Methylene blue induced morphological deformations in *Plasmodium falciparum* gametocytes: implications for transmission-blocking

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**Abstract**

**Background:** Malaria remains a global health problem despite availability of effective tools. For malaria elimination, drugs targeting sexual stages of *Plasmodium falciparum* need to be incorporated in treatment regimen along with schizonticidal drugs to interrupt transmission. Primaquine is recommended as a transmission blocking drug for its effect on mature gametocytes but is not extensively utilized because of associated safety concerns among glucose-6-phosphate dehydrogenase (G6PD) deficient patients. In present work, methylene blue, which is proposed as an alternative to primaquine is investigated for its gametocytocidal activity amongst Indian field isolates. An effort has been made to establish Indian field isolates of *P. falciparum* as in vitro model for gametocytocidal drugs screening.

**Methods:** *Plasmodium falciparum* isolates were adapted to in vitro culture and induced to gametocyte production by hypoxanthine and culture was enriched for gametocyte stages using N-acetyl-glucosamine. Gametocytes were incubated with methylene blue for 48 h and stage specific gametocytocidal activity was evaluated by microscopic examination.

**Results:** *Plasmodium falciparum* field isolates RKL-9 and JDP-8 were able to reproducibly produce gametocytes in high yield and were used to screen gametocytocidal drugs. Methylene blue was found to target gametocytes in a concentration dependent manner by either completely eliminating gametocytes or rendering them morphologically deformed with mean IC\textsubscript{50} (early stages) as 424.1 nM and mean IC\textsubscript{50} (late stages) as 106.4 nM. These morphologically altered gametocytes appeared highly degenerated having shrinkage, distortions and membrane deformations.

**Conclusions:** Field isolates that produce gametocytes in high yield in vitro can be identified and used to screen gametocytocidal drugs. These isolates should be used for validation of gametocytocidal hits obtained previously by using lab adapted reference strains. Methylene blue was found to target gametocytes produced from Indian field isolates and is proposed to be used as a gametocytocidal adjunct with artemisinin-based combination therapy. Further exploration of methylene blue in clinical studies amongst Indian population, including G6PD deficient patients, is recommended.

**Keywords:** Malaria, Gametocytes, Morphological deformations, Microscopy, *P. falciparum*, Gametocytocidal activity, Methylene blue, Primaquine, G6PD deficiency
Methods

**Cultivation of asexual stages and production of gametocytes**

Asexual blood stages from infected blood collected from patients living in malaria prone areas of India—Rourkela (Odisha), Jaisalmer (Rajasthan), Jagdalpur (Chhattisgarh) and Mangalore (Karnataka), were cultivated in vitro by following the procedures of Trager and Jensen [14, 15] with minor modifications. The parasites were cultivated in RPMI 1640 medium (with glutamine) containing 25 mM HEPES, 2 g/L D-glucose, 2 g/L sodium bicarbonate, 40 μg/mL gentamicin sulfate supplemented with 10% heat inactivated AB+ human serum. All culture adapted isolates were uniformly subjected to gametocyte production by following procedures of Ifediba and Vanderberg [16] with certain modifications. Parasites were maintained in culture starting from 0.5% parasitaemia (ring stage; sorbitol synchronized) and 10% haematocrit (day 0). These parasites were kept devoid of fresh erythrocytes throughout the course of 2 weeks of culture maintenance and daily replenished with complete RPMI-1640 media supplemented with hypoxanthine (50 µg/mL). Hypoxanthine provided additional purine source required for sexual differentiation as well as maturation of gametocytes. Haematocrit was reduced to 5% on day 8 and 50 mM N-acetyl-glucosamine (Sigma) was added on days 9–12 to eliminate asexual stage parasites. On day 14 onwards, a uniform population of gametocytes is obtained with majority of late stage (stage IV and V) gametocytes. Field isolates demonstrating higher gametocytaemia than the rest were classified as gametocyte producers and were selected for in vitro drug sensitivity testing. In separate set of experiments, these isolates were cultured continuously for a period of ~ 6 months from the date of revival of cryopreserved stablate to ascertain the effect of number of in vitro asexual cycles on gametocyte production. Stability of gametocyte production phenotype across multiple cryopreservation events was also investigated. Comparative analysis was carried out using unpaired t-test and p value < 0.05 was considered as statistically significant. Drug susceptibility (asexual stages) of selected isolates to chloroquine and artesunate was also ascertained by microscopy-based schizont maturation inhibition assay [17].

