Antimalarial activity of Malaria Box Compounds against *Plasmodium falciparum* clinical isolates

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**A B S T R A C T**

Malaria remains a major cause of childhood deaths in resource-limited settings. In the absence of an effective vaccine, drugs and other interventions have played very significant roles in combating the scourge of malaria. The recent reports of resistance to artemisinin necessitate the need for new antimalarial drugs with novel mechanisms of action. Towards the development of new, affordable and easily accessible antimalarial drugs for endemic regions, the Medicines for Malaria Venture (MMV) assembled a total of 400 active antimalarial compounds called the Malaria Box. The potency and the efficacy of the Malaria Box Compounds have been determined mainly using laboratory strains of *P. falciparum*.

This study investigated the potency of twenty compounds from the Malaria Box against four clinical isolates from Ghana, using optimized *in vitro* growth inhibitory assays. Seven out of the 20 compounds screened had 50% inhibitory concentration (IC50) below 500 nM. The most active among the selected compounds was MMV006087 (average IC50 of 30.79 nM). Variations in the potency of the Malaria Box Compounds were observed between *P. falciparum* clinical isolates and Dd2 strain. We also investigated the sensitivity of the clinical isolates to chloroquine and artesunate. The N093 clinical isolate was found to be resistant to chloroquine but showed high sensitivity to artesunate. The results underscore the importance of including clinical isolates with different drug-resistant backgrounds, in addition to laboratory strains, in validating potential compounds during antimalarial compound screening programs.

1. Introduction

*Plasmodium falciparum* related malaria is still a major threat to health systems in resource-limited settings. Since 2010, the global malaria incidence has declined by about 21%, however, malaria remains a major global health problem. An estimated 429,000 deaths were recorded in 2015 (WHO, 2016). About 90% of the global malaria-related deaths occurred in WHO Africa region with about 70% of these deaths occurring in children under five years (WHO, 2016). The global efforts to eliminate malaria over the past decades have largely been hampered by the development of insecticide and drug resistance by mosquitoes and *Plasmodium* respectively (Peacham et al., 2010; Ranson et al., 2009).

The emergence of resistant parasites to antimalarial drugs such as chloroquine (Moore and Lanier, 1961; Payne, 1987), sulfadoxine and pyrimethamine (Hurwitz, 1981) over the years, led to the recommendation for the use of artemisinin and artemisinin-based combination therapies (ACT), as first-line drugs for the treatment of malaria in all endemic regions (Smithuis et al., 2004; Valecha et al., 2010; WHO, 2001). Although there was a recent report *P. falciparum* isolate that was resistant to artemisinin in Equatorial Guinea (Lu et al., 2017), resistance to ACTs is still not wide-spread in Africa (WHO, 2016). Nevertheless, the recent emergence of ACT resistant *Plasmodium falciparum* strains in South-East Asia (Ashley et al., 2014; Dondorp et al., 2009; Noedl et al., 2008; Yeung et al., 2009), calls for new sets of antimalarial drugs with novel mechanisms of action.

Another important consideration is the lack of compliance to antimalarial drugs on the part of patients resulting in inadequate treatments, which may lead to selection and transmission of resistant parasites (White et al., 2009). The need for single dose drugs to promote patient compliance, with less ability to drive the development of resistance is therefore critical (Alonso et al., 2011). This not only offers an...
advantage of easy patient compliance, but also makes treatment less expensive (Burrows et al., 2013). In addition to possessing a long acting effect against drug resistant malaria parasites, such a drug should be able to reduce the parasite burden in asymptomatic individuals who serve as reservoirs for malaria transmission (Alonso et al., 2011; Diagana, 2015). Some recent studies have found a few antimalarial compounds with multi-stage activity (Baragana et al., 2015; Fidock, 2016; Kato et al., 2016), but there is the need to increase the repertoire of such compounds to provide the foundation for development of new drugs.

