The following full text is a publisher's version.

For additional information about this publication click this link. http://hdl.handle.net/2066/199051

Please be advised that this information was generated on 2019-05-05 and may be subject to change.
A multiplex assay for the sensitive detection and quantification of male and female *Plasmodium falciparum* gametocytes

Lisette Meerstein-Kessel¹,², Chiara Andolina¹, Elvira Carrio³,⁴, Almahamoudou Mahamar⁵, Patrick Sawa⁶, Halimatou Diawara⁵, Marga van de Vegte-Bolmer¹, Will Stone⁷, Katharine A. Collins¹, Petra Schneider⁸, Alassane Dicko⁵, Chris Drakeley⁷, Ingrid Felger³,⁴, Till Voss³,⁴, Kjerstin Lanke¹ and Teun Bousema¹*

Abstract

**Background:** The transmission of malaria to mosquitoes depends on the presence of gametocytes that circulate in the peripheral blood of infected human hosts. Sensitive estimates of the densities of female gametocytes (FG) and male gametocytes (MG) may allow the prediction of infectivity to mosquitoes and thus a molecular estimate of the human infectious reservoir for transmission.

**Methods:** A novel multiplex qRT-PCR assay with intron-spanning primers was developed for the parallel quantification of FG and MG. *CCp4* (PF3D7_0903800) transcripts specific for FG and *PfMGET* (PF3D7_1469900) transcripts specific for MG were quantified in total nucleic acids. The assay was validated on sex-sorted gametocytes from culture material and on samples from clinical trials with gametocytocidal drugs. Synthetic RNA standards were generated for the two target genes and calibrated against known gametocyte quantities.

**Results:** The limit of detection was determined at 0.1 male and 0.1 female gametocyte/µL, which was equal to the limit of quantification (LOQ) for MG, while the LOQ for FG was 1 FG/µL. Results from previously reported clinical trials that used separate gametocyte qRT-PCR assays for FG (targeting *Pfs25*) and MG (targeting *PfMGET*) were reproduced with the multiplex assay. High levels of agreement between separate assays and the multiplex approach were observed ($R^2 = 0.9473$, 95% CI 0.9314–0.9632, for FG measured by transcript levels of *Pfs25* in qRT-PCR or *CCp4* in multiplex; $R^2 = 0.8869$, 95% CI 0.8541–0.9197, for MG measured by *PfMGET* in either single or multiplex qRT-PCR). FG and MG transcripts were detected in pure ring stage parasites at 10,000- and 100,000-fold reduced frequency for *CCp4* and *PfMGET*, respectively. The *CCp4* and *PfMGET* transcripts were equally stable under suboptimal storage conditions.

**Conclusions:** Gametocyte densities and their sex ratios can be determined in the presented one-step multiplex assay with higher throughput than single assays. The interpretation of low gametocyte densities at asexual parasite densities above 1000 parasites/µL requires caution to avoid false positive gametocyte signals from spurious transcript levels in ring stage parasites.

*Correspondence: teun.bousema@radboudumc.nl
¹ Department of Medical Microbiology, Radboud University Medical Center, Nijmegen, The Netherlands
Full list of author information is available at the end of the article

© The Author(s) 2018. This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.
Background
The significant reduction in malaria mortality and morbidity seen in the last 10 years is the result of combined efficient control measures such as early diagnosis, effective treatment with artemisinin-based combination therapy (ACT), active surveillance and vector control. Nevertheless, in 2016, 216 million cases were still reported worldwide [1] and major concerns exist about the extent to which the emergence and spread of insecticide [2] and artemisinin resistance [3] may affect worldwide malaria control and elimination efforts. New strategies may be needed to sustain recent gains and accelerate malaria elimination initiatives. These new strategies include the development and deployment of transmission-blocking strategies that aim to reduce malaria incidence by targeting the infection reservoir involved in maintaining parasite transmission from humans to *Anopheles* mosquitoes.