**Gametocytocidal assays and data analysis**

Stock solutions of methylene blue (Sigma-Aldrich) and primaquine (Sigma-Aldrich) were prepared in double distilled water and RPMI-1640 media, respectively. Appropriate working solutions were made a fresh on the day of the experiment with complete culture medium.
Gametocytes were harvested on the day of experiment and thorough and systematic morphological examination was performed by microscopy before carrying out screening experiments. Pre-dosed culture plates were prepared by plating two fold dilutions of the drugs in duplicates to achieve the desired concentrations up to 5 µM for methylene blue and up to 25 µM for primaquine and incubated with blood containing 2–3% gametocytes. Control wells were also prepared containing drug free media along with gametocytes for calculation of untreated inhibition. Also, 0.5% DMSO and 50 µM thioestrepton (Sigma Aldrich) were used as negative and positive controls, respectively. Plates were incubated at 37 °C for 48 h in presence of 5% CO₂ [18]. After incubation period, thin smears were prepared, stained with 10% Giemsa and examined under a 100× oil immersion objective [19, 20]. Five thousand RBCs from each slide were counted to examine the gametocytaemia and gametocyte morphology at each concentration. Gametocytes observed were morphologically categorized into two groups, (1) normal morphology (NM) or (2) altered morphology (AM) and grouped either in early stage gametocytes (stages II and III) or late stage gametocytes (stages IV and V). Gametocytaemia for each concentration was expressed as percentage inhibition compared to drug-free control which was plotted against logarithm of drug concentration using a non-linear regression analysis (four parameter log dose with variable slope) to compute IC₅₀ values and 95% confidence intervals. Dose–response curves expressed as percentage inhibition vs. logarithm of drug concentration were generated by Graphpad prism 6 [21]. IC₅₀ values were calculated separately for early (stages II and III) and late (IV and V) stage gametocytes (Table 1) than other isolates in vitro and were therefore, deemed most suitable for stage specific drug screening experiments. Gametocytes produced from isolates RKL-9 and JDP-8 followed a very reproducible progression of gametocyte maturation from stage I to stage V. This was achieved in a period of ~ 12–14 days from the date of initiation of gametocyte culture (induction) and yielded > 70% of mature gametocytes. Both RKL-9 and JDP-8 were able to produce gametocytes after continuous maintenance in asexual culture for about 6 months as evidenced by statistically insignificant change (p > 0.05) in percentage gametocytaemia (Table 2). Gametocyte induction was not performed for the respective parasite lines beyond 6 months of continuous asexual culture due to technical reasons. Moreover, gametocyte production in these field isolates was found to be stable after multiple cryopreservation cycles. RKL-9 and JDP-8 collected from Rourkela and Jagdalpur, respectively, exhibited higher gametocytaemia (> 2%) (Table 1) than other isolates in vitro and were therefore, deemed most suitable for stage specific drug screening experiments. Gametocytes produced from isolates RKL-9 and JDP-8 followed a very reproducible progression of gametocyte maturation from stage I to stage V. This was achieved in a period of ~ 12–14 days from the date of initiation of gametocyte culture (induction) and yielded > 70% of mature gametocytes. Both RKL-9 and JDP-8 were able to produce gametocytes after continuous maintenance in asexual culture for about 6 months as evidenced by statistically insignificant change (p > 0.05) in percentage gametocytaemia (Table 2). Gametocyte induction was not performed for the respective parasite lines beyond 6 months of continuous asexual culture due to technical reasons. Moreover, gametocyte production in these field isolates was found to be stable after multiple cryopreservation cycles. RKL-9 and JDP-8 remained high

Table 1 Percentage of gametocytaemia in Indian field isolates and their asexual stage drug susceptibility profile

| Isolate | Place of origin | State | % Gametocytaemia | Chloroquine asexual stage IC₅₀ (nM) (95% CI) | Artesunate asexual stage IC₅₀ (nM) (95% CI) |
|---------|-----------------|-------|------------------|-----------------------------------------------|-----------------------------------------------|
| RKL-9   | Rourkela        | Odisha| Total G: (2.50 ± 0.27)% Early G: (0.58 ± 0.12)% Late G: (1.92 ± 0.35)% | 114.4 (77.70–168.4) | 3.7 (2.710–5.036) |
| JDP-8   | Jagdalpur       | Chhattisgarh | Total G: (2.18 ± 0.23)% Early G: (0.50 ± 0.17)% Late G: (1.68 ± 0.24)% | 266.7 (179.5–396.2) | 5.06 (4.107–6.229) |