A number of drug screening studies have identified potent antimalarial compounds from the Malaria Box (Spangenberg et al., 2013) against laboratory-adapted strains of P. falciparum (Fong et al., 2015; Lucantoni et al., 2013; Van Voorhis et al., 2016). However, not much is known of the activity of these compounds against clinical isolates. In this study, in vitro drug susceptibility screening and polymerase chain reaction-restriction fragment length polymorphism (PCR-RFLP) genotyping analysis were adapted to confirm the potency of twenty selected Malaria Box Compounds using P. falciparum clinical isolates from Ghana. The compounds selected were based on their high potency against P. falciparum laboratory strains that was previously established using the entire Malaria Box library (Creek et al., 2016; Tiwari et al., 2016; Van Voorhis et al., 2016). The isolates were also screened against chloroquine and artemesunate in defining their relative susceptibility to the different compounds. In addition, PCR-RFLP was used to investigate established drug associated mutations in four genes; P. falciparum chloroquine resistance transporter (pfCRT), P. falciparum multidrug resistance gene 1 (pfmdr1), P. falciparum dihydrofolate reductase gene (pfDHFR) and P. falciparum dihydropteroate synthase gene (pfDPS), known to mediate and/or modulate resistance to standard antimalarials.

2. Materials and methods

2.1. Clinical isolates

Four clinical isolates (K239, N093, A156 and A160) were randomly selected from archived samples from three endemic areas in Ghana, with different transmission intensities (Accra-low endemcity area, Kintampo-holoendemic area and Navrongo-hyperendemic area). The samples were collected as part of an ongoing study on erythrocyte invasion mechanisms (EIM). The samples were cryopreserved in liquid nitrogen immediately upon arrival from the field. The selected cryopreserved parasites were thawed and cultured with human blood group O+ erythrocytes using standard methods (Trager and Jensen, 1976) with slight modifications. P. falciparum clinical isolates were cultured to > 5% (approximately 10 cycles) parasitemia of ring stage parasites. Using 5% Sorbitol treatment, a synchronized culture of ring-stage parasites (Lambros and Vanderberg, 1979) was obtained and diluted to 1% parasitemia in 2% haematoctrit for the growth inhibition assays.

2.2. In vitro drug susceptibility assay

The library of 400 Malaria Box Compounds was provided by Medicines for Malaria Venture (MMV) at concentrations of 10 mM in dimethyl sulfoxide (DMSO) in 96-well microtiter plates (Spangenberg et al., 2013). A total of 20 compounds out of the 400 Malaria Box Compounds were selected for this study. The Malaria Box Compounds were serially diluted and screened against the clinical isolates at a concentration ranging from 0.064 nM to 25 μM. All the growth inhibition assays were set up in triplicate wells at final well volumes of 100 μL consisting of 10 μL of the test compound and 90 μL parasite culture at 1% parasitemia and 2% haematocrit in a 96-well flat bottom plate. The assay was incubated at 37 °C for 48 h. RPMI containing 0.25% DMSO was used as negative control whilst uninfected erythrocytes at 2% haematocrit were used as background control. After 48 h, 80 μL of the supernatant was taken out and replaced with 80 μL of SYBR Green I (Invitrogen, USA) stain, which was used to differentiate infected erythrocytes from uninfected erythrocytes by flow cytometry analysis. The plates were incubated in the dark with the SYBR Green I stain for 30 min prior to flow cytometry analysis.

The parasitemia corresponding to each culture well was quantified as previously described (Wirjanata et al., 2015) with modifications using BD FACS LSRSort™ X-20 flow cytometer with the BD FACSDiva Software (v8.0.1). The forward and side scatter parameters were used to gate erythrocytes and exclude debris. Photo multiplier tube (PMT) voltages of 200, 250 and 300 V were set for the forward scatter, side scatter and SYBR Green I, respectively. A dilution of 1:50 comprising 10 μL of packed erythrocytes to 490 μL of sheath fluid (BD Bioscience, USA) was used in the flow cytometry analysis. SYBR Green I positive erythrocytes corresponding with infected erythrocytes were used to determine the mean parasitemia levels. Data from 50,000 cells per well were recorded for all assays. The sensitivity of the clinical isolates to chloroquine and artemesunate was also evaluated. Stock solutions of 10 mM chloroquine solution was prepared and used to screen against the four clinical isolates at final well concentrations between 0.0064 nM and 25 μM. The negative control was sterile distilled water in place of chloroquine. Artesunate stock of 1 mM artemesunate (Sigma-Aldrich, USA) were similarly prepared in DMSO and tested at final well concentrations between 0.00248 nM and 32 nM. A 0.1% DMSO in RPMI was used in the negative control wells.