Transmission to mosquitoes is mediated by the presence of gametocytes in peripheral blood of a human host. Gametocytes are sexually dimorphic and both sexes are required to ensure the development of the parasite inside the mosquito. Although the likelihood of mosquito infection is largely dictated by gametocyte density [4], the gametocyte sex ratio may also play a significant role in ensuring fertilization [5–7]. Since one female gametocyte (FG) produces only one gamete, while a male gametocyte (MG) produces eight gametes [8, 9], the gametocyte sex ratio is usually female-biased in the proportion of 3–5 females: 1 male [10, 11] with indications from rodent malarias and natural *Plasmodium falciparum* infections that sex ratios may be adjusted in the presence of other parasite clones [6, 12], in relation to gametocyte density [7, 13] during infections and in response to environmental factors, such as anaemia [10]. Recent findings further suggest that anti-malarial drugs may have differential effects on MG and FG [14, 15]. Understanding gametocyte sex ratios is thus of interest to understand *Plasmodium* biology, better predict transmission potential during natural malaria infections and estimate the likelihood of onward transmission to mosquitoes after treatment with anti-malarial drugs.

Gametocytes usually circulate in blood at low levels as only 0.2–1% of asexual parasites commit to sexual development at every cycle of red blood cell invasion [16]. Several studies have observed infected mosquitoes after feeding on blood containing gametocyte densities as low as 0.25–0.3 gametocytes/microlitre of blood, well below the threshold for detection by routine microscopy [7, 17–21]. As a consequence of the abundant presence of submicroscopic densities of gametocytes in clinical and asymptomatic infections [22–24], they represent a silent infectious reservoir in the population. In the last 20 years, sensitive molecular techniques based on sexual stage-specific mRNA transcripts have been developed to improve the detection and quantification of both gametocyte sexes. *Pfs25* mRNA has been widely used as a mature gametocyte marker [19–21, 25] and was recently confirmed to be female-specific or at least consider-ably female-biased [26, 27]. Based on RNA-seq analysis, *PfMGET* was recently presented as a novel male-specific gametocyte marker [26, 28]. The use of intron-spanning primers allows for sensitive detection of MG in samples of naturally infected parasite carriers [28]. Thus far a combination of separate *Pfs25* and *PfMGET* qRT-PCR assays has been used to estimate gametocyte sex ratios in natural and controlled infections [7, 15, 28–30]. However, estimating sex ratios by using two separate qRT-PCR assays may affect assay precision and throughput. Here, a novel target for FG is proposed, *CCp4*, that was previously identified as a gametocyte-specific transcript [31] and allows for intron-spanning primer design. This manuscript describes a one-step multiplex qRT-PCR assay for robust assessments of gametocyte sex ratios at densities below the microscopic threshold for gameto-ocyte detection.

Methods
Selection of male and female marker transcript
The selection of the male marker *PfMGET* was described previously [28]. The female marker *CCp4* was identified by integrating transcriptomics data and validated as gametocyte-specific [31].

Preparation of gametocyte material
Sex-sorted gametocytes were generated as described previously [26, 28]. In brief, cultures of the PfDynGFP/P47mCherry line [26] were treated with N-acetyl glucosamine and stage V gametocytes were FACS sorted for their fluorescence signal (MG are sorted as GFP-positive/mCherry-negative, FG as mCherry-positive/GFP-negative) and afterwards counted with a Bürker-Türk counting chamber. For both MG and FG tenfold dilution series were prepared in whole-blood in the range of 10⁶/mL to 10⁴/mL and stored in RNAProtect to serve as standard curves for gametocyte quantification.

Preparation of ring stage parasites to assess transcript stage specificity
Asexual parasites of the NF54 strain were synchronized by the selection of late trophozoites and schizonts as described [28]. In brief, a 63% Percoll density gradient was followed by a 5% sorbitol treatment, killing the remaining schizonts after 5 h and ensuring tight synchronization. NF54 ring stage parasites were harvested 10–20 h after the Percoll treatment and stored in lysis.
buffer (5.25 M GuSCN; 50 mM Tris–HCl pH 6.4; 20 mM EDTA; 1.3% Triton X-100) for later analysis.

To obtain pure asexual stage reference material without plausible contamination by gametocytes, ring stage parasites of the gametocyte-deficient F12 clone were used that have a loss-of-function mutation in the gene encoding the gametocyte master transcription factor AP2-G [32]. In addition, ring stage parasites were generated from the recently described AP2-G knock-down line 3D7/AP2-G-GFP-DDgsmS that does not express AP2-G when grown in the presence of 2.5 mM (d)-glucosamine (GlcN) (Sigma Aldrich) [33]. Under these conditions both the ap2-g-fdp-dd transcript and AP2-G-GFP-DD protein are degraded, resulting in no gametocyte production. Parasites were released from infected RBCs by saponin lysis and total RNA was directly isolated using Ribozol (Amresco) according to the manufacturer’s manual.