Results

In vitro gametocyte production

In present study, different field isolates demonstrated varied gametocyte production. Two isolates, RKL-9 and JDP-8 collected from Rourkela and Jagdalpur, respectively, exhibited higher gametocytaemia (> 2%) (Table 1) than other isolates in vitro and were therefore, deemed most suitable for stage specific drug screening experiments. Gametocytes produced from isolates RKL-9 and JDP-8 followed a very reproducible progression of gametocyte maturation from stage I to stage V. This was achieved in a period of ~ 12–14 days from the date of initiation of gametocyte culture (induction) and yielded > 70% of mature gametocytes. Both RKL-9 and JDP-8 were able to produce gametocytes after continuous maintenance in asexual culture for about 6 months as evidenced by statistically insignificant change (p > 0.05) in percentage gametocytaemia (Table 2). Gametocyte induction was not performed for the respective parasite lines beyond 6 months of continuous asexual culture due to technical reasons. Moreover, gametocyte production in these field isolates was found to be stable after multiple cryopreservation cycles. RKL-9 and JDP-8 remained high.

Table 2 Percentage of gametocytes produced from RKL-9 and JDP-8

| Isolate | aDuration of asexual culture before induction (in months) | bNumber of cryopreservation events before induction |
|---------|---------------------------------------------------------|---------------------------------------------------|
| RKL-9   | 2.24 ± 0.21                                            | 2.50 ± 0.46                                        |
| JDP-8   | 2.02 ± 0.22                                            | 2.00 ± 0.39                                        |

Comparison of % gametocytaemia. aAt the interval of 6 months (expressed as mean ± SD of three separate induction experiments). bAfter three cryopreservation events (expressed as mean ± SD of four separate induction experiments). No significant difference (p > 0.05) in percentage gametocytaemia after induction was observed in RKL-9 and JDP-8 for both the experiments a(effect of duration of parasites in asexual culture) and b(effect of number of cryopreservation events). Percentage gametocytaemia calculations are per 5000 RBCs.
gametocyte producers even after three additional freeze–thaw cycles following initial cryopreservation of infected blood sample (time spent in asexual culture was up to 1 month before each cryopreservation event) (Table 2). Also, RKL-9 and JDP-8 were found to be chloroquine resistant with asexual stage mean IC₅₀ value 174.7 nM. Artesunate was also tested against both of these isolates and was found to effectively inhibit the parasite growth with asexual stage mean IC₅₀ of 4.32 nM. Individual asexual stage IC₅₀ values of field isolates RKL-9 and JDP-8 are mentioned in Table 1. The rest of the field isolates used in this study exhibited comparatively lesser gametocytæmia (< 0.5%) in vitro.

For evaluation of stage specific gametocytocidal activity of drugs discussed in subsequent section, gametocyte stages were categorized using a classification similar to one used by Carter and Miller [22] as seen from representative microscopic images shown in Fig. 1.

Gametocytocidal assays using methylene blue and primaquine
Methylene blue was effective in targeting both early and late stage gametocytes produced from field isolates RKL-9 and JDP-8 in a dose dependent manner with a mean IC₅₀ (NM) value of 424.1 nM (IC₅₀ Total = 958.1 nM) for early stage and 106.4 nM (IC₅₀ Total = 1060.2 nM) for late stage gametocytes. Individual IC₅₀ value for each field isolate in ‘NM’ and ‘Total’ category is mentioned in Table 3. Concentration–response curves of early and late stage gametocytes for NM category (normal morphology, data include healthy gametocytes with no visible drug induced deformation) and Total (data include gametocytes with normal morphology and altered morphology, both) are shown in Fig. 2. Methylene blue was more effective in inducing morphological deformations in late stage gametocytes group as compared to early stage gametocyte group as evident from

**Fig. 1** Representative images of different stages of gametocytes as observed under 100× objective bright-field light microscope. a–e Stage I–V gametocytes respectively (computer magnified image for better interpretation of morphology)
their IC50 values for NM category. These morphological abnormalities induced by methylene blue comprised of shrinkage and distortions that created degenerated or irregular gametocytes (Fig. 3). Moreover, many of the treated gametocytes appeared to have lost their outer membrane or developed membrane deformations. On the other hand, untreated gametocytes of the control group appeared perfectly healthy based on morphology and staining characteristics (Fig. 4a).

