Activity of *Plasmodium vivax* promoter elements in *Plasmodium knowlesi*, and a centromere-containing plasmid that expresses NanoLuc throughout the parasite life cycle

Roberto R. Moraes Barros¹,²*, Kittisak Thawnashom³,⁴, Tyler J. Gibson¹, Jennifer S. Armistead¹,⁵, Ramoncito L. Caleon¹, Miho Kaneko³,⁶, Whitney A. Kite¹, J. Patrick Mershon¹, Jacqueline K. Brockhurst¹, Theresa Engels², Lynn Lambert³, Sachy Orr-Gonzalez⁵, John H. Adams⁵, Juliana M. Sá¹, Osamu Kaneko³ and Thomas E. Wellems¹

Abstract

**Background:** *Plasmodium knowlesi* is now the major cause of human malaria in Malaysia, complicating malaria control efforts that must attend to the elimination of multiple *Plasmodium* species. Recent advances in the cultivation of *P. knowlesi* erythrocytic-stage parasites in vitro, transformation with exogenous DNA, and infection of mosquitoes with gametocytes from culture have opened up studies of this pathogen without the need for resource-intensive and costly non-human primate (NHP) models. For further understanding and development of methods for parasite transformation in malaria research, this study examined the activity of various trans-species transcriptional control sequences and the influence of *Plasmodium vivax* centromeric (*pvcen*) repeats in plasmid-transfected *P. knowlesi* parasites.

**Methods:** In vitro cultivated *P. knowlesi* parasites were transfected with plasmid constructs that incorporated *Plasmodium vivax* or *P. falciparum* 5′ UTRs driving the expression of bioluminescence markers (firefly luciferase or NanoLuc). Promoter activities were assessed by bioluminescence, and parasites transformed with human resistant allele dihydrofolate reductase-expressing plasmids were selected using antifolates. The stability of transformants carrying *pvcen*-stabilized episomes was assessed by bioluminescence over a complete parasite life cycle through a rhesus macaque monkey, mosquitoes, and a second rhesus monkey.

**Results:** Luciferase expression assessments show that certain *P. vivax* promoter regions, not functional in the more evolutionarily-distant *P. falciparum*, can drive transgene expression in *P. knowlesi*. Further, *pvcen* repeats may improve the stability of episomal plasmids in *P. knowlesi* and support detection of NanoLuc-expressing elements over the full parasite life cycle from rhesus macaque monkeys to *Anopheles dirus* mosquitoes and back again to monkeys. In assays of drug responses to chloroquine, G418 and WR9910, anti-malarial half-inhibitory concentration (**IC**₅₀) values of...
blood stages measured by NanoLuc activity proved comparable to IC_{50} values measured by the standard SYBR Green method.

**Conclusion:** All three *P. vivax* promoters tested in this study functioned in *P. knowlesi*, whereas two of the three were inactive in *P. falciparum*. NanoLuc-expressing, centromere-stabilized plasmids may support high-throughput screenings of *P. knowlesi* for new anti-malarial agents, including compounds that can block the development of mosquito- and/or liver-stage parasites.

**Keywords:** Heterologous transfection, Transgenic parasites, Genetic transformation, Luciferase expression, In vitro growth assays, Antimalarial drug response assays

**Background**

*Plasmodium knowlesi*, once thought to infect macaques almost exclusively, is now a major cause of human malaria in Malaysia [1]. Up to 10% of *P. knowlesi* human infections cause severe malaria, leading to death in 1–2% of those afflicted [2]. Increased prevalence of *P. knowlesi* infections represents an additional challenge to malaria elimination, especially in Southeast Asia, where *Plasmodium falciparum* and *Plasmodium vivax* infections are in decline [3, 4]. In some regions endemic to multiple *Plasmodium* species, malaria control work was found to decrease the prevalence of *P. falciparum*, but did not produce comparable decreases of *P. vivax* or *P. knowlesi* [3–5].

In the 1970s, laboratory research on *P. falciparum* was boosted by the development of in vitro cultivation methods [6, 7]. However, similar long-term cultivation of any other human malaria species was unavailable until the adaptation of *P. knowlesi* parasites to culture [8, 9]. As with *P. falciparum*, transfection methods and the ability to infect mosquitoes with cultivated *P. knowlesi* parasites followed [8–11], and these are supporting new experimental studies without the need for resource-intensive and costly non-human primate (NHP) models. Fundamental biological differences between the *Plasmodium* species and factors affecting malaria control that are now more accessible to investigation include: the period of the intraerythrocytic life cycle (24 h for *P. knowlesi* vs. 48 h for *P. falciparum*) [12, 13]; features of host cell preference and cell invasion mechanisms [14–16]; gametocyte development and transmission to mosquitoes [17]; and anti-malarial drug responses and mechanisms of resistance [18, 19].

Transfection and genetic transformation of malaria parasites with various bioluminescent proteins have been used for in vivo studies of the traversal and tissue invasion of sporozoites injected in mammalian hosts by mosquitoes [20]; functional analyses of promoters and introns [21–25]; screens for candidate anti-malarials and inhibitors of parasite growth [26–28]; and evaluations of compounds that block parasite infectivity to mosquitoes [29]. This report presents investigations of exogenous transcriptional control sequences in transfected *P. knowlesi* parasites. Results show that *P. knowlesi* parasites can recognize *P. vivax* promoters that are poorly recognized by *P. falciparum* parasites; further, *P. vivax* centromeric (pvcen) repeats can improve the stability of transgenic plasmids across the complete parasite life cycle of *P. knowlesi* in rhesus monkeys and *Anopheles dirus* mosquitoes.

**Methods**

**Parasites and culture**

*Plasmodium falciparum* parasites were maintained in human red blood cells (hRBCs) purchased from Interstate Blood Bank (Memphis, TN, USA). *Plasmodium knowlesi* parasites were cultivated in rhesus red blood cells (rRBCs) as described [10]; blood samples were obtained according to the National Institutes of Health (NIH) Guidelines for Animal Care and Use, under an Animal Study Proposal (ASP) approved by the National Institute of Allergy and Infectious Diseases (NIAID) Animal Care and Use Committee (ACUC); rhesus blood was obtained according to the National Institutes of Health (NIH) Guidelines for Animal Care and Use, under an Animal Study Proposal (ASP) approved by the National Institute of Allergy and Infectious Diseases (NIAID) Animal Care and Use Committee (ACUC); rhesus blood was processed and plasma was removed as described [10].