Samples from naturally infected gametocyte carriers

Samples from two previously published clinical trials in light microscopy-positive gametocyte carriers from Kenya [28] and Mali [15] were used to directly compare gametocyte density estimates using the PfMGET/CCp4 multiplex assay with the previous qRT-PCR assays targeting PfMGET (male) and Pfs25 (female) transcripts separately. The multiplex assay was performed on Kenyan samples collected prior to treatment (day 0; n = 31), and after treatment on day 2 (n = 16), day 7 (n = 46), and day 14 with dihydroartemisinin–piperaquine (DP) alone or with primaquine (n = 28; total n = 121). The samples from Mali that were included in the current study were collected on day 7 after treatment with either DP (n = 15) or DP + methylene blue (15 mg/kg given daily for the first 3 days of treatment; DP + MB, n = 19) or sulfadoxine–pyrimethamine and amodiaquine (SP–AQ, n = 19) or SP–AQ with a single dose of primaquine (0.25 mg/kg given together with the first dose of SP–AQ, SP–AQ + PQ, n = 19) [15]. In both studies considerable variation in gametocyte densities and sex ratios was previously reported.

Nucleic acid extraction and target amplification

Total nucleic acids were extracted from 50 µL whole blood in five volumes RNAProtect with the MagNAPure LC automated extractor (Roche) using the MagNAPure LC Total Nucleic Acid Isolation Kit—High Performance, with the exception of F12- and 3D7/AP2-G-GFP-DDgsmS-derived material (bulk Ribozol (Amresco) extraction according to the manufacturer’s manual). Total nucleic acids were eluted in 50 µL of MagNAPure elution buffer, of which 5 µL was used in the multiplex assay. For the multiplex assay, we used the Luna® Universal Probe One-Step RT-qPCR Kit (NEB). Gene IDs of the male and female markers and respective primer and probe sequences can be found in Table 1, for additional primer sequences see Additional file 1. Probe and primer concentrations were optimized (see Additional file 1) to obtain efficient amplification of both targets. The optimal conditions are summarized in Table 2. Negative controls were run to ensure there were no unspecific signals detected from no-template controls, or from total nucleic acids without reverse transcription (intron-spanning primers do not bind to genomic DNA); and melt curves were visually inspected.

Serial dilutions of all ring stage materials were used to detect ring stage transcripts (SBP-1) and early

<table>
<thead>
<tr>
<th>Gene ID</th>
<th>Name</th>
<th>Fwd primer seq</th>
<th>Rev primer seq</th>
<th>Probe seq</th>
<th>Fluoro-phore</th>
</tr>
</thead>
<tbody>
<tr>
<td>PF3D7_1031000</td>
<td>Pf25 [36]</td>
<td>GAAATCCCGTTTCTATACGGCTTG</td>
<td>AGTTTTACACAGAGATTGCCCTTGTACAA</td>
<td>6FAM-ccgcttcatagcttgtaa-MGB</td>
<td>FAM</td>
</tr>
<tr>
<td>PF3D7_1031000</td>
<td>Pf25 [34]</td>
<td>CACATGAATATGAGAATAAAATTG*</td>
<td>TAGGGCAACATGGGAAAG</td>
<td>AGCAACACCCGTATGTGGCCTTAAACGC</td>
<td>Texas Red</td>
</tr>
<tr>
<td>PF3D7_0903800</td>
<td>CCp4 (MPX)</td>
<td>CACATGAATATGAGAATAAAATTG*</td>
<td>TAGGGCAACATGGGAAAG</td>
<td>CAGCTCCAGCATTTAAAAACAC</td>
<td>FAM</td>
</tr>
<tr>
<td>PF3D7_0903800</td>
<td>CCp4 (qRT-PCR)</td>
<td>CACATGAATATGAGAATAAAATTG*</td>
<td>TAGGGCAACATGGGAAAG</td>
<td>CAGCTCCAGCATTTAAAAACAC</td>
<td>FAM</td>
</tr>
<tr>
<td>PF3D7_1469900</td>
<td>PfMGET (MPX)</td>
<td>CGGTCAACGATTAAAACCTGC*</td>
<td>TGTGTAACGATTACCTTTC</td>
<td>CAGCTCCAGCATTTAAAAACAC</td>
<td>FAM</td>
</tr>
<tr>
<td>PF3D7_1469900</td>
<td>PfMGET (qRT-PCR, [28])</td>
<td>CGGTCAACGATTAAAACCTGC*</td>
<td>TGTGTAACGATTACCTTTC</td>
<td>CAGCTCCAGCATTTAAAAACAC</td>
<td>FAM</td>
</tr>
</tbody>
</table>