Primaquine which was tested under similar conditions showed lack of any gametocytocidal activity even at micromolar concentrations. Healthy gametocytes (both

Table 3 Early and late stage IC50 values and log IC50 values along with 95% confidence intervals (CI) for methylene blue against gametocytes produced from RKL-9 and JDP-8

| Isolate | Early stage gametocytes | Late stage gametocytes |
|---------|-------------------------|------------------------|
|         | NM | Total | NM | Total |
| RKL-9   | IC50 (95% CI) | log IC50 (95% CI) | IC50 (95% CI) | log IC50 (95% CI) |
|        | 378.5 (239.1–599) | 2.578 (2.379–2.777) | 1069 (139.7–8180) | 3.029 (2.145–3.913) |
| JDP-8   | 475.2 (318.2–709.7) | 2.677 (2.503–2.851) | 858.8 (477.5–1545) | 2.934 (2.679–3.189) |

Methylene blue demonstrated submicromolar IC50 values (NM) for both early and late stage gametocytes produced from RKL-9 and JDP-8 (All IC50 values are in nanomolar). No significant difference was observed between logarithm of IC50 values (NM and total) obtained for methylene blue against RKL-9 and JDP-8 (both early and late stage gametocytes) (p > 0.05)

Fig. 2 Dose-response curves describing relationship between concentration of methylene blue (in logarithmic scale) and percentage inhibition of a early stage gametocytes of RKL-9 b late stage gametocytes of RKL-9 c early stage gametocytes of JDP-8 d late stage gametocytes of JDP-8 (Error bars denote SD of the mean of three independent experiments). Gametocytaemia at start of drug screening experiments [In (Mean ± SD)%, RKL-9: Total G: (2.65 ± 0.36)%, Early G: (0.79 ± 0.14)%, Late G: (1.86 ± 0.23)% and JDP-8: Total G: (2.57 ± 0.35)%, Early G: (0.61 ± 0.18)%, Late G: (1.95 ± 0.21)%] where Early G and Late G represents the percentage of early (Stage II and III) and late stage (stage IV and V) gametocytes respectively and Total G represents total gametocytes inclusive of early and late stages. % Gametocytaemia calculations are per 5000 RBCs
early and late category) were observed in primaquine treated group even at concentrations as high as 25 µM (Fig. 4b). Here, gametocytes appeared perfectly non-compromised and looked similar to untreated control group (Fig. 4a).

Discussion
This work highlights the applicability of culture adapted field isolates of *P. falciparum* in anti-gametocyte drug discovery. Here, we present a simple technique to produce gametocytes in high yield from gametocyte producing field isolates, useful for gametocytocidal drugs screening applications. To date, limited data is available involving field isolates for directly testing gametocytocidal activity. This might be because very few isolates can reproducibly generate gametocytes in high yield in vitro and also show a gradual loss of gametocyte producing potential in continuous culture [23]. This makes it difficult to study the process of gametocytogenesis [23, 24] and to perform gametocytocidal drug screening due to the dependency on cryopreserved stabilates with minimum passage in order to preserve gametocyte production phenotype [13]. The two field isolates, RKL-9 and JDP-8 used in present study did not show any significant reduction in gametocyte production potential in vitro for at least 6 months in asexual culture. This gives these isolates an additional advantage over clonal lines such as 3D7 in which ability to form gametocytes wanes in as little as 2 weeks [25]. Moreover, gametocyte production potential in RKL-9 and JDP-8 appeared to be stable after

![Microscopic images demonstrating morphological deformations in methylene blue treated early gametocytes (a–c) and late gametocytes (d–l)](image-url)
multiple freeze–thaw cycles. Other studies also reported no loss in gametocyte production upon maintenance of isolates in asexual culture for 18 months [22] and also after cryopreservation [13, 26, 27]. In vitro gametocyte production potential of parasite is strain specific [28], exhibited in response to ‘nonspecific’ stress in the form of environmental stimuli [22, 23, 29]. However, the definition of ‘stress’ as well as other triggers involved in pathway shift towards gametocytogenesis in *P. falciparum* are not precisely clear [30]. The stress on the parasite is not regulated by a single component but might be a collective contribution of multitude of factors, such as high parasite load, and decrease with haematocrit [23, 31]. Studies suggest that some clones show more preference towards production of gametocytes [26, 29, 32, 33] than others under similar conditions as a result of which gametocyte production in some isolates is upregulated [34, 35]. This is also evident from data reported here, as out of 15 culture adapted field isolates, only 2 (RKL-9 and JDP-8) were able to reproducibly produce > 2% gametocytes in vitro. Moreover, asexual stages cultivated from both RKL-9 and JDP-8 were found to be chloroquine resistant. Production of higher gametocyteaemia in vitro (and in vivo) by drug resistant parasites may be correlated with tendency to spread the resistant mutation as a part of parasite’s survival strategy [34, 36]. However, current study was not designed in that context and separate studies involving more number of field isolates are needed to be carried out before a link between drug resistance and in vitro gametocytogenesis can be established.