Cryopreserved stocks of a *P. falciparum* 3D7 clone [30] and the *P. knowlesi* H strain [31] were obtained from NIH inventory and thawed by standard methods [32]. *Plasmodium falciparum* cultures were propagated in *P. falciparum* complete RPMI (PF-cRPMI) consisting of RPMI-1640 (KD Medical, Columbia, MD, USA) supplemented with 25 mM HEPES, 50 μg/mL hypoxanthine, 0.21% sodium bicarbonate, 20 mg/L gentamicin, and 1% Albumax II (Life Technologies, Carlsbad, CA, USA). *Plasmodium knowlesi* cultures were maintained in *P. knowlesi* complete RPMI (PK-cRPMI), consisting of RPMI-1640 supplemented with 25 mM HEPES, 50 μg/mL hypoxanthine, 0.26% sodium bicarbonate, 10 mg/L gentamicin (KD Medical), and 1% Albumax II [32]. Cultures were maintained with RBCs up to 5% haematocrit, under a 90% N_{2}/5% CO_{2}/5% O_{2} gas mixture at 37 °C with daily media changes. Parasite development and propagation across the asexual blood-stage cycle were monitored by microscopy of thin blood films after fixation in methanol and staining for 15 min with 20% Giemsa’s solution.
Parasitaemia was estimated by counting the number of parasitized RBCs (pRBCs) per 1,000 RBCs. Cultures were maintained at parasitaemias between 0.5 and 10%.

**Rhesus infection using blood-stage *P. knowlesi***

Rhesus macaques (*Macaca mulatta*) were obtained from NIH-approved sources and housed in compliance with the Animal Welfare Act and the Guide for the Care and Use of Laboratory Animals (ILAR, 1996). All animal care and use in this study were performed in accordance with the NIH Animal Research Advisory Committee (NIH ARAC) Guidelines, under ASP protocols approved by the NIAID ACUC.

Animals were anesthetized intramuscularly (IM) with 10 mg/kg ketamine to allow infection by mosquito bites or by intravenous (IV) inoculation of cultivated blood stage parasites. For the IV inoculation, a culture sample containing ~2 × 10⁸ parasitized RBCs was pelleted by centrifugation (700 × g, 3 min). Pelleted cells were washed with 10 mL sterile and pure RPMI-1640, and resuspended to 1.5 mL volume of the same medium. After infection, parasitaemia was checked daily by Giemsa-stained thin blood films as described above.

*Plasmodium knowlesi* infections were cured with three daily administrations of oral chloroquine (total dose: 50 mg/kg body weight). Thin and thick blood films were checked up to 30 days after treatment to confirm cure.

**Mosquito feedings and assessment of oocysts and sporozoites**

Mosquito feedings were performed using 3–5-day-old female *Anopheles dirus* B and *An. dirus* (X strain) mosquitoes [31] raised in the LMVR/NIAID insectary. Mosquitoes were starved for 24 h in secure pint containers with up to 100 female mosquitoes per container. Feedings were performed at night (12:00 AM), as recommended for *P. knowlesi* transmission [12]. Two pints of mosquitoes were used in each experiment, and mosquitoes were allowed to feed through a double safety net (bridal veil) for ~20 min directly onto the inner arm or thighs of the anesthetized rhesus macaque. Fed mosquitoes were maintained with 10% corn syrup in water at 26 °C for up to 18 days in the LMVR human malaria-secure insectary.

Mosquitoes were randomly selected and dissected 6–9 days post-feeding and examined for oocysts in the midgut; additional mosquitoes were randomly selected and dissected after 10 days to examine for sporozoites in the salivary glands. To assess oocyst presence and number, midguts from 16–20 dissected mosquitoes per container were stained with 0.05% mercurochrome for 10 min and then examined under the microscope (20×–40× objective lens). Individual mosquito midguts containing oocysts were transferred to 1.5 mL tubes and used for NanoLuc assays. Sporozoites isolated from dissected salivary glands were combined and counted using a haemocytometer under the microscope at 40× magnification. The sporozoites were transferred to 1.5 mL tubes and used for NanoLuc assays. Remaining infected mosquitoes were used 14 days post-feeding to transmit parasites to an uninfected rhesus macaque as described above.

**Plasmids**

Plasmid pD-pFcam-Luc [10] contains the fLuc (firefly luciferase) cassette under control of a 0.6 kb 5′ untranslated region (UTR) from *P. falciparum* calmodulin (*pfcam*) and a 0.8 kb 3′ UTR sequence from *P. falciparum* heat shock protein 86 (*pfhsp86*). pD-pFcam-Luc also contains the hdlfr cassette under control of a 0.6 kb 5′ UTR from *Plasmodium chabaudi* dhfr-ts (*pcdts*) and a 0.8 kb 3′ UTR from *P. falciparum* heat shock protein 70 (*pfhsp70*) (Fig. 1).

*Plasmodium vivax* and *P. knowlesi* 5′ UTR sequences were amplified from genomic DNA of *P. vivax* NIH-1993 [33] or *P. knowlesi* H strains using the primers listed in Additional file 1: Table S1. Amplified sequences of 0.9 kb from the *P. vivax* calmodulin (*pvcam*) 5′ UTR, 0.9 kb of the *P. vivax* chloroquine resistance transporter (*pfvhr)* 5′ UTR, 1.4 kb of the *P. vivax* heat shock protein 70 (*pfhsp70*) 5′ UTR, and 0.6 kb of the *P. knowlesi* elongation factor-1 alpha (*pkef1-alpha*) 5′ UTR were isolated from agarose gels and cloned into the pGEM-T vector (Promega, Madison, WI, USA). The 5′ UTR sequences selected were used before and reported as capable to drive transcription in *P. vivax* (*pvcam*) [34], *Plasmodium yoelii* (*pfvhr*) [35], or *P. knowlesi* (*pfcam* and *pkef1-alpha*) [9, 10]. The *pfvhr* 5′ UTR sequence was selected by size similarity with the *pvcam* 5′ UTR sequence (0.9 Kb). To replace the *pfcam* 5′ UTR sequence that drives luciferase transcription in the pD-pFcam-Luc, the pGEM-T clones of the *pvcam*, *pfvhr*, and *pfhsp70* 5′ UTRs were digested using BplI and SpeI restriction enzymes (New England Biolabs, Ipswich, MA), and agarose gel-purified fragments were subcloned into the plasmid pD-pFcam-Luc also digested with these enzymes. Resultant plasmids were named pD-pVcam-Luc, pD-pVhr-Luc and pD-pVhsp70-Luc. To replace the *P. chabaudi* dhfr-ts 5′ UTR that drives the hdlfr transcription in the pD-PFcam-Luc, the pGEM-T clones of the *pfhsp70* and *pkef1-alpha* 5′ UTRs were digested using BplI and NcoI, and the resulting fragments were cloned into the plasmid previously digested with the same enzymes, thus generating *pfhsp70D-pFcam-Luc* and *pkef1D-pFcam-Luc*.

