A semi-automated luminescence based standard membrane feeding assay identifies novel small molecules that inhibit transmission of malaria parasites by mosquitoes

Martijn W. Vos1, 2, *, Will J. R. Stone1, 2, *, Karin M. Koolen1, Geert-Jan van Gemert2, Ben van Schaijk2, Didier Leroy3, Robert W. Sauerwein1, 2, Teun Bousema2 & Koen J. Dechering1

Current first-line treatments for uncomplicated falciparum malaria rapidly clear the asexual stages of the parasite, but do not fully prevent parasite transmission by mosquitoes. The standard membrane feeding assay (SMFA) is the biological gold standard assessment of transmission reducing activity (TRA), but its throughput is limited by the need to determine mosquito infection status by dissection and microscopy. Here we present a novel dissection-free luminescence based SMFA format using a transgenic Plasmodium falciparum reporter parasite without resistance to known antimalarials and therefore unrestricted in its utility in compound screening. Analyses of sixty-five compounds from the Medicines for Malaria Venture validation and malaria boxes identified 37 compounds with high levels of TRA (>80%); different assay modes allowed discrimination between gametocytocidal and downstream modes of action. Comparison of SMFA data to published assay formats for predicting parasite infectivity indicated that individual in vitro screens show substantial numbers of false negatives. These results highlight the importance of the SMFA in the screening pipeline for transmission reducing compounds and present a rapid and objective method. In addition we present sixteen diverse chemical scaffolds from the malaria box that may serve as a starting point for further discovery and development of malaria transmission blocking drugs.

The introduction of Artemisinin combination therapy (ACT) as the first line treatment for malaria has contributed substantially to recent declines in child mortality across Africa1, 2. ACT rapidly clear the Plasmodium life stages responsible for clinical disease but at the dose used to clear asexual blood stage parasites have only limited effect against mature gametocytes, the stage transmitted to mosquitoes3. Though immature gametocytes may be cleared shortly after their production4 resulting in reduced gametocyte burden and transmission potential5, current dosing regimens of ACT do not radically clear mature transmission stages and do not prevent transmission shortly after treatment6. Expanding efforts to achieve local malaria elimination and contain transmission in areas of emerging ACT resistance have stimulated interest in combining standard treatment with compounds active against mature gametocytes7 and have highlighted the need to screen new antimalarial drug candidates for their transmission reducing activity (TRA).

Currently the only available drug that radically prevents malaria transmission is primaquine8, but its wide-scale deployment is hampered by issues of safety and dosing9,10. Though the results of numerous ongoing trials give reason to expect wider scale use of single-low dose primaquine for reducing P. falciparum transmission, there is growing interest in identifying novel transmission reducing compounds7.

For screening new compounds, several assays determining gametocyte viability activity have been developed. These have been based on the expression of metabolic markers or reporter genes, or more recently on gametocyte
activation in vitro measured by the presence of female and male gametocytes. Such assays are, however, not always predictive of blocking transmission of the parasite to the mosquito, and agents with putative TRA are best tested in the standard membrane feeding assay (SMFA). In the SMFA cultured gametocytes are mixed with blood and test compounds, and this mix is fed to laboratory reared mosquitoes to determine the effect of the compound on the parasites establishment in the mosquito. As the outcome of the SMFA is mosquito infection and the timing and duration of compound exposure is controlled, the assay is easily adapted to assess effects active at any stage of parasite development – from gametocyte inhibition, to inhibition of sporogony. The SMFA has numerous pitfalls though, first among these being the need to dissect mosquitoes and count oocysts on the mosquito midgut by microscopy; a labour intensive and subjective task, which imposes significant limits on the assays scalability. We previously described the use of a parasite line expressing a fusion of the green fluorescent and firefly luciferase proteins (NF54HT-GFP-Luc) in a new SMFA format based on the measurement of luciferase activity in groups of homogenised individual or pooled mosquitoes. Although very effective, this strain may be sub-optimal for drug screening; introduction of the human dihydrofolate reductase (hDHFR) selection marker during integration of the GFP-Luc expression cassette renders it resistant to antifolates and makes it necessary to maintain exposure to antifolate compound WR99210 during parasite culture to prevent reversion to wild-type (WT), possibly modifying its response to other inhibitors by affecting parasite metabolism. Here we report the development of a parasite line that constitutively expresses GFP-Luc and is free of limiting selection markers, for use in an efficient, dissection-free SMFA screen. Our approach used frozen mosquitoes and machine homogenisation in a 96-well format to improve the efficiency of indiscriminate drug screening. After validation and comparison of luciferase expression with microscopy based oocyst counts we used our luminescent SMFA to screen 47 marketed and experimental antimalarials and compared the results with data from the gametocytocidal lactate dehydrogenase (pLDH) activity assay and Pfs25 AlphaLISA assay. Furthermore, the luciferase screen was used to evaluate the TRA of 18 compounds selected from the Medicines for Malaria Venture (MMV) malaria box specifically for their potential activity as gametocytocidal. In this final screen, gametocytocidal or permanent inhibitory activity was confirmed by incubating gametocytes with compounds before washing out the compound prior to mosquito feeding, while specific activity against the parasite stages emerging in the mosquito gut was confirmed by adding compounds to the gametocyte culture immediately prior to mosquito feeding. These assays identified novel transmission blocking molecules and provided new insight into the predictive value of in vitro metabolic and gamete formation inhibition assays, showing they can miss potent compounds, and highlighting the advantages of the gold-standard SMFA for screening transmission blocking compounds. Importantly, our more scalable luciferase based read-out increases throughput of the assay and makes compound screening with the SMFA less resource intensive than when performed with microscopy.

Results

Generation of a GFP-Luc expressing parasite line free of selection markers. To allow unrestricted evaluation of new compounds we developed a new parasite line, free of drug selection markers (full title: NF54-DP47-5’hsp70-GFP-Luc; hereafter: NF54-HGL), that stably expresses a GFP-Luciferase fusion protein under control of the constitutive P. falciparum hsp70 promoter (Fig. 1A,B).

To confirm the proposed effect of removal of the hDHFR selection marker on antifolate sensitivity, we compared sensitivity of wildtype NF54, intermediate clone NF54-HGL-hDHFR (created prior to FLP recombinase-mediated excision of the hDHFR maker) and NF54-HGL asexual blood stage parasites to dihydroartemisinin (DHA), chloroquine, and two antifolate drugs: WR99210 and Pyrimethamine. The half maximal inhibitory concentration (IC50) for DHA and chloroquine are comparable for all three parasite lines. As predicted, presence of the hDHFR selection marker in NF54-HGL-hDHFR renders this parasite resistant to WR99210 and Pyrimethamine. Excision of the hDHFR open reading frame effectively restores sensitivity to these antifolates for NF54-HGL (Fig. 1D–G). Since pLDH is a stable enzyme and antifolates have relatively slow parasite killing kinetics, the inhibition by WR99210 and pyrimethamine did not reach 100% within the timeframe of the assay.