MPX multiplex assay, qRT-PCR quantitative real time reverse transcription-PCR

*Intron-spanning
Table 2 Multiplex conditions for male–female assay

<table>
<thead>
<tr>
<th></th>
<th>Concentration</th>
<th>Program</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female primers CCp4</td>
<td>900 nM</td>
<td>55 °C 15 min</td>
</tr>
<tr>
<td>Female probe—Texas Red</td>
<td>200 nM</td>
<td>95 °C 1 min</td>
</tr>
<tr>
<td>Male primers PfMGET</td>
<td>225 nM</td>
<td>95 °C 10 s</td>
</tr>
<tr>
<td>Male probe—FAM</td>
<td>200 nM</td>
<td>60 °C 1 min</td>
</tr>
<tr>
<td>Input total nucleic acid</td>
<td>5 µL</td>
<td></td>
</tr>
</tbody>
</table>

Synthetic RNA standard curve material
Linear dsDNA templates for the target regions of CCp4 and PfMGET were synthesized by BaseClear B.V. the Netherlands (for sequences, see Additional file 2) and purified by agarose gel electrophoresis. Bulk RNA was transcribed with the MEGAShortscript T7 high yield transcription kit (Invitrogen) at 10–50 nM dsDNA input according to the manufacturer’s instructions. Transcription samples were DNase I treatment (Promega). cDNA was prepared with the High Capacity cDNA Reverse Transcription Kit (Applied BioSystems) and 2 µL of cDNA was run in the GoTaq qPCR Master Mix (Promega). All primers were used at 900 nM, except for SBP1, PfMGET and PfS25 which were run at 225 nM primers.

Gametocyte transcript stability testing
Synchronized mature gametocytes (NF54, 1.8% parasitaemia) were diluted in whole EDTA-blood starting at concentrations of 10⁵ gametocytes/mL. These samples were either stabilized by adding five volumes of RNAProtect (Qiagen) or left unstabilized (no protective buffer added) for further treatments: One to three aliquots were kept at room temperature (22–25 °C) for 0 h, 1 h, 2 h, 4 h, 6 h, 8 h or 24 h before freezing them at −80 °C and subsequent processing. A subset of samples (n = 3) in RNAProtect (added at 0 h) were additionally freeze–thawed five times (37 °C/−80 °C cycling for at least 1 h each) before extraction of nucleic acids. The stability of the transcripts after extraction was also tested by freeze–thaw cycles at which the samples were left at room temperature (22–25 °C) or 37 °C for 1 h, interspersed by at least 1 h at −20 °C.

Statistical analysis
Graphs and statistical analyses were made with GraphPad Prism (version 5.0.3) or R statistical software (version 3.4.0). The concordance between separate qRT-PCR assays and the multiplex approach was assessed by estimating the slope and 95% confidence interval (95% CI) in linear regression. No statistical comparisons were made on the CT values of different marker genes since this was beyond the scope of the current manuscript and meaningful comparisons would require a larger number of replicates. Where transcript abundance differences were estimated, this was based on the assumption of doubling of transcript after each cycle (1 Ct difference), regardless of reaction efficiencies.

Results
Male and female gametocyte transcripts are detected in low densities of gametocytes
A multiplex assay was developed to target the female- and male-specific gametocyte transcripts CCp4 and PfMGET simultaneously by intron-spanning primers. Amplification from genomic DNA was absent when qRT-PCRs were performed without reverse transcription. Protocol optimization included limiting the primer concentration for the male target, which is explained in detail in Additional file 1 and resulted in an assay with two equally efficient amplification reactions for both target transcripts. Sorted MG and FG of the NF54-derived fluorescent reporter line PfDynGFP/PfP47mCherry [26] were used in dilution series to determine the limit of detection (LOD), limit of quantification (LOQ) and variation in the multiplex assay. In an octuplicate run of female and male standard curve material, the coefficient of within-run variation was assessed (Fig. 1a, b), which is very low (2–3%) for both target transcripts at high gametocyte densities, but increasing to 6% or 29% at densities of 100 sex-sorted gametocytes/mL for PfMGET and CCp4, respectively. The LOD is 100 MG or FG/mL (0.1/µL) with 96.2% or 100% of all experiments at this density (n = 26) positive for PfMGET or CCp4, respectively. Even at 10 MG/mL (0.01 MG/µL), the male signal was detected in the majority of experiments (18 of 26, 69.2%), but with an increased variation coefficient (18%). The LOQ was hence equal to the LOD for MG (10² MG/mL or 0.1 MG/µL) while for FG the LOD was 10² FG/mL (0.1 FG/µL) and the LOQ was higher (10³ FG/mL or 1 FG/µL).
Low densities of male and female gametocyte transcripts are detected in high densities of asexual parasites