Plasmid pvcen-pVhsp70-D-NanoLuc was developed from pDST-PvCEN1133, pENT41-PvHSP70-5U-1 k,
pENT12-hDHFR, and pENT23-NLuc by MultiSite Gateway LR reaction (Invitrogen, Carlsbad, CA, USA) [35]. pENT23-NLuc was generated by Gateway BP reaction between pDONR P2R-P3 plasmid (Invitrogen) and a DNA fragment encoding NanoLuc Luciferase amplified from pNL1.1 (NanoLuc) vector (Promega) with primers Nluc.B2F and Nluc.B3R (Additional file 1: Table S1). The plasmid contains the centromeric sequences of *P. vivax* chromosome 11 and harbours a 1 kb 5′ UTR sequence from the *P. vivax heat shock protein* driving the expression of hdhfr. The plasmid also contains 800 bp of the 3′ UTR sequence from the *Plasmodium berghei* bifunctional dihydrofolate reductase-thymidylate synthase gene.

**Parasite transformation**

Parasites were transformed by the spontaneous DNA uptake method as described [10, 36]. Briefly, 300 µL of pelleted rhesus RBCs were washed twice by centrifugation and resuspended in 5 mL cytomic (120 mM KCl, 0.15 mM CaCl, 2 mM EGTA, 5 mM MgCl, 10 mM K_2HPO_4/KH_2PO_4, 25 mM Hepes, pH adjusted to 7.6 with KOH). The RBCs were recovered and mixed with 100 µL of cytomic containing 40 µg of plasmid DNA, and transferred to a 0.2 cm electroporation cuvette (Bio-Rad, Hercules, CA, USA). Uninfected RBCs were electroporated in a Bio-Rad Gene Pulser II at 310 V and 975 µF, with resulting time constants of 20–35 ms. The plasmid-loaded RBCs were washed twice with 5 mL of incomplete RPMI (cRPMI without Albumax II) and used immediately. Plasmid-loaded RBCs were added to 10 mL of PK-cRPMI in a 25 cm² flask. A 50 µL suspension of 3% asynchronous pRBCs at 5% haematocrit was added to the flask for a final parasitaemia of 0.5%. After 24 h in culture, fresh plasmid-loaded RBCs were again added to the flask for *P. knowlesi* transfections. After

![Fig. 1 Plasmid constructs used to evaluate Plasmodium promoter sequences in *P. knowlesi*. Expression of hdhfr from the plasmids confers resistance to WR99210 and a fluc or NanoLuc reporter. The pD-pfcam-Luc plasmid has the fluc cassette driven by the *P. falciparum* calmodulin 5′ UTR (pfcam 5′) and the hdhfr cassette driven by the *P. chabaudi* dts 5′ UTR (pcdts 5′). pD-pvcam-Luc, pD-pvcrt-Luc and pD-pvhsp70-Luc have the 5′ UTR sequences of *P. vivax* calmodulin (pvcam 5′), chloroquine resistance transporter (pvcrt 5′) and heat shock protein 70 (pvhsp70 5′) driving fluc expression, respectively, replacing the pfcam promoter from the pD-pfcam-Luc. Plasmids pvhsp70D-pfcam-Luc and pkef1D-pfcam-Luc have the 5′ UTR sequences of *P. vivax* heat shock protein 70 (pvhsp70 5′) and *P. knowlesi* elongation factor 1 α (pkef1-α 5′), respectively, replacing the pcdts promoter from the pD-pfcam-Luc to drive the expression of hdhfr. The pvcen-pvhsp70D-NanoLuc plasmid includes *P. vivax* centromeric sequence repeats from chromosome 11 (Pvcen repeat) and has hdhfr-NanoLuc fusion expression driven by the *P. vivax* heat shock protein 70 5′ UTR (pvhsp70 5′). The arrows indicate directions of transcription.
incubated for 10 min at room temperature. The result -

Luciferase assays

The fLuc and NanoLuc enzymatic assays were performed using blood- and mosquito-stage parasites. To remove haemoglobin from RBCs and avoid a possible quenching effect of luciferase assay signal, RBCs were treated with saponin: up to 250 µL of packed RBCs were pelleted from cultures by centrifugation (800 × g, 3 min), overlying supernatant culture medium was removed, and the cells were twice suspended and re-pelleted in 1,000 µL of 0.15% saponin in PBS (10 mM PO₄³⁻, 137 mM NaCl, 2.7 mM KCl). After washing twice with 1 mL of PBS, the parasites were pelleted, resuspended in 50 µL (fLuc assays) or 100 µL (NanoLuc assays) of 1× Cell Culture Lysis Reagent (Luciferase Assay System; Promega) and incubated for 10 min at room temperature. The resulting lysate was centrifuged for 1 min at 10,600 × g and the supernatant (20 µL upper liquid layer) was used for the enzymatic reaction.