Assessment of GFP-Luc expression. Analyses of luminescence signals confirmed expression of the GFP-Luc protein in asexual blood stage parasites (Fig. 1C). In line with previous reports on different expression levels in the various life cycle stages of the parasite, expression was higher in day 8 gametocytes than in mature gametocytes and asexual blood stages. To assess GFP-Luc expression during the parasites development in mosquitoes, Anopheles stephensi were fed a blood-meal containing NF54-HGL gametocytes and dissected at various time-points post infection (PI) to allow visual inspection of the mosquito midgut. No fluorescence was observed during ookinete formation (24 h PI, data not shown), indicating that GFP expression levels may have been under the limit for microscopic detection. From day 6 PI positive oocysts were easily observable (Supplementary Fig. S1). Luciferase expression was weak during early sporogonic development (2–6 days PI, data not shown), but robust signal in low intensity infections (mean oocysts = 1.28, range = 0–7) was detectable from day 7 PI onwards (Supplementary Fig. S2). Uninfected negative control mosquitoes showed low levels of luminescence (n = 617; mean RLU = 3.2 RLU; standard deviation = 1.9 RLU). Day 8 PI showed a clear separation of positive and negative mosquitoes and was chosen for subsequent assessments of infection intensity (i.e. number of oocysts per midgut) and prevalence (i.e. percentage of mosquitoes with oocyst stage infection) based on luminescence intensity.

Microscopy and luciferase based assessments of infection and TRA. To confirm that measurements of luciferase activity can replace manual dissection-based oocyst counts, as was previously shown for NF54HT-GFP-Luc, 172 experimental mosquito feeds (hereafter ‘feeds’) were performed using NF54-HGL gametocyte culture. From each feed, separate mosquito samples were assayed by standard microscopy and by luminescent SMFA (microscopy, n = 2557; luminescence, n = 3570). Thirty-two of these feeds were performed as controls allowing for direct comparison of infection intensity and prevalence based on oocyst and luminescence
readouts. These were performed either without test compounds (0.1% DMSO, n = 16) or with control compound DHA that shows gametocytocidal activity at high concentration (10 μM DHA, n = 16)13. Mean oocyst counts for control feeds were between zero and 29 oocysts (mean 4.9 oocysts), and had a positive linear relationship with mean luminescence intensity (R² = 0.86, p < 0.0001). Similarly, oocyst prevalence was positively associated with luminescence based prevalence (R² = 0.95, p < 0.0001) (Supplementary Fig. S3). Combining all mosquitoes from control feeds, prevalence estimates for all mosquitoes from control feeds were 43.33% (208/480) for microscopy, and 49.22% (350/711) for luminescence.

To further validate the luminescent SMFA we tested seven compounds from the MMV validation box in full dose response in an ‘indirect’ luminescent SMFA format. In the indirect assay, gametocytes were incubated with compounds for 24 hours, and mosquitoes were fed a blood-meal containing this gametocyte culture/compound mix. Microscopic examination of oocyst densities in midguts and luminescence measurements were performed on distinct cohorts of mosquitoes from the same cage. As shown in Fig. 2, all compounds dose-dependently reduced oocyst formation and luminescence intensity. Data were fitted to a Hill equation using Maximum Likelihood Estimation to find the best fit. The dose response curves fitted on the microscopy and luminescence data aligned very well for artemisone, atovaquone, mefloquine, NPC1161B, pyrimethamine and pyronaridine. The curves for lumefantrine appeared to be less well aligned due to a distinct dispersion of oocyst counts and RLU values at the highest concentration tested. For all other compounds the derived IC50 values differ by no more than 2–3 fold when comparing the two read-outs.

The luminescent SMFA identifies compounds with different modes of TRA. Using only luminescence readouts, we screened 47 established antimalarial compounds in the indirect luminescent SMFA format at a 5 μM concentration. Samples of 22–24 mosquitoes were analysed for each compound (n = 46*24; 1*22). The results of this rapid screen are shown in Fig. 3A, with TRA (%) calculated using generalised linear mixed models (GLMM) from differences in the intensity of infection between test and vehicle (0.1% DMSO) controls. Error bars indicate standard deviations.

Figure 1. (A) Schematic of the gene targeting strategy to insert a GFP-Luc reporter gene under control of the hsp70 promoter in the pf47 locus, showing the genomic locus of the wild-type (WT) and gene deletion mutants Δpf47 containing the GFP-luciferase ORF before and after removal of the hdhfr resistance marker. P1, P2: primers MWV300 and MWV301, respectively, used for PCR analysis; X (XcmI), N (NcoI): restriction sites used for conformation of LR-PCR analysis; pf47: pf47 locus; hsp: heat shock protein; hdhfr: human dihydodrofolate reductase coding region; GFP-Luc: GFP-luciferase fusion protein; fcu: cytosine deaminase/uracil phosphoribosyl transferase; pbdt: P. berghei dhfr transcription termination region; kb: kilo basepairs. (B) Verification of the integration site by PCR and restriction digestion. Genomic DNA fragments from the parental NF54 strain (WT), intermediate transgene NF54-HGL-hDHFR and final transgene NF54-HGL were amplified by PCR using primers MWV300 and MWV301 (P1 and P2) and analysed by DNA gel electrophoresis. Where indicated, the PCR fragment was subjected to restriction digestion with XcmI/NcoI (X/N) (C) Luminescence intensity (relative light units, RLU) as a function of cell number. The figure shows a comparison between uninfected red blood cells (RBC), immature gametocytes (Day 8 GCT) and mature gametocytes (Day 14 GCT) and asexual blood stage parasites. (D–G) FLP-FRT mediated excision of the selection marker restores sensitivity to antifolates. Drug sensitivity analysis of WT, ΔPf47GFP-luc-hdhfr and ΔPf47GFP-luc to DHA (D), Chloroquine (E), Pyrimethamine (F), and WR99210 (G) in a dose dependent asexual growth inhibition assay using pLDH enzyme activity as a readout. Depicted data represents a triplicate measurement and was normalized to the MIN (10 μM DHA) and MAX (0.1% DMSO) controls. Error bars indicate standard deviations.
Thirty-four compounds had statistically significant TRA based on reductions in luminescence intensity; 21/34 with ≥ 80% TRA.

To provide more information on the mechanism of active compounds, all compounds showing ≥ 80% TRA on the intensity scale in the indirect assay were re-tested, this time adding compounds to the blood/gametocyte culture mix immediately prior to mosquito feeding (hereafter, the ‘direct’ assay variant). Fourteen compounds had >80% TRA in the indirect assay but not in the direct assay, suggesting the absence of strong activity against
the parasite life stages occurring in the mosquito gut. Amino alcohol halofantrine, antifolates P218, pyrimethamine and chlorproguanil, synthetic dye methylene blue, BC1 inhibitor atovaquone, and protein synthesis inhibitor cycloheximide had comparable TRA in the indirect and direct SMFA formats, indicating activity against mosquito stage parasites, but not excluding the possibility of concurrent gametocytocidal action.