Whilst previous studies reported negligible $Pfs25$ and male gametocyte transcripts in asexual (ring stage) parasites [25, 34, 35], low levels of $CCp4$, $Pfs25$ and $PfMGET$ transcripts were detected in pure asexual parasite material (Fig. 2). About 10,000 ring stage parasites from the gametocyte-deficient F12 $P. falciparum$ line [32] provided a similar quantity of $CCp4$ transcript as one FG and 100,000 F12 ring stages carried $PfMGET$ transcript equivalent to one MG. In addition, ring stage parasites also contained $Pfs25$ transcripts. Their concentration was about 100,000-fold less than FG (Additional file 1: Figure S2A). Interestingly, compared to F12 ring stages the expression of these gametocyte markers in ring stages of the 3D7/AP2-G-GFP-DDglmS line [33] was slightly lower when AP2-G was not expressed. Under these conditions, the fold-change in transcript between asexual rings and FG increased to 50,000-fold for $CCp4$ and to 200,000- to 300,000-fold for $PfMGET$ (MG over asexual rings). Ring stage material of the F12 line and 3D7/AP2-G-GFP-DDglmS was then compared with the gametocyte-producing NF54 line for the expression of markers that are transcribed from or at onset of sexual differentiation ($Pfg27$, $Pfs16$, $Pfg14-744$ and $Pfg14-748$). The results indicate low-level contamination with early gametocytes in NF54 compared to both gametocyte-non producing lines. Consistently, all
markers were expressed at higher levels in NF54 ring stages compared to F12 and 3D7/AP2-G-GFP-DDglmS (Additional file 1: Figure S3).

The measured background expression of PfMGET and CCp4 transcripts in asexual ring stages leads to the following precautions, based on a conservative estimate of fold change of transcript levels (compared to F12 rings): for the detection of MG and FG, the presence of asexual parasites at densities above 1000 parasites/µL can result in a false-positive signal for FG whilst asexual parasites at densities above 10,000 parasites/µL can also result in a false-positive signal for MG (Table 3).

Both CCp4 and PfMGET were also moderately expressed in the opposite-sex gametocyte, detectable at concentrations of 10,000/mL or higher (Fig. 3). The opposite sex (MG) accounted for 0.1% (1 in 1000) false-positive FG with CCp4 (Fig. 3a), and about every 100 FG accounted for 1 MG (1% false positives) in PfMGET-based detection (Fig. 3b). Pfs25 shows higher background than CCp4 in the opposite sex: 0.5% (1 in 200, Additional file 1: Figure S2B).

The male/female gametocyte multiplex assay corroborates earlier density estimates in naturally infected individuals. Estimates of MG and FG are highly correlated with the estimates obtained by Pfs25 qRT-PCR (FG for Kenya and Mali combined, R² = 0.9473, p < 0.001) and PfMGET qRT-PCR (MG combined, R² = 0.8869, p < 0.001) (Fig. 4a, b) using a selection of samples from clinical trials in Kenya (n = 84 samples) [28] and Mali (n = 72 samples) [15]. In the Malian study [15], extreme gametocyte sex ratios were reported and multiplex qRT-PCR-estimates for sex-specific gametocyte densities (Fig. 4a, b), prevalence (Fig. 4c) and sex ratios (Fig. 4d) were in agreement with the conventional quantification by qRT-PCR targeting Pfs25 for FG [36] or PfMGET transcripts for MG [28].