Midguts containing P. knowlesi oocysts were isolated from Anopheles mosquitoes. Each isolated midgut was re-suspended in 20 µL of PBS, mixed with 80 µL of Passive Lysis Buffer (Promega) and incubated for 10 min at room temperature for cell lysis. The lysate was centrifuged for 1 min at 10,600 × g and the supernatant (upper liquid layer) was used for the NanoLuc assays. Sporozoites isolated from 8 salivary gland pairs were combined and re-suspended in 40 µL of PBS. The sporozoites were mixed with 60 µL of Passive Lysis buffer and incubated for 10 min at room temperature for cell lysis. After centrifugation (1 min at 10,600 × g) the supernatant was diluted tenfold in Passive Lysis Buffer and the dilution was utilized for NanoLuc assays.

Firefly luciferase assays were performed using the Luciferase Assay System (Promega) following manufacturer’s instructions. For this assay, 20 µL of the lysate supernatant was mixed with 100 µL of Promega Luciferase Assay Reagent and luminescence was assessed with a GloMax 20/20 Luminometer (10 s integration time). NanoLuc assays were performed using the Nano-Glo Luciferase Assay System (Promega) following manufacturer’s instructions. For the enzymatic reaction, 40 µL of the lysate supernatant were mixed with the same volume of Nano-Glo reaction solution (Promega), incubated for 3 min at room temperature, and assessed with a GloMax 20/20 Luminometer (1 s integration time). Measurements were collected in technical duplicates and averaged for analysis.

Quantitative real-time PCR

Quantitative real-time PCR (qPCR) reactions to evaluate plasmid copy number were performed using the Rotor-Gene SYBR Green PCR Kit according to manufacturer’s instructions (Qiagen, Hilden, Germany). For each reaction a total of 50 ng of genomic DNA and 200 nM of each oligonucleotide primer were mixed with the Rotor-Gene PCR SYBR Green mix and the reaction was performed in the Rotor-Gene Q instrument, following the PCR cycle: 10 s at 96 °C, 30 s at 60 °C, repeated 35 times. Primers NanoLuc RT-F-2 and NanoLucRT-R-2 were designed to amplify a 133 bp sequence of the NanoLuc sequence. Primers Pk-Aldolase-F-2 and Pk-Aldolase-R were designed to amplify a 147 bp sequence of the single copy gene Pk aldolase (access number: PKNH_1237500.1). The threshold cycle value (Ct) for each gene fragment was used to estimate the ΔCt of each sample compared with that of the single copy gene Pk aldolase, and the ΔΔCt [37] was used to calculate relative proportions of NanoLuc copies in the different samples.

The samples used in this analysis were collected from in vitro cultures containing 1 nM WR99210 (Jacobus Pharmaceuticals, Princeton, NJ) for selective pressure, without drug pressure for 14 days, and from two infected rhesus macaques (rhesus #1 and rhesus #2). One sample was collected at day four of the first infection (rhesus #1) and the second sample was collected at day six of the second infection (rhesus #2). The in vitro culture sample under drug pressure was used as reference sample.

Anti-malarial half-maximum inhibitory concentration (IC₅₀) determinations

Blood-stage P. knowlesi cultures were diluted to 0.5% parasitaemia and brought to 2% haematocrit for growth in 96-well plates containing the desired drug concentration series (200 µL of culture/well). Plates were incubated for 40 h at 37 °C in a humidified chamber containing 5% CO₂, 5% O₂, and 90% N₂. Parasite growth in the plates was determined by quantification of double stranded DNA in each well (SYBR Green method) [38] or by the luminescence measured in each well (NanoLuc method).

For the SYBR Green method, the plates were removed from the incubator and frozen overnight. DNA amounts representing parasite growth were quantified by the SYBR Green I intercalating fluorescent dye (Life Technologies) as described [39]. Plates were protected from light during the dye incubation period (30–60 min) and fluorescence intensities were measured using a FLU-Ostar Optima Microplate Reader (BMG Labtech, Ortenberg, Germany) with excitation and emission settings of 485 nm and 535 nm, respectively.
For the NanoLuc method, plates were removed from the incubator and 100 µL of NanoGlo IC₅₀ solution (Promega NanoGlo reaction mixture diluted tenfold in PBS) were added to each well and mixed by pipetting. The plates were then incubated for 3 min at room temperature and the luminescence was assessed using a plate luminometer (Molecular Devices, San Jose, CA, USA) with 1 s integration time. Fluorescence and luminescence readings were normalized to values from control wells containing no drug. IC₅₀ values were determined from fitted response curves (non-linear regression with variable slope, GraphPad Prism Software), and data from at least two independent assays were used to calculate the average IC₅₀ value of the P. knowlesi pvcen-pvhs70-D-NanoLuc transgenic line with each method.

Plasmodium knowlesi red blood cell invasion assays

Plasmodium knowlesi tightly synchronized schizonts were obtained by magnetic purification using MACS column (Miltenyi Biotec, Bergisch Gladbach, Germany) [40], adjusted to a 0.5% parasitaemia in the target RBCs and incubated for 16 h at standard culture conditions. Rhesus RBCs were used as positive controls and mouse RBCs as negative controls. Invasion of human, Aotus nancymae, and Saimiri boliviensis RBCs was evaluated in the experiments. Invasion was determined by counting the number of ring-infected RBCs per 5000 RBCs (microscopy counts), and expressed relative to the counts observed in rhesus RBCs (positive control). Assay statistics were obtained from at least three independent experiments. For NanoLuc evaluation of RBC invasion, pvcen-pvhs70-D-NanoLuc-transformed parasites were used for the invasion assays and the NanoLuc activity was assessed using the NanoGlo 20/20 luminometer as described above.

Results

Plasmodium vivax 5' UTR are broadly active in P. knowlesi but limited in P. falciparum

To evaluate the cross-species activity of different Plasmodium gene regulatory sequences, a variety of plasmids were constructed for expression of firefly luciferase (fLuc) under control of P. vivax or P. falciparum 5' UTR sequences. Plasmids pD-pvcam-Luc, pD-pvcrt-Luc and pD-pvhs70-Luc contain 5' UTR sequences from the genes for P. vivax calmodulin (pvcam), chloroquine resistance transporter (pvcrt), and heat shock protein 70 (pvhs70), respectively (Fig. 1). Plasmid pD-pfcam-Luc (Fig. 1) contains a 5' UTR sequence of the P. falciparum calmodulin gene (pfcam) and is a flanking region previously found to have cross-species promoter activity in P. knowlesi [10].