**The luminescent SMFA identifies compounds with TRA that are missed by gametocyte and gametogenesis assays.** The results of the indirect and direct assays were compared with previously described data showing the effect of the same 47 compounds at 5 μM in a gametocyte viability assay with pLDH readout, and a homogeneous immunoassay detecting the Pf525 antigen on emerging female gametes (Fig. 3B). At a threshold of ≥80% inhibition, the pLDH and gametogenesis assays identified 6/47 and 13/47 compounds respectively as having gametocytocidal activity. The activity of all but one of the six compounds identified by the pLDH assay (napthoquine) was confirmed by the gametogenesis assay, while all hits from either viability assay were confirmed in the indirect luminescent SMFA. However, high TRA was observed for a number of compounds where the inhibition assays showed low activity, indicating that some compounds had activity only identifiable with transmission as the definitive readout. Total false negative rates for the pLDH and gametogenesis assays were 32% (15/47) and 17% (8/47), respectively. Excluding compounds that blocked transmission in the direct luminescent SMFA, which we presumed to have activity downstream of the gametocyte, the SMFA still identified a larger number of transmission-blocking compounds than either gametocytocidal assay, with false negativity rates of 25% (10/40) in the pLDH assay and 10% (4/40) in the gametogenesis assay. This is not surprising since the SMFA covers a higher biological content than either of the other two methodologies.

**Identification of novel transmission-blocking small molecules.** In recent years, a number of gametocyte assays have been developed that attempt to predict gametocyte viability and infectivity based on metabolic activity, reporter gene expression, or changes in cell shape or antigen expression. In an effort coordinated by the Bill & Melinda Gates Foundation and the Medicines for Malaria Venture (MMV), a number of these and additional novel assays selected a subset of compounds from the MMV Malaria box as potentially transmission blocking. All of these molecules kill asexual blood stage parasites with submicromolar IC50s (Table 1). Here, we evaluated 18 of these selected compounds in full dose-response experiments using the luminescent SMFA. In order to confirm a gametocyte-based mechanism of action, compounds were washed out following incubation with stage V gametocytes to prevent activity against parasite stages emerging in the mosquito midgut. Effectiveness of the washing step was tested by evaluating the activity of pyrimethamine, which has potent activity against the parasite stages that develop in the mosquito midgut (Fig. 2) but not against gametocytes. Indeed, washing a culture of gametocytes exposed to pyrimethamine restored subsequent oocyst development in the mosquito, confirming that this format of the assay is likely restricted to the analyses of gametocytocidal modes of actions (Supplementary Fig. S4). This might however be different for more lipophilic drugs, which might be washed out with less efficiency. Sixteen out of the 18 Malaria box compounds tested in this wash-out mode of the SMFA dose-dependently reduced oocyst development in the mosquito midgut (Fig. 4). Compounds MMV665941, MMV667491, MMV019918 and MMV665827 were among the most potent transmission-reducing molecules, with IC50 values of 0.04, 0.06, 0.07 and 0.1 μM, respectively. To reveal whether these compounds act against the parasite stages that develop in the mosquito midgut on top of their gametocytocidal activity, all compounds were tested in the direct assay variant at the same maximum concentration as in the wash-out assay (10 μM), confirming that 10/18 compounds significantly reduced mosquito oocyst intensity. Compounds MMV020492 and MMV665882 that did not show gametocytocidal activity were equally inactive in the direct SMFA. To further address the mechanism of action of compounds with functional TRA, the SMFA data were compared to data from published studies that addressed different aspects of gametocyte biology (Table 1). MMV019918 was the only compound consistently showing activity across the different assay formats, although IC50 values ranged from 0.07 μM to 0.9 μM, with large differences between the two assays that interrogated the ability of a gametocyte to undergo exflagellation. Conversely, MMV007116 and MMV665827 showed potent activity in the SMFA but very low activity in the published gametocyte assays, with the exception of a luciferase reporter assay that revealed the effect of these compounds on gametocyte viability. In general, the luminescent SMFA identified more compounds with functional TRA than predicted by each of the individual published gametocyte assay formats.

**Discussion**

The firefly luciferase expressing parasite line NF54-HGL enables the SMFA to be performed for indiscriminate compound screening with a semi-automated luciferase based read-out. The results of the current study demonstrate the versatility of the SMFA for screening compounds with activity against different transmission stages of the malaria parasite, and validate the luminescent assay format in combination with microtiter plate-based processing of mosquitoes for improving the efficiency of the biological gold-standard for assessing *Plasmodium* transmission in controlled experiments.

We presented a screening pipeline for the SMFA that allowed us to investigate drug mechanism of action while retaining transmission as the assay end-point. The indirect assay, in which compounds were incubated with gametocytes for 24 hours prior to mosquito feeding, represents the simplest and most sensitive screen and the first step in the pipeline for screening large compound libraries. The direct assay, in which the compound is added at the moment the gametocyte activates to form a gamete, interrogates compound action in the mosquito midgut. Pharmacokinetics in the mosquito are largely unknown, but detoxifying enzymes may limit the compound’s action in the midgut, restricting the direct SMFA to monitoring effects on gametogenesis, ookinete formation and perhaps early sporogony. Lastly, the wash-out format of the SMFA addresses a mode of action that is likely restricted to the gametocyte. High-throughput screening is impossible with the SMFA, but the luminescent SMFA represents a significant improvement in the techniques scalability. The replacement of direct oocyst counts with
Table 1. Compound activity as determined in the luminescent SMFA compared to activities reported by published gametocyte viability assays and asexual replication assays. The table shows IC50 values where available (NA: not available) or the percentage inhibition at the highest concentration tested, as indicated in the header row. IC50 data for the exflagellation (imaging) and Pfs25 assays were generated in a washout format, whereas percentage inhibition values represent data from a continued exposure experiment. For SMFA data, 95% confidence intervals are indicated between brackets where possible. For MMV019881, and MMV665941 the model did not converge. Asterisks (*) indicate statistically significant inhibition (p < 0.05) in the direct SMFA. The last column shows activity against asexual blood stages of *P. falciparum* strain 3D7 as deposited at the ChEMBL repository under accession number CHEMBL2028071. ND: no dose response relationship was detected in the data.