2.3. Determination of the molecular markers of drug resistance

P. falciparum genomic DNA was extracted from the clinical isolates (N093, A156, A160 and K239) using the QIAamp blood midi kit (QIAGEN, USA) and stored at −20 °C. Each of the isolates was analyzed for the putative point mutations that have been shown to mediate antimalarial drug resistance in the pfCRT (K76T, pfmdr1 (N86Y and Y184F), pfDHFR (N51I, C59R and S108N) and pfdhps (A437G) genes. Regions flanking these point mutations were amplified by polymerase chain reaction (PCR) using previously reported primer sets (Djimdé et al., 2001; Duraisingham et al., 1998). One microlitre of the outer PCR products was used as template DNA in the nested PCR. Both the outer and the nested PCRs were set at a final volume of 25 μL containing 1X of Maxima Hot Start Green PCR master mix (Thermo Scientific, USA) and primers at a final concentration of 250 nM. Five microlitres of the nested PCR products were run on ethidium bromide-stained 2% agarose and then visualized using the Amersham Imager 600 (General Electric Healthcare Life Sciences, USA). Five microlitres of the remaining products were digested with restriction enzymes (New England BioLabs, USA) specific for the mutations, in a total volume of 15 μL. The restriction digest products were run on ethidium bromide-stained 2% agarose gels and visualized using the Amersham Imager 600. Purified P. falciparum genomic DNA from laboratory-adapted strains (DD2, 3D7, K1, W2, FCR3 and 7G8) were used as controls.

2.4. Data analysis

The data from the flow cytometry analysis was first formatted in Microsoft Excel by subtracting the background fluorescence from all the data. GraphPad Prism (Version 6.01) was then used to generate sigmoidal dose-response curves by fitting a non-linear regression curve to the data. The 50% inhibitory concentration (IC50) values were then estimated from the dose response curves. Each data point on the dose response is presented as the mean ± (SEM) of two experiments set up in triplicate.

2.5. Ethics

The studies were approved by the ethics committees of the Ghana Health Service, Navrongo Health Research Centre, Kintampo Health
Table 1
Summary of information on the 20 Malaria Box Compounds screened. The structures and IC50 values for all the 20 compounds against the four clinical isolates and Dd2 are all shown.

| 1 | MMV009015 | MMV006278 | MMV000753 | MMV019555 |
|---|---|---|---|---|
| **N093:** | 275.60 nM | 1900 nM | 1122 nM | 1122 nM |
| **A156:** | 804.40 nM | 18410 nM | 769.60 nM | 811.10 nM |
| **A160:** | 758.80 nM | 9460 nM | 940.70 nM | 940.70 nM |
| **K239:** | 1009 nM | 4340 nM | 769.60 nM | 811.10 nM |
| **Dd2:** | | | 811.10 nM | 811.10 nM |

| 2 | MMV006087 | MMV019555 | MMV006787 | MMV085203 |
|---|---|---|---|---|
| **N093:** | 659.90 nM | 3318 nM | 190 nM | 55.83 nM |
| **A156:** | 701.10 nM | 280.70 nM | 939.60 nM | 280.70 nM |
| **A160:** | 835.80 nM | 9460 nM | 811.10 nM | 811.10 nM |
| **K239:** | 801.60 nM | 399.90 nM | 811.10 nM | 811.10 nM |
| **Dd2:** | | | 811.10 nM | 811.10 nM |