For measurements of luciferase activity, pRBC were taken from asynchronous P. falciparum or P. knowlesi cultures and transformed by introduction to culture with plasmid-loaded RBCs (spontaneous DNA uptake method) [10, 36]. Figure 2 shows the relative luciferase levels from the different transfected parasite cultures normalized to the levels from parasites transfected with the plasmid pD-pfcam-Luc. Each of the four plasmid constructs yielded signals in P. knowlesi ranging from 0.5 to 4.4 × of the pD-pfcam-Luc signal. In contrast, only two of the four plasmids provided luciferase signals in P. falciparum; no signals were obtained from P. falciparum parasites transfected with the P. vivax promoter constructs pD-pvcam-Luc and pD-pvcrt-Luc (Fig. 2, Additional file 1: Table S2). These results are consistent with a previous report suggesting that P. falciparum is unable to recognize certain P. vivax promoters for expression [22]. Nevertheless, P. falciparum parasites transfected with the plasmid pD-pvhs70-Luc presented luciferase levels similar to those from P. knowlesi transformants, 4.55 ± 0.18–fold higher (average ± standard error of the mean) than the level from parasites transfected with the control plasmid pD-pfcam-Luc.

The P. vivax hsp70 promoter sequence was also tested in P. knowlesi for expression of a drug resistance sequence used in selection of transgenic parasites. Figure 1 presents schematics of the plasmids for this purpose: pvhs70D-pfcam-Luc contains the pvhsp70 promoter sequence driving hdhfr expression that confers resistance
to antifolates such as WR99210 and pyrimethamine; pkef1D-pfcam-Luc contains the pkef1-alpha promoter previously used to drive hdblur expression for selection of transgenic parasites in vitro [9]. Both plasmids contain the fLuc reporter sequence under control of the pfcam promoter (Fig. 1). Figure 3A shows the parasite growth observed by thin blood film microscopy in the cultures transfected with the different plasmids. Parasites were not detected by thin blood films in either culture 4 days after the start of drug selection with 1 nM WR99210, but they became detectable on day 14 (parasitaemias \(= 0.1\%\)) (Additional file 1: Table S3). Figure 3B shows the luciferase activity obtained from 80 µL of RBCs collected from the cultures under selection. The luminescence signal of pvhsp70D-pfcam-Luc transgenic parasites increased an average of 1.34-fold per day for the first 12 days under selection, similar to the pkef1D-pfcam-Luc culture (average increase of 1.46-fold/day) (Fig. 3B, Additional file 1: Table S3). Over a period of three weeks of drug pressure, the transformed parasites were observed to grow at a rate similar to untransfected cultures in vitro (three to fivefold/day; data not shown).

**Rapid selection of transgenic parasites containing plasmids with P. vivax centromeric sequences**

The inclusion of centromeric sequences in plasmids and artificial chromosomes can support maintenance of exogenous DNA in transfected parasites, with proper segregation through the multiple stages of their life cycle. Examples of success with this strategy have been reported with P. berghei, P. falciparum, and Plasmodium cynomolgi [41, 42]. To improve the maintenance and segregation of a selectable marker with very high sensitivity in detection assays, a plasmid containing P. vivax centromeric sequences (pvcen) was developed and tested in P. yoelii [35]. Here, this plasmid was modified to generate pvcen-pvhsp70-D-NanoLuc, which expresses a fused-NanoLuc reporter cassette driven by the P. vivax hsp70 promoter sequence (Fig. 1).

*Plasmodium knowlesi* parasites were transfected with pvcen-pvhsp70-D-NanoLuc by the spontaneous DNA uptake method and drug-resistant parasites were selected by 1 nM WR99210 exposure. In three independent experiments, transgenic parasites containing the centromeric plasmid were detected after 8 days of drug pressure (parasitaemia \(\geq 0.1\%\)) (Fig. 3C, Additional file 1: Table S4), 6 days earlier than the detection of transgenic parasites containing episomal constructs without the centromeric sequences (Fig. 3A, Additional file 1: Table S3). The time parasites transfected with non-centromeric plasmids re-emerged under drug pressure was consistent with our previous report [10] that showed a 14–20 days period to detect drug resistant parasites after introduction of plasmid-loaded uninfected RBCs. This
period can be shortened to 4–6 days by electroporation of tightly synchronized schizonts [9, 10]. In luminescence assays, parasites transfected with the centromeric plasmid presented an average reporter activity increase of 2.73-fold/day during eight days of drug selection (Fig. 3D, Additional file 1: Table S3), greater than the rate of signal increase observed during selection of parasites transfected with the non-centromeric plasmid that also express *hdhfr* under control of the *pvhsp70* promoter (average increase of 1.34-fold/day, Fig. 3B, Additional file 1: Table S3). Over a period of two weeks’ drug pressure, transgenic parasites were observed to have a growth rate similar to untransfected parasites (three to fivefold/day) and the NanoLuc signal increased linearly with the number of parasites (Additional file 1: Table S4). In serial dilution experiments, as few as 100 transformants could be detected by NanoLuc signal above background control from uninfected RBCs (Additional file 1: Table S4).

*Plasmodium knowlesi* transformed with a *P. vivax* centromere-containing reporter plasmid completes the parasite life cycle in rhesus macaques and *An. dirus* mosquitoes

In experiments to evaluate the ability of *pvcen-pvhs*70-D-NanoLuc transformants to complete the full life cycle while expressing NanoLuc in vivo under no drug selection pressure, the parasites were inoculated into a spleenectomized rhesus macaque (rhesus #1). Parasites were observed in Giemsa-stained thin blood films at day 3 after inoculation (Fig. 4A, Additional file 1: Table S5), after which parasitaemia rapidly increased up to 2% at day 5. At midnight of day 5, when the rhesus parasitaemia reached 1.23%, *An. dirus* B and *An. dirus* X mosquitoes [31] were fed on the infected monkey; the procedure was repeated the next night at a similar parasitaemia of 1.20% (Fig. 4A; Feeds #1 and #2). Seven days after each feed, 16–20 mosquitoes were randomly selected from the individual batch for dissection and midgut isolation. As shown in Fig. 4B and Additional file 1: Table S6, Feed #1 yielded a smaller number of infected mosquitoes (11 of 16 dissected) than Feed #2 (19 of 20 dissected). Oocyst counts per mosquito were also lower in Feed #1 than in Feed #2 (Fig. 4B; 2.5 vs. 16.6 oocysts per mosquito on average, respectively). NanoLuc determinations from individual isolated midguts confirmed higher signals in proportion to the number of oocysts per midgut (Fig. 4C, Additional file 1: Table S6) ($R^2 = 0.6903$).