| Compound | IC50μM (95% CI) | slope (95% CI) | % inh. 10μM | exflagellation\(^2\) (imaging) IC50μM (%) | Pfs25\(^3\) IC50μM (%) | alamarBlue\(^5\) IC50μM (%) | Imaging\(^6\) IC50μM (%) | exflagellation\(^7\) (SYBR green) IC50μM (%) | asexual blood stage \(^8\) |
|----------|-----------------|----------------|-------------|--------------------------------------|------------------------|----------------------------|----------------------------|----------------------------------------|-------------------------------|
| MMV000248 | 1.64 (1.26 – 2.13) | -2.44 (-3.07 – -1.81) | 48.76\(^9\) | NA (-66%) | NA (8%) | 0.31 | 1.1 | NA (95%) | 0.72 |
| MMV000442 | 0.59 (0.35 – 1.01) | -1.18 (-1.58 – -0.78) | 5.50 | NA (12%) | NA (6%) | NA (72%) | 0.92 (55%) | NA (59%) | 0.36 |
| MMV000448 | 1.18 (1.07 – 1.31) | -1.63 (-1.591 – -7.34) | 97.93\(^9\) | 0.25 | 4.02 | 1 | 0.71 | 5.4 | 0.23 |
| MMV007116 | 0.11 (0.06 – 0.19) | -1.62 (-2.15 – -1.08) | 99.22\(^9\) | 20.75 | NA (10%) | NA (40%) | 0.26 | NA (67%) | 0.35 |
| MMV011438 | 8.58 (7.95 – 9.36) | -8.47 (-12.62 – -4.32) | -3.29 | NA (-77%) | NA (-31%) | 0.82 | 1.13 | NA (69%) | 0.32 |
| MMV019266 | 1.29 (1.21 – 1.38) | -9.38 (-12.41 – -6.35) | 7.07 | NA (31%) | NA (-11%) | 0.31 | 0.33 | NA (81%) | 0.62 |
| MMV019881 | 1.17 (–) | -15.81 (–) | 88.15 | NA (57%) | NA (7%) | NA (76%) | NA (32%) | 5.5 | 0.65 |
| MMV019918 | 0.07 (-0.13) | -1.36 (-1.80 – -0.93) | 88.55\(^9\) | 0.07 | 0.52 | 0.32 | 0.7 | 0.9 | 0.79 |
| MMV020492 | ND | ND | -16.63 | NA (36%) | NA (-12%) | NA (70%) | NA (-5%) | NA (59%) | 0.03 |
| MMV396797 | 3.62 (8.17 – 15.18) | -4.10 (-9.87 – 1.67) | -16.44 | NA (-14%) | NA (-17%) | NA (90%) | 0.01 | 8.8 | 0.48 |
| MMV665827 | 0.10 (-0.05 – 0.15) | -2.23 (-3.05 – -1.42) | 98.25\(^9\) | 16.55 | NA (-15%) | NA (41%) | 0.34 | NA (66%) | 0.12 |
| MMV665882 | ND | ND | 4.92 | NA (-12%) | NA (-48%) | NA (92%) | 0.07 | NA (75%) | 0.47 |
| MMV665941 | 0.04 (–) | -10.29 (–) | 99.84\(^9\) | 0.42 | 0.3 | NA (86%) | 0.32 | 1.8 | 0.26 |
| MMV665971 | 3.22 (1.45 – 7.15) | -1.50 (-2.57 – -0.44) | -7.04 | NA (-29%) | NA (-13%) | NA (73%) | 0.23 | NA (61%) | 0.49 |
| MMV665980 | 1.76 (1.57 – 1.96) | -11.52 (-25.29 – 2.24) | 64.17 | 8.07 | NA (74%) | NA (27%) | NA (91%) | NA (76%) | 0.21 |
| MMV666021 | 1.25 (1.17 – 1.34) | -13.30 (-17.52 – -9.28) | 94.02 | NA (30%) | NA (-20%) | NA (58%) | 0.61 | NA (68%) | 0.09 |
| MMV666125 | 1.40 (1.11 – 1.76) | -2.86 (-3.97 – -1.76) | -24.69 | NA (-3%) | NA (-17%) | 0.48 | 0.75 | NA (74%) | 0.38 |
| MMV667491 | 0.06 (-0.03 – -0.11) | -0.61 (-0.73 – -0.50) | 89.05\(^9\) | 0.09 | 0.18 | NA (91%) | 1.06 | 4.5 | 1.23 |

Luciferase measurements from homogenised mosquitoes in the SMFA was described previously for parasite line NF54HT-GFP-luc, but aspects of the methodology described here differ. The current reporter parasite is devoid of the DHFR selection marker, allowing for precise description of the effects of antifolates including pyrimethamine on oocyst development, and expediting the evaluation of a wide variety of compound libraries. Mosquito sample sizes were limited to increase throughput, and freezing mosquitoes prior to further processing increased flexibility of laboratory operations. Furthermore, semi-automated processing of mosquitoes significantly reduced the time needed for assessment of infection status. In all, the luminescent format for individual mosquitoes reduces hands-on time 2-fold in comparison with the microscopical readout, which is a tremendous step up from a previously published ‘industrialized’ SMFA format. For even further increased throughput, screening in the luminescent SMFA may be performed by pooling larger numbers of mosquitoes, achieving an approximate four-fold increase in throughput over the standard microscopical readout if only infection intensity is the desired output.

Our data from a large number of infections indicate that oocyst intensity is a simple linear function of luminescence intensity. This is in keeping with our previous work that showed a linear relationship between oocyst counts and luminescence intensity, using a different promoter to drive expression of the reporter gene. Oocysts are an intermediate parasite life-stage which even when developing concurrently in a host mosquito may differ in size and sporozoite content. Whereas microscopy produces discrete data, luminescence measurements provide continuous data that reflect actual numbers of developing sporozoites, which may be more directly relatable to mosquito infectivity, for which oocyst counts are an accepted proxy. An added advantage of the luminescent assay is that it is less subjective than microscopy based assessments of oocyst numbers. We cannot exclude the possibility that in heavily infected mosquitoes, with higher oocyst intensities than those reported here, nutrient and space competition may lead to lower *per capita* sporozoite production, resulting in a non-linear relationship between luminescence intensity and lower luminescence based TRA estimates than would be determined from oocyst number alone. Though we advise caution in extrapolating actual oocyst number from luminescence values, we believe luminescence based TRA estimates may be more accurate than those made using oocyst data.

In the direct mode of the SMFA the compound remains present in the bloodmeal, and there is a remote possibility that it interferes with activity of the luciferase reporter rather than with oocyst development. We have been unable to investigate the potential effect of compounds interacting with *hsp70* expression, as has been observed in other eukaryotes, or the possibility that compounds may directly inhibit luciferase protein after its production.
We consider both events highly unlikely to affect the luciferase readout in the current SMFA format, as compounds are at latest added to the mosquito bloodmeal 8 days before luminescence measurements. Furthermore, analyses of a subset of the MMV malaria box compounds in a luciferase inhibition assay did not reveal inhibitory activity at the concentrations that reduce oocyst development. The possibility that some compounds may effect mosquito metabolism, and by proxy parasite developmental success, can also not be excluded. Readouts based on oocyst number may be insensitive to such subtle effects, while the more quantitative luminescence readout may not be. Additional experiments will be required to elucidate the interaction of mosquito and parasite fitness in relation to compound exposure.