A linear regression curve was fitted for single versus multiplex assay density estimates. For the MG quantification, there were indications that the slope deviated

### Table 3 The consequences of low level gametocyte transcripts in asexual parasites and gametocytes of the opposite sex for assessing gametocyte prevalence and density

<table>
<thead>
<tr>
<th>Asexual parasite density</th>
<th>False positive signal in gametocyte quantification</th>
<th>Trust prevalence/estimated density if density is significantly above</th>
</tr>
</thead>
<tbody>
<tr>
<td>Log10/mL</td>
<td>FG</td>
<td>MG</td>
</tr>
<tr>
<td>4</td>
<td>10/µL</td>
<td>0.001/µL</td>
</tr>
<tr>
<td>5</td>
<td>100/µL</td>
<td>0.01/µL</td>
</tr>
<tr>
<td>6</td>
<td>1000/µL</td>
<td>0.1/µL</td>
</tr>
<tr>
<td>7</td>
<td>10,000/µL</td>
<td>1/µL</td>
</tr>
<tr>
<td>8</td>
<td>100,000/µL</td>
<td>10/µL</td>
</tr>
</tbody>
</table>

Numbers in italics are above the limit of detection. Other estimates are presented to illustrate the marginal impact of signal derived from asexual parasites on gametocyte quantification. The false positive signal strength for female (FG, CCp4) and male target (MG, PfMGET) is based on the comparison of NF54 (sorted) gametocytes and F12 ring stages.
from 1 (95% CI 0.7744–0.9507 for the study in Kenya and 0.8633–0.9800 for the study in Mali) but resulting estimates in sex ratio were highly similar to those obtained after separate qRT-PCR (Fig. 4d).

**Male and female targets show similar stability under sub-optimal sample storage conditions**

Differences in transcript stability upon delays in sample processing or following freeze–thaws may affect gametocyte quantification of field samples. To resemble freshly taken blood samples with delays in sample processing, whole blood samples with NF54 parasites containing a mix of male and female gametocytes were stored at room temperature (22–25 °C) without an RNA-protecting agent. Transcript abundance of *CCp4*, *PfMGET* and *Pfs25* was determined at baseline (0 h) and after 1 h, 2 h, 4 h, 6 h, 8 h or 24 h. All targets had stable Ct values after 1 h without RNA-protecting agent (n = 1). *PfMGET* and *CCp4* showed delayed Ct values after 2 h without protection (n = 2, Additional file 1: Figure S4). To study the effect of freezing/thawing after storage, blood was stored in RNAProtect and subjected to five freeze–thaw cycles at 37 °C prior to nucleic acid extraction. A minimal transcript loss was observed (0.523 and 0.245 Ct difference to baseline for *PfMGET* and *CCp4*, respectively, Additional file 1: Table S1). Extracted total nucleic acids were more stable and showed no loss in signal (0.08 and 0.02 Ct reduction) compared to baseline for *PfMGET* and *CCp4*, after five freeze–thaw cycles at room temperature for the sorted trend line material (Additional file 1: Table S1).

**Synthetic RNA standards can be used to estimate gametocyte densities and copy numbers per gametocyte**

In vitro-synthesized RNA of the target regions of *CCp4* and *PfMGET* was used in serial dilutions alongside sorted gametocyte standards (see Additional file 1: Figure SSA, B). The synthetic RNA standards were used to estimate the copy numbers in the gametocyte reference material (see Additional file 1: Figure SSC–E). For *CCp4*, estimated transcript copies were 4 (95% CI 3.2–5.1) mRNA copies per female gametocyte and for *PfMGET* 9.8 (95% CI 8.9–10.2) copies per male gametocyte. For
comparison, female gametocytes had an average of 231.7 (95% CI 199.1–269.8) Pfs25 copies per gametocyte.

**Discussion**

In the current study, a multiplex assay for the rapid quantification of female and male gametocytes is reported. The assay utilizes a new female gametocyte marker CCp4 in conjunction with the reported male gametocyte marker PfMGET [28]. The use of intron-spanning primers allows simultaneous quantification of male and female-specific transcript levels in total nucleic acids without prior DNase I treatment. The presented analysis concludes low but non-negligible gametocyte transcripts in gametocytes of the opposite sex and gametocyte-free ring-stage asexual parasites. The stability of CCp4 and PfMGET transcripts was similar under suboptimal storage conditions; gametocytes can be reliably detected and quantified at densities 0.1–1 gametocyte/μL.