| 3 | MMV000248 | MMV006587 | MMV089856 |
|---|---|---|---|
| **N093:** | 31.60 nM | 4164 nM | 940.70 nM |
| **A156:** | 26.82 nM | 8100 nM | 1122 nM |
| **A160:** | 42.39 nM | 212.50 nM | 1122 nM |
| **K239:** | 42.39 nM | 1122 nM | 1122 nM |
| **Dd2:** | 74.34 nM | 178.80 nM | 1122 nM |

| 4 | MMV007275 | MMV0065843 | MMV006455 |
|---|---|---|---|
| **N093:** | 554.10 nM | 1166 nM | 153.50 nM |
| **A156:** | 716.10 nM | 776.90 nM | 239.57 nM |
| **A160:** | 759.80 nM | 713.50 nM | 239.57 nM |
| **K239:** | 327.50 nM | 921.20 nM | 239.57 nM |
| **Dd2:** | 1119 nM | 21930 nM | 1122 nM |

| 5 | MMV665949 | MMV665977 | MMV007199 |
|---|---|---|---|
| **N093:** | 9764 nM | 282.90 nM | 19.02 nM |
| **A156:** | 4138 nM | 144.20 nM | 19.02 nM |
| **A160:** | 8351 nM | 79.08 nM | 19.02 nM |
| **K239:** | 3991 nM | 1334 nM | 19.02 nM |
| **Dd2:** | 1309 nM | 7769 nM | 19.02 nM |

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Research Centre, and Noguchi Memorial Institute for Medical Research, University of Ghana, Legon. All samples were collected after obtaining written informed consent from the parents/guardians of participating children who were aged ≤10 years. For children older than 10 years, additional assent was obtained from the donor, following receipt of parental consent. Blood used in this study for culturing was obtained from donors with informed consent.

3. Results

3.1. Determining the potency of Malaria Box Compounds against clinical and laboratory isolates of *P. falciparum*

A subset of 20 Malaria Box Compounds (Table 1) that have been found to be potent against laboratory strains of *P. falciparum* (Fong et al., 2015; Tiwari et al., 2016; Van Voorhis et al., 2016) were selected and evaluated against four clinical isolates of *P. falciparum* (N093, A156, A156 and K239). Out of the 20 compounds screened, the dose response curves of the four most potent compounds to all the four clinical isolates have been shown (Fig. 1). In total, seven compounds had IC$_{50}$ values below 500 nM, six compounds had IC$_{50}$ values between 500 nM and 1 μM and the remaining seven had IC$_{50}$ values greater than 1 μM (Fig. 2). Considering compounds with IC$_{50}$ values below 500 nM; three compounds (MMV008956, MMV085203 and MMV006087) were active against all four clinical isolates (Figs. 1 and 2). Three compounds (MMV006787, MMV006455 and MMV665977) were active against three clinical isolates (Figs. 1 and 2 and S1A) whilst two compounds (MMV019555 and MMV665878) were active against two clinical isolates (Fig. 2 and S1C). In addition, four compounds (MMV009015, MMV00248, MMV006764 and MMV007199) showed activity against one clinical isolate (Fig. 2, S1A to S1C). The rest of the compounds (MMV396797, MMV665843, MMV000753, MMV006764, MMV006913, MMV007275, MMV665949, MMV006278 and MMV008416) were less active against the clinical isolates with IC$_{50}$ values greater than 500 nM (Fig. 2, S1B to S1D).

The most potent compound in this screen was MMV006087, with an average IC$_{50}$ of 30.79 nM, followed by MMV085203, with an average IC$_{50}$ of 63.83 nM across the four clinical isolates (Fig. 2, Table 1). MMV006087 and MMV085203 generally showed comparable IC$_{50}$ values to that of chloroquine in three of the clinical isolates (A156, A160 and K239), but exhibited about three-fold higher potency against N093 than chloroquine (Fig. 2 and S1E). Artesunate (Fig. 2) showed about eight-fold higher potency against the clinical isolates compared to MMV006087 and MMV085203 (Fig. 1).