Figure 4. Dose response experiments for 18 compounds from the MMV malaria box, assessed in the luminescent SMFA in a wash-out assay variant. All compounds were tested in duplicate in two independent experiments. The figure shows oocyst intensities relative to vehicle control (0.1% DMSO) as a function of concentration for the compounds indicated at the top of the panels.
Apart from oocyst intensity, oocyst prevalence (the proportion of mosquitoes that develop an oocyst-stage infection), is an important parameter determining malaria transmission. Whereas inter-experimental variation in control mosquito oocyst burden has no effect on TRA expressed as a reduction in oocyst intensity, it heavily influences the outcome based on a reduction in oocyst prevalence13. Since our experiments comprised a large set of experimental infections that each showed a distinct distribution of oocyst densities we have, therefore, limited our analyses to a comparison of compound effects on oocyst intensity, as this is the only parameter that is directly comparable between experiments. Nevertheless, a complete assessment of the transmission blocking potential of an antimalarial would require a description of the effects on infection prevalence, preferable at infection intensities that mimic those observed in naturally infected mosquitoes36. An experimental and computational strategy to achieve such an assessment has been implemented in our laboratory and will be described elsewhere.

Compounds reducing malaria parasite transmission to mosquitoes may kill gametocytes directly (gametocytocidal activity), irreversibly impair the parasites ability to develop once ingested by the mosquito, or else inhibit gametocyte developmental viability only while drug pressure is maintained in the mosquito environment12. Because mature gametocytes persist at low densities for several weeks after ACT treatment4, and because submicroscopic gametocyte densities may be highly transmissible57,58, efforts to develop transmission-reducing compounds are focused on identifying activity against mature gametocytes59. In vitro assays based on markers of metabolic activity (pLDH13,19, Alamar Blue reduction25) or reporter genes21,22,40 can be used to assess the effects of gametocytocidal compounds but may overlook compounds that manifest their action downstream of the gametocyte. Gametocyte infectivity may be assessed in greater detail with gamete formation assays, which can include gametocyte washing steps to discriminate between reversible and irreversible compound activity12,13,23. In vitro ookinete formation assays can also be used to identify compounds whose presence in the mosquito blood meal inhibits parasite fertilisation60, but these assays cannot exclude the possibility of compounds reducing ookinete viability or oocyst formation. Assays measuring the viability or development of gametocytes without mosquito feeding are comparatively inexpensive and highly scalable, so will continue to be required as a prioritization step in the screening pipeline for transmission blocking drugs. The data presented here indicate, however, that increased throughput of in vitro gametocyte assays goes at the cost of a considerable number of false negatives. Importantly, as no in vitro assay or combination of assays is able to fully interrogate the effect of compounds on all parasite developmental stages exposed to test compounds, our results underline that the SMFA remains the gold standard assessment of gametocyte infectivity. Our indirect screen of a varied set of 47 antimalarials identified compounds preventing mosquito infection by mechanisms targeting any stage of the parasites sexual development or early sporogony. Assays based on parasite metabolic processes (pLDH) and on the expression of female gamete upregulated protein Pfs25 cumulatively failed to identify seven compounds which, incubated with cultured gametocytes (NF54–HGL) in the same concentration, caused marked reductions in oocyst stage productivity as assessed by the luminescent SMFA. Though we anticipated identifying compounds with additional activity downstream of gamete formation in the SMFA, the results of the direct assay indicate that the pLDH and gamete formation assays failed to identify compounds with effects restricted to the gametocyte; three of the seven false negatives had no activity on mosquito stage parasites in the direct SMFA format. The effect of the remaining four compounds may rely on a sporontocidal rather than a gametocytocidal activity. In keeping with previous reports of their sporontocidal activity (reviewed in Butcher61), pyrimethamine and atovaquone blocked oocyst development but were not active in the pLDH and gamete formation assays. Atovaquone has demonstrated a long-lasting transmission-blocking effect in ex vivo studies with sera from drug-treated volunteers62, but the effect on transmission in malaria-endemic settings has not been described in detail. The sporontocidal effect of pyrimethamine most likely represents a class effect of dihydrofolate reductase (DHFR) inhibitors, as it was also observed for P218 and chlorproguanil. Proguanil was understandably not active in the SMFA as its only active metabolite cycloguanil binds to DHFR63.

Analyses of eighteen compounds from the MMV malaria box revealed sixteen compounds with functional TRA that were not necessarily identified by individual gametocyte based assay formats12,24–26. These 18 compounds were flagged as potential transmission blockers by a joint effort coordinated by the Bill and Melinda Gates Foundation and the Medicines for Malaria Venture under the ‘Gametocyte Assays for Plasmodium falciparum’ program. Under this program, five consortia consisting of two to five malaria laboratories each screened the MMV malaria box in a variety of gametocyte-based assays. All consortia proposed a shortlist of active molecules for further evaluation in the SMFA. The final selection of 18 molecules, governed by MMV, was then based on potency, diversity in chemical scaffolds and presumed tractability. As these compounds were expected to act at the level of the gametocyte, we applied a wash-out format to restrict the analysis to the sexual blood stage of the parasite when testing these compounds in the luminescent SMFA. For the 18 compounds subjected to this more rigorous investigation, 16 were able to reduce onward oocyst development in the mosquito with potencies ranging from 8.6 to 0.04 μM. The luminescent SMFA did not reveal TRA for MMV020492, which is in line with its lack of activity in published gametocyte assays. Equally inactive in the SMFA, MMV020492 showed a lack of activity in most published gametocyte assays, with the exception of a luciferase assay that indicated an IC50 of 62 nM. Overall, from a qualitative perspective the luciferase assay showed the best agreement with the SMFA, as it identified 14 out of the 16 compounds with functional TRA. Assays that interrogate the ability of a gametocyte to form a male or female gamete did not identify MMV000442 as a potential transmission reducing compound12, whereas it showed an IC50 of 590 nM in the SMFA. This indicates that gametocyte infectivity is defined by a greater set of functions than described by its sole ability to form a gamete. Quantitatively, neither of the published gametocyte assays correlated with efficacy. Although mouse models have been proposed to determine the efficacious dose for blockage of malaria transmission, these are restricted to P. berghei parasites and have limited value for predicting efficacy on human
malaria. In addition, *P. berghei* infection intensities are much higher than *P. falciparum* infection intensities, which leads to an underestimation of TRA when estimated from differences in infection prevalence\(^5\)\(^6\).