At day 10 after the feeds, eight mosquitoes from each batch were randomly selected and dissected for salivary glands and sporozoite detection. Total sporozoites from each group were counted, and, as expected from the oocyst data, higher sporozoite numbers were observed in mosquitoes from Feed #2 than Feed #1 (29,688 vs. 5953 sporozoites per mosquito on average, respectively); no significant difference was found between the *An. dirus* B and *An. dirus* X strains (Additional file 1: Table S7). NanoLuc activity was correspondingly higher from the greater number of sporozoites in Feed #2 mosquitoes (Fig. 4D; Additional file 1: Table S7) ($R^2 = 0.9516$).

The remaining sporozoite-infected mosquitoes were used to infect a second non-naïve macaque (rhesus #2). Six days after the mosquito bites and sporozoite injection, parasites were detected in Giemsa-stained thin blood films of peripheral blood (Fig. 4E; Additional file 1: Table S5). Development of the infection was similar to previous reports of rhesus infections using sporozoites (via mosquito bites) [31, 43]. NanoLuc activity was readily detected in blood samples, even when taken at a low 0.03% parasitaemia (data not shown).

Centromeric plasmids are present at reduced number in transformed parasites that complete the full life cycle without drug selection

During the course of rhesus macaque infection #1, transmission to mosquitoes, and infection of rhesus macaque #2, the *pvcen-pvhs*70-D-NanoLuc transformants were not subjected to drug pressure. To assess the plasmid maintenance over the life cycle in this absence of drug selection, including mitosis during the asexual parasite multiplication and meiosis after zygote formation, DNA was extracted from parasite samples and qPCR analysis was performed to quantify copy number of the NanoLuc cassette relative to the endogenous chromosome copy of the *aldolase* gene. The results indicated plasmid copy number loss during in vivo infections: parasites isolated from rhesus #1 at infection day 4 presented only half of the plasmid copy numbers of the parasites from in vitro cultures maintained for 14 days without drug pressure (Fig. 4F, Additional file 1: Table S8). Parasites isolated from rhesus #2 at infection day 6, showed a 20-fold plasmid copy number loss compared to parasites from the unpressured in vitro culture. In marked contrast, parasites from the in vitro cultures showed no plasmid copy number loss relative to the *aldolase* gene after 14 days without drug pressure (Additional file 1: Table S8).

NanoLuc facilitates high-throughput assays of anti-malarial compounds and parasite invasion

To evaluate for potential use of NanoLuc-transformed *P. knowlesi* line in assays of anti-malarial compounds, $IC_{50}$ drug response measures assessed by NanoLuc signals were compared to the measures from standard SYBR Green assays. Results were obtained from transgenic and non-transgenic (“wild-type”) parasites exposed to chloroquine, WR99210, and G418 (Table 1, Additional file 1: Table S9). No significant differences were observed
between the IC_{50} values from the two methods, and \emph{P. knowlesi} transformants presented chloroquine and G418 IC_{50} values similar to untransfected parasites (Table 1, Additional file 1: Table S9). WR99210 IC_{50} values of the transformants were about 100-fold higher, consistent with \textit{hdhfr} expression from the pvcen-pvhs70-D-NanoLuc plasmid.

Invasion assays were also performed using NanoLuc–transformed parasites. Tightly synchronized in vitro-cultivated \emph{P. knowlesi} schizonts were used to invade RBCs from different animal sources—rehesus macaque, human, \emph{Aotus nancymaeae}, \emph{Saimiri boliviensis} and mouse. Table 2 and Additional file 1: Table S10 show that no significant difference in the results was observed between the Giemsa-stained thin blood film counts and the luminescence activity of the NanoLuc-transformed parasites, confirming that luminescence can be used to evaluate invasion of RBCs.

**Fig. 4** Bioluminescent \emph{P. knowlesi} transformants complete the parasite life cycle in vivo. **A** Development of blood stage parasitaemia in a splenectomized non-naïve rhesus macaque (rhesus #1) infected with pvcen-pvhs70-D-NanoLuc-transformed \emph{P. knowlesi}. The red arrows indicate when mosquito feedings were performed; blue arrows indicate administration of chloroquine (50 mg/kg, oral). **B** Counts of oocysts in the mosquito midgut 7 days after blood feeding. **C** NanoLuc activities from midguts isolated from Feed #2 infected mosquitoes. **D** NanoLuc activities from sporozoites isolated from mosquitoes 10 days after either Feed #1 or Feed #2 on rhesus macaque DCID. **E** Blood stage parasitaemia developed in a splenectomized non-naïve rhesus macaque (rhesus #2) after bites of mosquitoes carrying infectious sporozoites. The blue arrows indicate chloroquine treatments (50 mg/kg, oral). **F** Copy numbers of the NanoLuc coding sequence relative to those of the single-copy \emph{P. knowlesi aldolase} gene in blood-stage parasites from in vitro cultures and from blood samples of rhesus monkeys #1 and #2. Copy number results from each sample are presented relative to the copy number of pvce-pvhs70-D-NanoLuc-transformed \emph{P. knowlesi} parasites cultivated under WR99210 selection pressure (black); cultivated parasites maintained 14 days in vitro without drug pressure (gray); parasites from rhesus #1 (blue); parasites from rhesus #2 (red). Error bars represent standard error of the mean; LUs, Luminescence units.
Table 1 Antimalarial half‑maximum inhibitory concentration (IC50) values obtained by SYBR Green and NanoLuc assays

| P. knowlesi parasite line | Drug     | SYBR Green method | NanoLuc method |
|--------------------------|----------|-------------------|---------------|
|                          |          | IC50   | SEM  | N   | IC50   | SEM  | N   |
| H strain (untransfected) | Chloroquine | 17.06 nM | 1.82a | 2   | NA    | NA   | NA  |
|                          | WR92910  | 0.07 nM | 0.01 | 3   | NA    | NA   | NA  |
|                          | G418     | 213.85 μM | 60.17a | 2   | NA    | NA   | NA  |
| pvccen‑pvhsp70‑D‑NanoLuc | Chloroquine | 16.81 nM | 1.91 | 3   | 16.13 nM | 1.54 | 4   |
|                          | WR92910  | 6.59 nM | 1.54 | 3   | 7.36 nM | 2.14 | 5   |
|                          | G418     | 270.80 μM | 57.13a | 2   | 239.63 μM | 22.59 | 3   |