CCp4 is a member of the LCCL-domain containing adhesion protein family and orthologous to LAP6 in *Plasmodium berghei*, where this gene is reported to be translationally repressed with protein expression occurring at the ookinete stage only [37]. Earlier, a DOZI (development of zygote inhibited) knock out indicated that LAP6 transcripts (then called PB000955.03.0) are accumulated in an mRNA storage complex [38]. In *P. falciparum*, the CCp4 protein is predominantly expressed at the gametocyte stage, with only minor expression in male gametocytes [39] and no evidence for translational repression. Gametocyte-specific CCp4 transcripts were reported in an integrated analysis of eight *Plasmodium* transcriptomes [31]. The initial validation in qRT-PCR experiments confirmed CCp4 expression to be at least 1000-fold upregulated in gametocytes of different *P. falciparum* strains compared to asexual blood stages. The 1000-fold higher mRNA levels in FG over MG reported here confirm and exceed the previous estimates by RNAseq (38-fold higher in females by RPKM values) [26].

Unlike the commonly used female marker Pfs25, CCp4 allows the design of intron-spanning primers. The current assay utilizes intron-spanning primers of both male and female reporter genes with two marker-specific probes. Importantly, the multiplex gametocyte assay can be performed on total nucleic acids without DNase I treatment, which may affect gametocyte detection at low densities [28, 36]. The multiplex male–female assay is thus faster than separate assays. Sensitivity for detecting female gametocyte was lower than for the single qRT-PCR targeting Pfs25, which is at least in part explained by a lower estimated number of CCp4 transcripts per female gametocyte as compared to Pfs25. The LOD for female and male gametocytes in the multiplex assay is 0.1/μL, well below the limit of microscopic detection and in the same range as other molecular sex-specific assays [28, 34, 35]. More sensitive total gametocyte assays have been reported [40] but the current multiplex LOD allows for the detection of infections that are likely to be transmissible to mosquitoes. An increasing likelihood of mosquito infections is consistently observed at gametocyte densities above 1–5 gametocyte per microlitre [7, 29, 41]. The current assay reliably quantifies gametocytes at these densities. The lower sensitivity to detect female gametocytes as compared to Pfs25 may be a concern in studies where very low overall gametocyte densities are observed [29, 30] but the operational attractiveness of a multiplex assay that does not require DNase treatment is considerable for many other studies.

Previous work indicated that the stability of transcripts is a relevant concern when estimating gametocyte prevalence or density [36, 41]. When assessing gametocyte sex ratio, transcript stability is a particular concern since differences in the stability between target transcripts may affect bias estimates. In a limited set of experiments there were no indications for differences in the stability of PfMGET and CCp4 transcripts, provided samples are transferred to RNA-protective buffer within 1–2 h of blood collection. Freeze–thaw cycles resulted in limited RNA loss once blood samples are in this protective buffer. A similar stability of male or female signal is of particular relevance for studies conducted in low-resource settings where there may be challenges in ensuring optimal storage conditions. Repeated or prolonged freeze–thaw cycles may thus affect overall gametocyte detection or quantification [41, 42] but current results indicate they would not disproportionally affect MG or FG quantification and thus sex-ratio estimates.

The presented multiplex assay is a fast route to accurate *P. falciparum* sex ratio determination, saving about 25% of the time—with similar material costs—compared to two separate assays of which one requires a DNase treatment step. Medium sample throughput in 96-well format is the recommended application, providing accurate gametocyte quantification and sex ratio determination for blood and culture samples.