Our studies identified four compounds from the malaria box with IC50 values below 100 nM in the luminescent SMFA (Fig. 5). Compounds MMV665827, MMV665941, and MMV667491 lack structural properties that meet Lipinski’s rule of five that predicts oral absorption of a candidate drug\(^6\). Therefore, these compounds are considered tool compounds rather than drug-like molecules\(^6\). Nevertheless, these compounds may have value in the identification of mechanisms underlying transmission-blocking activity, which remain poorly understood. Importantly, these molecules have activity against asexual blood stage parasites, which opens the possibility of selecting drug resistant mutants followed by whole genome sequencing. This strategy has been successfully employed in the identification of PI4K, and eEF2 and confirmation of DHODH as bona fide drug targets\(^48\)\(^50\). From the group of compounds with potent transmission blocking activity, MMV19918 appears to qualify as a drug-like molecule and may, besides serving as a tool in unravelling mechanism of action, act as a starting point for further optimization. This compound appeared 10-fold more active in the SMFA as in assays interrogating activity against the asexual blood stages. Such a transmission-selective profile was also observed for MMV665941 and MMV667491. To the best of our knowledge these are the first molecules described to date that show selectivity towards the transmission stages of the parasite. Recently, the malaria drug development pipeline was enriched with molecules like OZ439, DSM265 and DDD107498, which unlike the majority of marketed antimalarials have activity against the transmission stages of the parasite\(^51\). However, for all these molecules this activity is at par or less potent than the activity observed against asexual blood stage parasites\(^13\)\(^49\)\(^50\)\(^52\). The observation of a transmission-selective profile in the study presented here suggests a novel mechanism of action outside the range of targets described to date. In view of the need for molecules that address a target candidate profile for prevention of transmission\(^53\), these mechanisms and the molecules identified here warrant further investigation.

**Conclusion**

The results of the current study demonstrate the sensitivity and flexibility of the SMFA for determining the effect of small molecules on *P. falciparum* gametocyte transmission to mosquitoes, while validating a method that increases the assays scalability and ease of use. Comparison to gametocyte inhibition assays indicates that parasite development in the mosquito may be required to ensure assay sensitivity to transmission-blocking small molecules. Of the 18 compounds selected in this analysis based on the results of previous gametocyte assays, our data indicate that 3 compounds have greater activity on transmission stage parasites than asexual stage parasites. These molecules may be of particular interest for further drug discovery and elucidation of novel drug targets.

**Methods**

**Parasites and culture.** Asexual blood stages of all *P. falciparum* parasite lines were cultured in RPMI 1640 medium supplemented with 367 μM hypoxanthine, 25 mM HEPES, 25 mM sodium bicarbonate and 10% human type A serum, in a semi-automated system under standard conditions and induction of gametocyte production was performed as previously described\(^54\)\(^56\).

**Generation of NF54-HGL-hDHFR and NF54-HGL parasite lines.** The GFP::LUC fusion protein under control of the *hsp70* promoter was stably integrated into the genome of asexual stage NF54 (WT) parasites at the *pfs47* locus (PF13_0248) by standard double crossover recombination and positive/negative selection\(^57\)\(^59\). The targeting plasmid was based on the gene deletion construct pHHT-FRT-PF5236 described previously\(^46\). The 5′ *pfs47* target region in pHHT-FRT-PF5236 was replaced by the 3′ *pfs47* target region amplified by PCR using primes MW257 and MW259 (primer sequences in supplemental Table S1) and introduced in the plasmid by *MluI*/SacII restriction digestion and ligation. The 5′ p52 region was replaced with the 5′ *pfs47* target region amplified by PCR using primes MW262 and MW267 and introduced in the plasmid by *Xmal*/NarI restriction digestion and ligation. A reporter gene cassette was constructed by subcloning a GFP::LUC fragment from plasmid PL1063\(^46\) in pCR2.1-BLUNTII-TOPO after *AfIII/XbaI* restriction digestion. An *Agel* restriction site was introduced immediately upstream of the start codon by introducing a PCR fragment amplified with primes MW252 and MW253 through *KpnI/Hpal* restriction digestion and ligation. The promoter region of the *hsp70* gene (PF3D7\(_2\)0818900) was placed upstream of GFP through *sacII/Agel* restriction digestion and ligation after amplification with primes MW198 and MW200 and subcloning in pEGFP-N1 (Clontech laboratories, Mountain view, CA) through *BamHI/SacII* digestion and ligation. The 3′ transcription termination region from the *hsp86* gene (PF3D7\(_2\)0708400) was introduced following PCR amplification using primes MW268 and MW271 and *XbaI/Apal* restriction.
digested, followed by ligation and finally the reporter cassette was introduced into the parasite genome using the plasmid by *KpnI/NgoMIV restriction digestion and ligation yielding plasmid pMV163.

Pls/a target regions, 5 hsp70 and 3 hsp86 fragments were amplified by PCR amplification (Phusion, Finnzymes) from genomic *P. falciparum* DNA (NF54 strain) and all PCR fragments were sequenced after TOPO TA (Invitrogen) sub-cloning.

Transfection of wild type NF54 asexual parasites with DNA construct pMV163 was performed as described previously.[59] using a Cell Electroporation 600 (BTX, Holliston, USA) Double crossover mutants were selected by positive and negative selection as described,[67] yielding parasite line NF54-HG1-HDFR. Cloning of transgenic parasites was performed by limiting dilution in 96-well plates as described.[62] Parasites in positive wells were transferred to the semi-automated culture system and cultured for further phenotype and genotype analyses.

Removal of the hdhfr selection marker was next performed by transfection of the mutants with construct pMV-FPLec and selection on blastidin.[60, 63] resulting in parasite line NF54-HGL. Integrity of the transgen was analysed by PCR using primers MVW300 and MVW301 and XcmI/NcoI restriction digestion.

To confirm functional FRT-mediated excision of the hDHF selection marker, sensitivity to anti-folates and a number of non-folate reference compounds were performed on NF54 (WT), NF54-HG1-HDFR, and NF54-HGL using a modified pLDH assay. Briefly, non-synchronised asexual parasites (~50% ring stages, 25% trophozoites and 25% schizonts) were seeded at a density of 0.5% parasitaemia in 3% haematocrit in a black 96-well plate in 50 µl of culture medium, with 50 µl of diluted test compound. Following a 72 hour incubation at 37 °C, 98% humidity, 93% N2, 4% CO2 and 3% O2, the increase in parasitaemia was assessed using the modified pLDH assay as described previously.[13] Assays were performed in triplicate (three serial dilutions for each compound) and all plates included triplicate controls to assess minimum (MIN control, 1 µM DHA) and maximum (MAX control, 0.1% DMSO) signal. To compare the three different strains data were normalized and expressed as the percentage reduction in replication relative to the blocking and non-blocking controls.

**GFP fluorescence analysis.** The midguts of mosquitoes fed previously with NF54-HGL gametocyte culture were mounted on glass slides in a droplet of PBS, and GFP expression was visualized and photographed on a Leica fluorescence microscope with digital camera.