Herein, gametocytocidal activity of methylene blue which is primarily used for treatment of methemoglobinemia is evaluated amongst Indian field isolates of *P. falciparum*. In present study as well as other in vitro studies carried out using standard laboratory strains, methylene blue was able to effectively target gametocytes, especially relatively less metabolizing stage V [8–13, 28]. However, IC₅₀ values obtained for methylene blue were inconsistent across all these studies (varied from 29.5 nM in [10] to 490 nM in [8] and 106.44 nM (late stage mean IC₅₀, present study)). In spite of differences in drug efficacies in multiple studies (might be due to variation in culture parameters including length of drug exposure, type of screening assay, and difference in parasite strain used [28]), methylene blue was effective in killing gametocytes across all these studies. In the present work, morphological alterations induced by methylene blue are described. These alterations comprise of shrinkage, distortions and membrane deformations clearly representing unhealthy gametocytes. However, it is difficult to directly correlate morphology with viability. Therefore, viability and infectivity of these morphologically deformed gametocytes remains to be evaluated. Moreover, primaquine is a gametocytocidal drug having in vivo activity [5] but data reported here identifies it as non-gametocytocidal. This disparity between efficacy data highlights the absence of any metabolic activation in vitro because of lack of liver enzymes activity required for generation of active metabolites of primaquine [37]. However, identity of the metabolites and mode of action of primaquine is not fully elucidated [38]. Although, primaquine was not expected to show any significant potency in vitro but was added in present study to validate earlier studies [8, 9] and also served as an additional negative control for methylene blue other than DMSO. A major advantage that methylene blue confers over its alternatives is that, it is the only registered non 8-aminoquinoline having late stage gametocytocidal activity, which is inexpensive and currently, a suitable alternative to primaquine. So evaluation of gametocytocidal activity of methylene blue amongst Indian field isolates of *P. falciparum* has utmost importance. However, more evidence is needed to ascertain a dose that is safe for both G6PD deficient and G6PD non-deficient population as well as effective for stopping transmission. The study sets the stage for further basic and clinical research required for consideration of methylene blue as a gametocytocidal adjunct along with standard ACT in India and developing recommendations for future use.

![Fig. 4](image-url)
Conclusions
This is the first study as far as authors know which establishes culture adapted Indian field isolates as in vitro drug sensitivity model for screening gametocytocidal compounds. It is believed by the authors that field isolates should be utilized for validation of gametocytocidal hits (obtained by high throughput drug screening experiments using reference strains) and gametocyte producing field isolates described in this study such as RKL-9 and JDP-8 might play an important role in anti-gametocyte drug discovery. Methylene blue, which is currently, a suitable alternative to primaquine as a transmission blocking drug showed remarkable gametocytocidal effect in vitro, thereby inducing morphological deformations in treated gametocytes. This study highlights gametocytocidal properties of methylene blue amongst Indian field isolates and emphasizes the utility of field isolates in gametocytocidal drug screening. Transmission blocking potential of methylene blue should be further explored in ex vivo standard membrane feeding studies using Indian field isolates as well as clinical studies amongst Indian population. This will encourage future research that will help in forming recommendations for use of methylene blue as a transmission blocking drug in India.

Abbreviations
G6PD: glucose-6-phosphate dehydrogenase; ACT: artemisinin combination therapy; NM: normal morphology; AM: altered morphology.

Authors’ contributions
IW and CRP designed the drug screening experiments. IW performed the experiments and wrote the major portion of manuscript. IW and ARA analyzed and interpreted the data. MN and NV critically reviewed the manuscript. IW and CRP designed the drug screening experiments. IW performed the experiments and wrote the major portion of manuscript. IW and ARA analyzed and interpreted the data. MN and NV critically reviewed the manuscript. IW and A5 revised the manuscript critically. All authors read and approved the final manuscript.

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Competing interests
The authors declare that they have no competing interests.

Availability of data and materials
All data generated or analyzed during this study are included in this published article.

Consent for publication
Not applicable.

Ethics approval and consent to participate
This study has been approved by Institutional Ethics Committee (IEC) of ICMR-National Institute of Malaria Research (NIMR), Delhi and bears Ethical Committee Reference Number: ECR/NIMR/EC/2016/276. Informed consent was obtained from all the participating patients before collection of infected blood samples.

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