N Number of independent biological replicates performed, SEM Standard Error of the Mean, NA not applicable
a Standard Error is presented for the IC50 values obtained using two biological replicates

Discussion

Previous evaluations of *P. vivax* regulatory sequences in *P. falciparum* showed that many were inactive [22], suggesting significant differences in the utilization of promoter elements by these species. The *P. vivax* 5′ UTRs studied in this work functioned in *P. knowlesi* even though some were inactive in *P. falciparum*. The 5′ UTRs of *pvcam* and *pvcrt* are two examples of flanking regions capable of driving the fLuc in *P. knowlesi* but not in *P. falciparum*. In contrast, the 5′ UTR of *pvhsp70* has activity in both parasites, driving at least fourfold greater expression than the control *pfcam* 5′ UTR. The longer length of the *pvhsp70* 5′ UTR (1.4 Kb) may contribute for this stronger promoter activity. Highly conserved promoter elements in the *P. vivax* heat-shock gene evidently support similar levels of transcription in transformed *P. knowlesi* and *P. falciparum* parasites. In this regard, the presence of five copies of a GCATAT element in the 5′ UTR of *pvhsp70* may be relevant as this element has been identified as a feature of some deeply conserved genes in divergent branches of *Plasmodium* evolution [22]. Absence of GCA TAT from the 5′ UTRs of *pvcam* and *pvcrt* is consistent with less conserved promoters that are operative in both *P. knowlesi* and *P. vivax* but are not recognized in *P. falciparum*. High transcription of the *hsp70* gene has been observed in *P. falciparum*, with transcripts around tenfold higher than transcripts detected of *pfcam* and *pfert* [44]. The results presented a similar pattern of transcription/expression driven by *pvhsp70*, *pvcam* and *pvcrt* 5′ UTRs, with the *pvhsp70* 5′ UTR driving a tenfold more luciferase activity than *pvcrt* and *pvcam* 5′ UTRs in *P. knowlesi*. These results also indicate that a possible activity of *pvcrt* and *pvcam* 5′ UTRs in *P. falciparum* would be detected in the experiments performed, even tenfold lower than *pvhsp70* 5′ UTR activity.

Episomal plasmids replicate by a rolling-circle mechanism leading to the formation of concatamers, resulting in variable copy numbers of the plasmid [44] and wide ranges of reporter expression. Furthermore, while episomal plasmids are useful tools for genetic studies; their maintenance requires drug pressure to prevent plasmid loss during cell division. To obviate these difficulties with copy numbers and generate transgenic lines that express reporter sequences through the complete parasite life cycle, plasmids containing centromeric sequences have been developed and used in *P. berghei*, *P. yoelii*, *P. falciparum*, and *P. cynomolgi* [35, 41, 44, 45]. Previous reports showed transgenic parasites containing centromeric plasmids were selected faster than parasites transfected with regular episomal constructs [35, 41]. Therefore, in the present study cultivated *P. knowlesi* parasites were transfected with the pvccen-pvhsp70-D-NanoLuc plasmid that contains *P. vivax* centromeric sequences and an *hdhfr* selection cassette. The resulting parasites expanded more than twice as fast as transformants containing non-centromeric plasmids, consistent with improved segregation and maintenance of the centromeric plasmid. Expression of the ultra‑bright NanoLuc reporter sequence was confirmed throughout the in vivo life cycle, in blood-stage parasites, oocysts and sporozoites, and this expression was linearly related to the number of parasites in

Table 2 Plasmodium knowlesi pvccen-pvhsp70-D-NanoLuc red blood cell invasion, determined by microscopy and NanoLuc activity

| RBC source | Microscopy | NanoLuc method |
|------------|------------|---------------|
|            | RBC invasion relative to rhesus RBCs | SEM | N   | RBC invasion relative to rhesus RBCs | N |
| Human      | 0.74       | 0.11 | 3   | 0.84 | 1   |
| Aotus nancymae | 0.53   | 0.04 | 3   | 0.34 | 1   |
| Saimiri boliviensis | 0.34 | 0.01 | 3   | 0.40 | 1   |
| Mouse      | 0.03       | 0.02 | 3   | 0.06 | 1   |

N Number of independent biological replicates performed, SEM Standard Error of the Mean
the assays. This linearity is an advantage of centromeric plasmids, that has been exploited successfully in mosquito stages, allowing its use in studies of mosquito-stage gene control and transmission blocking strategies [41]. However, without drug selection, the episomal plasmid showed copy number loss over the life cycle, particularly in vivo. This relative loss of plasmid was above expectations. Two factors could have roles in this plasmid loss. First, difference in the growth speed of asexual stage parasites in vitro vs in vivo: P. knowlesi presents a proliferation rate of three to fourfold/cycle in vitro, while the proliferation can go up to tenfold/cycle in vivo (rhesus macaques). This increase in the in vivo growth may result in a higher number of parasites without plasmid, a number that increases exponentially, explaining in part the faster decay. Second, loss of plasmid from mosquito stage parasites: the possibility that centromeric plasmids may be lost during meiosis cannot be ruled out. Nevertheless, even with the plasmid loss, the correlation between NanoLuc signals and oocyst numbers is high, and in vitro or in vivo transmission experiments can be performed using bioluminescence assays through at least in one cycle of sexual transmission. This finding highlights an advantage of transgenic lines that contain reporter sequences integrated into chromosomes of the parasite genome, achievable for example by recently developed zinc-finger gene modification methods [34, 46] or by CRISPR/Cas9 technology [47, 48]. Whether the persistence of centromere-containing plasmids in P. knowlesi can be improved by use of homologous pvcen instead of pvcen elements remains to be investigated.