**Standard Membrane Feeding Assays.** In the ‘indirect’ mode of the SMFA, gametocytes were pre-incubated with test control compounds prior to mosquito feeding. To this end, compounds were dissolved in DMSO to achieve a stock solution of 10 mM that was serially diluted in DMSO to achieve concentrations 1000-fold above the final test concentration. Subsequently, 10 µl of diluted compound in DMSO was added to 990 µl pre-warmed RPMI 1640 medium supplemented with 367 µM hypoxanthine, 25 mM HEPES and 25 mM NaHCO3 (‘incomplete medium’). 40 µl of this intermediate dilution was added to 360 µl of parasite culture and incubated for 24 hrs at 37 °C in Eppendorf tubes, resulting in 0.1% final DMSO concentration. Hereafter, 300 µl of the gametocyte culture/compound mix was added to 180 µl of packed red blood cells and centrifuged for 20 seconds at 10,000 × g. After carefully aspirating the supernatant, 200 µl of human serum type A was added to the pellet. Finally, 300 µl of this mix was immediately injected into an individual membrane covered minifeeder, where 50 female *Anopheles stephensi* mosquitoes were allowed to feed for 10 minutes. In the ‘wash-out’ mode of the SMFA, compounds were washed out following pre-inoculation by adding 14 ml incomplete medium supplemented with 10% human type A serum (‘complete medium’) to the gametocyte suspension, followed by centrifugation (2000 × g) for 7 minutes. The supernatant was removed and the gametocytes were washed again with 14 ml of complete medium. The pellet was resuspended in 300 µl of complete medium and combined with 180 µl packed red blood cells. All media used for washing were prewarmed to 37 °C and the subsequent preparation of the blood meal was as described above. For the ‘direct’ mode of the SMFA, 300 µl parasite culture was added to 180 µl red blood cells, centrifuged for 20 seconds at 10,000 g and the supernatant was carefully aspirated. Meanwhile, 100 µl of compound diluted in incomplete medium was added to 490 µl human serum type A (0.1% DMSO final). 200 µl of this human serum/compound mix was added to the gametocyte culture/compound mix, which was subsequently used for the SMFA as described above.

For dose response analyses, compounds were diluted in DMSO with subsequent dilution in culture medium to a final DMSO concentration of 0.1%. Controls included vehicle (0.1% DMSO) and DHA at its approximate IC50 (10 µM). For all assay types, unfed and partially fed mosquitoes were removed from the cage after feeding, and fed mosquitoes were maintained at 26 °C, 80% humidity. Mosquitoes were either dissected and examined for midgut oocysts on day 6–8 PI as described,[64–67] or processed in the luciferase activity assay on day 8 PI.

**Mosquito processing and luciferase assay.** To enhance flexibility, and to ensure that individual mosquitoes were safely and efficiently transferred from feeding cages to 96-well plates for homogenisation, mosquitoes were immobilised at day 8 PI and stored at −20 °C in sealed 15 ml falcon tubes for at least 24 hours and up to 2 weeks, before being processed in the luciferase assay. Upon removal from storage at −20 °C mosquitoes were thawed and individually added to wells in shallow 96-well plates containing 60 µl PBS with Complete protease inhibitor cocktail according to the instructions of the manufacturer (Roche, Switzerland), and 0.2 gram of 1.0 mm diameter zirconia beads (Lab Services BV). Plates were next placed in a Mini-Beadbeater-96 (Biospec, Bartlesville, OK) and rocked for 2×15 seconds, with 1–2 minutes on ice between rounds of homogenisation. Based on visual inspection, this procedure achieved a homogenous consistency in line with that achievable using a hand-held pestle rotator as described previously.[13] For the individual mosquitoes 45 µl of homogenised material was transferred to wells in a Krystal 2000 black & white 96-well plate (Porvair, United Kingdom). Next 45 µl of Bright-Glo luciferase assay substrate (Promega, Madison, WI) was added to each well using a Cybi Selma dispensing station (Cybio, Germany) and the plate was incubated for 3 minutes at room temperature. Luciferase activity was measured using a Synergy 2 multi-purpose plate reader (Biotek, Winooski, VT). Background was determined by measuring uninfected mosquitoes that had undergone the same treatment.
pLDH and Pfs25 assays. Assays were performed as described previously.13

Data analysis. Statistical analysis was conducted using R (Foundation for Statistical Computing, Vienna, Austria), STATA 12 (StataCorp, TX, USA) and GraphPad Prism 5.0 (GraphPad Software Inc., CA, USA).

All comparisons of microscopy and luciferase outcomes were conducted with separate groups of mosquitoes for each measure sampled from the same feed. Comparisons between oocyst and luminescence based measures of mosquito infection prevalence were made with Chi-squared tests with Bonferroni correction. Associations between oocyst and luminescence intensity were quantified with linear regression. 95% confidence intervals around mean luminescence and oocyst intensity data was calculated by bootstrapping (1000 repetitions). Luminescence and microscopic TRA estimates were made using generalised linear mixed models (GLMMs)35,36. For assessment of infection prevalence, the cut-off for RLU based infection positivity was determined as absolutely or mean luminescence intensity of all uninfected controls plus five standard deviations (12.7 RLU), as described previously.15. For gametocyte viability and gametogenesis assays, data were expressed as the percentage effect relative to the MIN and MAX controls13. For SMFAs, data were expressed relative to the negative (vehicle) controls for oocyst density and oocyst prevalence (i.e. the proportion of infected mosquitoes). IC50 values were calculated by applying a four-parameter logistic regression model using a Maximum Likelihood Estimation algorithm to find the best fit.