As a reference, IC$_{50}$ values for the 20 compounds against the laboratory strains of *P. falciparum* (either screened in our laboratory or from published studies) were compared to the IC$_{50}$ values observed against the clinical isolates (Fig. 2). Thirteen of the 20 compounds (MMV006787, MMV006455, MMV00753, MMV006764, MMV006087, MMV006913, MMV00248, MMV006455, MMV007199, MMV665977, MMV007275, MMV009015 and MMV665843) were more potent against the clinical isolates compared to the laboratory-adapted strains (Fig. 2 and Table 1). The remaining seven compounds (MMV006278, MMV008416, MMV085203, MMV396797, MMV019555, MMV665878 and MMV665949) showed higher potency against laboratory strains of *P. falciparum* compared to the clinical isolates (Fig. 2). Also, compounds such as MMV665878 (Fig. S1C) and MMV665949 (Fig. S1D) showed higher potencies against some of the clinical isolates compared to Dd2.

3.2. Responses of clinical isolates to standard antimalarial drugs

To understand the drug sensitivity background of the clinical isolates used in this study, the clinical isolates were also screened against chloroquine and artesunate. The observed IC$_{50}$ values were then compared with the standard IC$_{50}$ values (Pradines et al., 2011) for both chloroquine and artesunate as well as the selected Malaria Box Compounds. The clinical isolate N093 was found to be resistant to chloroquine (IC$_{50}$ > 100 nM), but was sensitive to artesunate (IC$_{50}$ < 10.5 nM) (Table 1 and Fig. S1E). Relative to the Malaria Box compounds and chloroquine, artesunate was more potent against all the four clinical isolates with IC$_{50}$ ranging from 2.69 to 12.68 nM (Fig. 2 and S1E). Artesunate showed less activity against A156 with an IC$_{50}$ value of 12.68 nM which is about four-fold higher than the average for the other three isolates (Table 1 and Fig. S1E).

3.3. Drug resistance background of the clinical isolates

The PCR-RFLP analysis was used to determine molecular markers that are commonly used to detect antimalarial drug resistance. Four different genes with their corresponding mutations as have been implicated for drug resistance in *P. falciparum* were assessed. The respective mutations studied are: *pfcrt* codon 76 (Djimde et al., 2001), *pfmdr1* codons 86 and 184 (Foote et al., 1990), *pfldhps* codons 437 (Wang et al., 1997) and *pfldhfr* codons 51,108 and 59 (Peterson et al., 1998). Notably, N093 which exhibited resistance to chloroquine but was most sensitive to artesunate was found to harbour the *pfcrt* 76T mutant allele, whereas A156, A160 and K239 were found to contain the wild-type *pfcrt* K76 allele (Fig. S2 and Table 2). All the four clinical isolates had the mutant *pfmdr1* 184F allele (Fig. S2 and Table 2), whereas only N093 and K239 were found to contain the *pfmdr1* 86Y mutant allele (Fig. S2 and Table 2). For the *pfldhps*, all the clinical isolates had a mutant 437G allele with K239 showing mixed genotypes (Fig. S2 and Table 2). Two of the clinical isolates (A156 and N093) had the mutant *pfldhfr* S11 allele, whilst K239 was again found with mixed genotypes (Fig. S2 and Table 2).

4. Discussions

The ultimate target for any new antimalarial compounds will be
clinical isolates. Studies have shown that clinical isolates from endemic areas have multiple clones of *P. falciparum* parasites, some of which might be drug resistant (Farnert et al., 2002; Ofosu-Okyere et al., 2001). Therefore, the use of clinical isolates of *P. falciparum* is necessary for drug discovery and susceptibility studies. Much screening activities have been conducted using the Malaria Box Compounds against laboratory strains of *P. falciparum*, but not much has been reported on the potency of these compounds against clinical isolates. It was therefore

