymer which is central to activity, rather than any secondary effect on heme polymerization. This is in keeping with many of the hypotheses put forward to explain the antimalarial activities of these drugs over the years (5, 7, 13) and argues against the need to invoke different mechanisms of action for the 4-aminoquinolines and the quinoline or phenanthrene methanols, as has been suggested (10, 14). The facts that PQ does not inhibit hemein polymerization (12) and that it most likely exerts its antimalarial action via a heme-independent mechanism (12) give support to the use of PQ as a control in this study. As predicted, there was no antagonism between PQ and Ro40-4388. The observations of antagonism between the cysteine proteinase inhibitor E64 highlight, we believe, the importance of the cysteine proteinase falcipain in hemoglobin degradation and help to further confirm the view that all of these drugs (CQ, AQ, QN, MQ, and HF) exert their antimalarial effects via a common heme-dependent mechanism.

We have confirmed the incorporation of radiolabelled CQ into the growing hemozoin reported initially by Sullivan et al. (42) and have extended these observations to QN and MQ. The fact that this incorporation can be reduced with the proteinase inhibitors further supports a role for hemoglobin degradation in their antimalarial activity. It could be argued that, as incorporation was reduced by approximately 50% by E64 at its IC_{50}, some of this effect could be the indirect result of parasite death. However, exposure to this concentration of E64 for an equivalent period has been shown to have little effect on hypoxanthine uptake (1) and, by implication, on parasite viability.

The data presented here confirm the central and common role of hemoglobin degradation in the mechanisms of action of the 4-aminoquinolines, the quinoline methanols, and the phenanthrene methanols. This supports the view that all of these compounds are acting through the same process without the need to invoke additional targets. The data confirm that proteinase inhibition may be a rational target for antimalarial chemotherapy. If future strategies include the use of these inhibitors in combination with other antimalarial drugs, the
antagonism seen here would argue against combinations with quinoline type compounds.

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