References

1. World Health Organization. World Malaria Report 2013. (World Health Organization, 2013).
2. Okiro, E. et al. The decline in paediatric malaria admissions on the coast of Kenya. Malar J. 6, 151 (2007).
3. Peatey, C. L. et al. Effect of Antimalarial Drugs on Plasmodium falciparum Gametocytes. J. Infect. Dis. 200, 1518–1521 (2009).
4. Group, I. A. S. Artesunate combinations for treatment of malaria: meta-analysis. The Lancet 363, 9–17 (2004).
5. Okell, L., Drakeley, C., Ghani, A., Bousema, T. & Sutherland, C. Reduction in malaria patients by artemisinin combination therapies: a pooled analysis of six randomized trials. Malar J. 7, 125 (2008).
6. Bousema, T. et al. Revisiting the circulation time of Plasmodium falciparum gametocytes: molecular detection methods to estimate the duration of gametocyte carriage and the effect of gametocytocidal drugs. Malar J. 9, 136 (2010).
7. Abdul-Ghani, R. & Beier, J. Strategic use of antimalarial drugs that block falciparum malaria parasite transmission to mosquitoes to achieve local malaria elimination. Parasitol. Res. 113, 3335–3346 (2014).
8. White, N. J. Primaquine to prevent transmission of falciparum malaria. Lancet Infect Dis 13, 175–181 (2013).
9. Graves, P. M., Gelband, H. & Garner, P. Primaquine or other 8- aminoquinolines for reducing P. falciparum transmission. Cochrane Database Syst Rev 6, CD008152 (2014).
10. Ezekiel, A. et al. Rationale for short course primaquine in Africa to interrupt malaria transmission. Malar J. 11, 360 (2012).
11. Delves, M. J. et al. Male and Female Plasmodium falciparum Mature Gametocytes Show Different Responses to Antimalarial Drugs. Antimicrob Agents. Chemother. 57, 3268–3274 (2013).
12. Ruecker, A. et al. A Male and Female Gametocyte Functional Viability Assay To Identify Biologically Relevant Malaria Transmission-Blocking Drugs. Antimicrob Agents. Chemother. 58, 7292–7302 (2014).
13. Balsch, J. M. et al. A combination of new screening assays for prioritization of transmission-blocking antimalaria reveals distinct dynamics of marketed and experimental drugs. J. Antimicrob. Chemother. 70, 1357–1366 (2015).
14. Wang, J. et al. Anopheles gambiae Circumsporozoite Protein–Binding Protein Facilitates Plasmodium Infection of Mosquito Salivary Glands. J. Infect. Dis. 208, 1161–1169 (2013).
15. Stone, W. J. et al. A scalable assessment of Plasmodium falciparum transmission in the standard membrane-feeding assay, using transgenic parasites expressing green fluorescent protein-luciferase. J Infect Dis 210, 1456–1463 (2014).
16. Vaughan, A. M. et al. A transgenic Plasmodium falciparum NF54 strain that expresses GFP-luciferase throughout the parasite life cycle. Mol. Biochem. Parasitol. 186, 143–147 (2012).
17. Sanz, L. et al. P. falciparum in vitro killing rates allow to discriminate between different antimalarial mode-of-action. PloS one 7, e39049 (2012).
18. Limás, M., Bozdech, Z., Wong, E. D., Adai, A. T. & DeRisi, J. Comparative whole genome transcriptome analysis of three Plasmodium falciparum strains. Nucleic Acids Res. 34, 1166–1173 (2006).
19. D’Alessandro, S. et al. A Plasmodium falciparum screening assay for anti-gametocyte drugs based on parasite lactate dehydrogenase detection. J. Antimicrob. Chemother. 68, 2048–2058 (2013).
20. Tanaka, T. Q. & Williamson, K. C. A malaria gametocytocidal assay using oxidoreduction indicator, alamarBlue. Mol. Biochem. Parasitol. 177, 160–163 (2011).
21. Lelièvre, J. et al. Activity of clinically relevant antimalarial drugs on Plasmodium falciparum mature gametocytes in an ATP bioluminescence “transmission blocking” assay. PloS one 7, e35019 (2012).
22. Lucantoni, L., Duffy, S., Adjalley, S. H., Fidock, D. A. & Avery, V. M. Identification of MMV Malaria Box Inhibitors of Plasmodium falciparum Early-Stage Gametocytes Using a Luciferase-Based High-Throughput Assay. Antimicrob Agents. Chemother. 57, 6050–6062 (2013).
23. Delves, M. J. et al. A high-throughput assay for the identification of malarial transmission-blocking drugs and vaccines. Int. J. Parasitol. 42, 999–1006 (2012).
24. Bowman, J. D. et al. Antipicoplas and gametocytocidal screening to identify the mechanisms of action of compounds within the malaria box. Antimicrob Agents. Chemother. 58, 811–819 (2014).
25. Duffy, S. & Avery, V. M. Identification of inhibitors of Plasmodium falciparum gametocyte development. Malar J. 12, 408 (2013).
26. Sanders, N. G., Sullivan, D. J., Mlambo, G., Dimopoulos, G. & Tripathi, A. K. Gametocytocidal screen identifies novel chemical classes with Plasmodium falciparum transmission blocking activity. PloS one 9, e103817 (2014).
27. Dechering, K. J. et al. Isolation and functional characterization of two distinct sexual-stage-specific promoters of the human malaria parasite Plasmodium falciparum. Mol. Cell. Biol. 19, 967–978 (1999).
28. Faucon, E. et al. Identifying genomic changes associated with insecticide resistance in the dengue mosquito Aedes aegypti by deep targeted sequencing. Genome Res. 25, 1347–1359 (2015).
29. Li, T. et al. Robust, reproducible, industrialized, standard membrane feeding assay for assessing the transmission blocking activity of vaccines and drugs against Plasmodium falciparum. Malar J. 14, 150 (2015).
30. Zollner, G. E. et al. Population dynamics of sporogony for Plasmodium vivax parasites from western Thailand developing within three species of colonized Anopheles mosquitoes. Malar J. 5, 68 (2006).
31. Stone, W. J. R. et al. The relevance and applicability of oocyst prevalence as a read-out for mosquito feeding assays. Sci. Rep. 3, doi: 10.1038/srep03418 (2013).
32. Rosenberg, R. & Rungswongne, J. The number of sporozoites produced by individual malaria oocysts. Am J Trop Med Hyg 45, 574–577 (1991).
33. Yoon, Y. J. et al. KRIBB11 inhibits HSP70 synthesis through inhibition of heat shock factor 1 function by impairing the recruitment of positive transcription elongation factor b to the hsp70 promoter. J. Biol. Chem 286, 1737–1747 (2011).
Author Contributions
M.V., W.S. and K.D. wrote the main manuscript text. M.V., W.S., K.K., G.G. and B.S. performed experiments and data analyses. D.L. selected compounds and performed data analyses. R.S., T.B. and K.D. conceived the studies. All authors reviewed the manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep

Competing financial interests: KJD and RWS own stock in TroplIQ Health Sciences. All other authors have no competing financial interests to declare.

Acknowledgements
We want to thank Chris Janse and Shahid Kahn for providing us with plasmid PL1063. In addition, we thank Wouter Graumans, Jacqueline Kühnen, Laura Pelser-Posthumus, Astrid Pouwels, and Jolanda Klaassen for mosquito breeding, and for their assistance during SMFA evaluation. Hans-Peter Duerr is gratefully acknowledged for statistical analyses, and Paul Willis for selection of compounds for SMFA evaluation. This work was supported by a grant from the Bill and Melinda Gates Foundation (OPP1040394). W.S. and T.B. are supported by PATH Malaria Vaccine Initiative and the European Community's Seventh Framework Programme (SIGNAL; Marie Curie Career Integration Grant PCIG12-GA-2012-333936).

Author Contributions
M.V., W.S. and K.D. wrote the main manuscript text. M.V., W.S., K.K., G.G. and B.S. performed experiments and data analyses. D.L. selected compounds and performed data analyses. R.S., T.B. and K.D. conceived the studies. All authors reviewed the manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep

Competing financial interests: KJD and RWS own stock in TroplIQ Health Sciences. All other authors have no competing financial interests to declare.
How to cite this article: Vos, M. W. et al. A semi-automated luminescence based standard membrane feeding assay identifies novel small molecules that inhibit transmission of malaria parasites by mosquitoes. Sci. Rep. 5, 18704; doi: 10.1038/srep18704 (2015).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/