![Fig. 1. Dose-response curves of the four most potent Malaria box compounds against four clinical isolates of *P. falciparum*. Panel A–D are dose-response curves showing the response of clinical isolates (A) N093, (B) A156, (C) A160 and (D) K239 to MMV006087, MMV085203, MMV008956 and MMV665977 at concentrations from 0.064 nM to 25 μM. Each data point represents the mean ± SEM (n = 3). The plot shows percentage parasitemia against the log of the concentration of the compound. MMV006087 was the most potent compound followed by MMV085203 and these two compounds had IC₅₀ values less than 100 nM.](image-url)
imperative to validate the most potent Malaria Box Compounds that were previously reported against *P. falciparum* clinical isolates from Ghana. In this study, we present evidence for the efficacy of Malaria Box Compounds against clinical isolates of *P. falciparum* from Ghana. Our data shows that MMV008956, MMV085203, MMV006087 and MMV65977 were the most active compounds by IC₅₀ values. However, MMV006087 was the most potent compound against all the isolates used in the screen. The high potency of this compound against the field isolates compared to the Dd2 strain implies a more susceptible phenotype in the clinical isolates, though they seem to have varied susceptibility signatures (Table 2). This compound has been shown to affect protein degradation pathways, similar to the action of chloroquine and piperaquine (Creek et al., 2016), whilst exhibiting fast-killing activity similar to that of artemisinin (Van Voorhis et al., 2016). MMV006087 has also been found to be active against early ring stage *P. falciparum* parasites as well as gametocytes (Van Voorhis et al., 2016). These unique properties of MMV006087 prioritize this compound as a suitable antimalarial candidate against blood stage and transmission stage *P. falciparum* parasites.

The glutathione and thioredoxin systems may provide a way for malaria parasites to maintain redox homeostasis and antioxidant defense, considering that *P. falciparum* lacks glutathione peroxidase and catalase (Jortzik and Becker, 2012). Studies using genetic and chemical tools have shown that *P. falciparum* thioredoxin reductase (PTrxR) is a target necessary for the survival of the parasite (Muller, 2003). In a study by Tiwari et al. MMV085203 and MMV008956 were both found to target PTrxR, but MMV085203 was more potent at inhibiting this target than MMV008956. Based on the IC₅₀ values observed in this study, MMV085203 was also found to be more potent at killing the parasite than MMV008956. The efficient killing with lower IC₅₀ values of these two compounds indicates the possible essentiality of the proteins involved in the antioxidant defense system (Arner and Holmgren, 2000; Gilberger et al., 2000). The compound MMV00753, an inhibitor of hemozoin formation (Fong et al., 2015), was found in this study to be more effective against the clinical isolates than was reported for D6; a chloroquine sensitive strain, and C235, a multi-drug resistant strain, (Fong et al., 2015). These differences in potency observed in both the clinical isolates and laboratory-adapted strains suggest the need for thorough screening with clinical isolates in antimalarial drug discovery studies.

Studies have shown that the presence of mutations in the *pfcrt* and *pfdmrd1* genes of *P. falciparum* parasites not only decrease their susceptibility to quinine and halofantrine, but increase their sensitivity to artemisinin and lumefantrine (Gresty et al., 2014; Reed et al., 2000; Sidhu et al., 2005). In this current study, the N093 clinical isolate was the only parasite that was found to have polymorphisms in both the *pfdmrd1* and *pfcrt* genes. The N093 clinical isolate was also found to be the only isolate that was very sensitive to artesunate but resistant to chloroquine in this study. It therefore suggests that the presence of SNPs in *pfdmrd1* and *pfcrt* genes might play a key role in modulating the sensitivity of this isolate to artesunate, and its resistance to chloroquine.
as was observed by other studies (Gresty et al., 2014; Reed et al., 2000; Sidhu et al., 2005).

5. Conclusion

The findings suggest that the Malaria Box Compounds have varied activities against different clinical isolates. It has been shown that of all the compounds screened, MMV006087 had the best IC50 across the different clinical isolates, with N093 being a chloroquine resistant parasite. From the data, having monoclonal populations also presents a risk of interest. The authors declare no conflict of interest.

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Conflict of interest

The authors declare no conflict of interest.
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