Previous studies concluded no or negligible Pfs25 transcript numbers in asexual parasites. In the current set of experiments, we detected gametocyte transcripts in different preparations of ring-stage asexual parasite material. Whilst low-level contamination of gametocytes in supposedly pure asexual parasites may have contributed to the detection of CCp4, Pfs25 and PfMGET transcripts in asexual parasite material from the NF54 strain, the detection of these transcripts in the gametocyte-deficient F12 line and under knock-down conditions for AP2-G in a more recent gametocyte-less line 3D7/AP2-G-GFP-DDglmS [33] provides convincing evidence for low-level
expression of \( \textit{Pfs25} \), \( \textit{CCp4} \) and \( \textit{PfMGET} \) in asexual ring stage parasites. The current findings of detectable transcript expression for all gametocyte markers in high densities of asexual blood stages despite different strategies to avoid gametocyte contamination have implications for past and future gametocytaemia estimates. Whilst the difference in transcript abundance between gametocytes and asexual parasites is sufficiently pronounced to conclude a marginal impact on gametocyte density estimates, gametocyte prevalence estimates may be inflated in some populations. Given the high parasitaemia of some acute malaria infections (with densities typically above 10,000 parasites/µL [43] as opposed to asymptomatic infections where densities commonly lie below 10 parasites/µL [44]), earlier studies recruiting clinical malaria cases may have overestimated gametocyte prevalence by molecular assays. In studies with asymptomatic parasite carriers and in low-endemic settings, where lower asexual parasite densities dominate, this overestimation will be less pronounced and often negligible. A previously reported rapid decline in gametocyte prevalence based on \( P/25 \) mRNA detection in the first 3 days following treatment of high-density asexual infections [45] may thus be (partially) explained by the detection of \( Pfs25 \) transcripts arising from asexual parasites, whilst the gradual decline in gametocyte prevalence following treatment of lower-density asexual parasite carriers [15, 46] or gametocyte transcript kinetics in the period following asexual parasite clearance may better reflect gametocyte clearance and gametocyte half-life [47]. With a better appreciation of caveats in gametocyte detection, the molecular tools for gametocyte detection are of value for studies aiming to quantify the human infectious reservoir for malaria, the kinetics of gametocyte production and the impact of interventions on gametocyte carriage. As a consequence of the detection of low level transcripts of gametocyte markers in rings, it is advised to report gametocyte prevalence in samples with parasite densities > 1000 parasites/µL together with a qualifying remark on the reliability of gametocyte prevalence and quantification. The presented results suggest that gametocyte prevalence determined in samples below assay-specific cut-off values indeed can be trusted (Table 3). Stating the limitations of molecularly determined gametocyte prevalence for densities, if required, will re-confirm the validity of molecular gametocyte detection.

With a cautious interpretation of low gametocyte density estimates in samples with high concurrent asexual parasite densities, molecular gametocyte diagnostics such as the multiplex assay presented in this manuscript are valuable tools to obtain sensitive and robust estimates of gametocyte prevalence and density. With these tools, gametocyte densities and sex ratios can be assessed across the gametocyte density range that is likely to contribute to onward transmission to mosquitoes [7, 19], which in many settings is well below the threshold for detection by microscopy.

**Conclusion**

The presented multiplex qPCR assay is a valuable addition to gametocyte diagnostic tools. A new female gametocyte marker gene, \( \textit{CCp4} \) was introduced and benchmarked against \( \textit{Pfs25} \) transcript-based quantification. The use of \( \textit{CCp4} \) and \( \textit{PfMGET} \) as targets has the following advantages: throughput is facilitated by the use of intron-spanning primers which allow amplification of mRNA only without a DNA digestion step, sensitivity is sufficiently high to detect and quantify all potentially transmitting gametocyte densities. The target mRNAs result in a similar detectability of male and female gametocytes and show similar stability under suboptimal storage conditions, allowing robust gametocyte sex-ratio estimates in field studies.

### Additional files

- **Additional file 1.** Optimization of the multiplex amplification assay and different gametocyte targets.
- **Additional file 2.** Protocol for multiplex amplification assay.

### Authors’ contributions

Work was conceptualized and overseen by TB, KL, LMK, KC and TB planned and conducted experiments; additional samples were prepared and provided by EC, AM, PSA, HD, MVB. CA and LMK wrote the initial manuscript, editing and conducted experiments, additional samples were prepared and provided by EC, AM, PSA, HD, MVB. CA and LMK wrote the initial manuscript, editing was done by TB, KC, WS, CD, AD, TV, EC, IF, PSc and KL. All authors read and approved the final manuscript.

### Author details

1. Department of Medical Microbiology, Radboud University Medical Center, Nijmegen, The Netherlands. 2. Centre for Molecular and Biomolecular Informatics, Radboud Institute for Molecular Life Sciences, Radboud University Medical Center, Nijmegen, The Netherlands. 3. Swiss Tropical and Public Health Institute, Basel, Switzerland. 4. University of Basel, Basel, Switzerland. 5. Malaria Research and Training Centre, University of Science, Techniques and Technology, Mbita Point, Kenya. 6. Department of Immunology and Infection, Faculty of Infectious and Tropical Diseases, London School of Hygiene and Tropical Medicine, London, UK. 7. Institute of Evolutionary Biology and Institute of Immunology and Infection Research, School of Biological Sciences, University of Edinburgh, Edinburgh, UK.

### Acknowledgements

We are thankful to Roly Gosling and Michelle Roh from UCSF (California, USA) for their agreement to use samples from the clinical trial in Mali. We further thank Pietro Alano for his contributions to the manuscript, Wouter Graumans and Rianne Stoter for their support in parasite culture.

### Competing interests

The authors declare that they have no competing interests.