Rapid assay determinations and sensitivity allowing the use of low parasite numbers are among the advantages of NanoLuc-expressing parasites for high-throughput screening platforms. In the present work, NanoLuc expressing P. knowlesi were successfully used in drug response tests of blood-stage parasites in vitro, and the IC50 values obtained by luminescence assays were comparable to IC50 values obtained by the SYBR green method. Additionally to their use in drug response, centromeric plasmids such as pvcen-pvhsp70-D-NanoLuc may be useful in other experimental applications such as in vivo imaging. The P. knowlesi transformants can also be useful for validation of plasmid constructs prior to their use in transfection of P. vivax, which requires expensive non-human primate resources. The ultrabright and stable NanoLuc reporter has been previously used in P. falciparum [49] and P. berghei [29, 50], producing luminescence up to 150-fold higher than firefly luciferase, and enabling detection of parasites at low densities and as single sporozoite [50]. Recently developed methods of mosquito infection from in vitro P. knowlesi cultures [11] may thus provide useful strategies in high-sensitivity screens for transmission blocking vaccines and inhibitors of liver-stage parasites [29, 49, 50].

**Conclusions**

*Plasmodium knowlesi* parasites can drive expression of P. vivax promoters that are poorly recognized by P. falciparum, consistent with the closer evolutionary relationship of P. vivax to P. knowlesi than to P. falciparum. However, more conserved promoters like that of pvhsp70 retain similar recognition by P. vivax and P. falciparum. Centromeric sequences from P. vivax can improve the efficiency of genetic transformation and may promote the stability of transfected plasmids across the complete P. knowlesi life cycle, through rhesus monkeys and An. dirus mosquitoes. NanoLuc-expressing, centromere-stabilized plasmids may be useful for high-throughput anti-malarial screenings as well as biological imaging of P. knowlesi in the mosquito and vertebrate hosts.

**Supplementary Information**

The online version contains supplementary material available at https://doi.org/10.1186/s12936-021-03773-4.

**Additional file 1:** Table S1. Oligonucleotide primers used in this work.

Table S2. Transient transfection of Plasmodium parasites. Table S3. Plasmodium knowlesi growth and luminescence signal during selection using WR99210. Table S4. NanoLuc signal of in vitro cultivated transgenic blood-stage parasites. Table S5. Rhesus infections with transgenic luminescent P. knowlesi. Table S6. Number of oocysts detected in individual mosquitoes dissected 7 days after the feeds and Luminescence signals obtained (only midguts isolated from mosquitoes from Feed #2). Table S7. Total sporozoites isolated from 8 mosquitoes from each species, infected with transgenic bioluminescent P. knowlesi, 10 days after the feeds and Luminescence signals. Table S8. pvcen-pvhsp70D-NanoLuc plasmid copy number variation in blood-stage parasites. Table S9. Half-maximum inhibitory concentration (IC50) measures of drug response. Table S10. Plasmodium knowlesi red blood cell invasion assays.

**Acknowledgements**

We thank Patrick E. Duffy for providing the P. knowlesi H strain, rhesus blood samples for in vitro cultivation of parasites, and animal infections; Billeta Lewis, Milton J. Herrera, and Kelly E. Dicken, for monkey care; and Andre Laughing-house and Kevin L. Lee, for providing and maintaining mosquitoes.

**Authors’ contributions**

RRMB, JSA, JMS, JHA, OK, and TEW designed the study. RRMB, KT, JSA, TIG, RLC, MK, WAK, JPM, JKB, TE, LL, SOG performed experiments. RRMB, JSA, TIG,
JMS and EW wrote the manuscript. All authors read and approved the final manuscript.

Funding
This study was supported by the Intramural Research Program of the National Institute of Allergy and Infectious Diseases, National Institutes of Health and Bill & Melinda Gates Foundation (OPP1023643 to JHA), Grants-in-Aid for Scientific Research 23659215 (OK), MEXT, Japan and CNPq Universal Grant (434011/2018-5) and FAPESP JP award 2018/06219-8 (RMB), Brazil. This work was partly conducted at the Joint Usage/Research Center for Tropical Disease, Institute of Tropical Medicine, Nagasaki University, Japan.

Availability of data and materials
All data generated or analysed in this study are provided in the main text of this published article or as additional files.

Declarations

Ethics approval and consent to participate
All Non-human primate care and use in this study were performed in accordance with the NIH Animal Research Advisory Committee (NIH ARAC) Guidelines, under an Animal Study Proposal approved by the NIAID Animal Care and Use Committee (NIAIDLMIV 9E).

Consent for publication
Not applicable.

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

Author details
1 Laboratory of Malaria and Vector Research, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, MD, USA. 2 Departamento de Microbiologia, Imunologia e Parasitologia, Escola Paulista de Medicina, Universidade Federal de São Paulo, São Paulo, SP, Brazil. 3 Departamento de Protozoologia, Institute of Tropical Medicine (NEKKEN), Nagasaki University, Nagasaki, Japan. 4 Department of Medical Technology, Faculty of Allied Health Sciences, Naesuan University, Phitsanuklok, Thailand. 5 Center for Global Health and Infectious Diseases Research, College of Public Health, University of South Florida, Tampa, FL, USA. 6 Department of Molecular Microbiology and Immunology, Graduate School of Biomedical Sciences, Nagasaki University, Nagasaki, Japan. 7 Division of Veterinary Research, National Institutes of Health, Bethesda, MD, USA. 8 Laboratory of Malaria Immunology and Virology, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, MD, USA.

Received: 6 January 2021 Accepted: 16 May 2021 Published online: 05 June 2021

References
1. Yusof R, Lau YL, Mahmud R, Fong MY, Jelip J, Ngian HU, et al. High proportion of knowlesi malaria in recent malaria cases in Malaysia. Malar J. 2014;13:168.
2. Singh B, Daneshvar C. Human infections and detection of Plasmodium knowlesi. Clin Microbiol Rev. 2013;26:165–84.
3. William T, Jelip J, Menon J, Andeiro F, Mohammad R, Awang Mohammad TA, et al. Changing epidemiology of malaria in Sabah, Malaysia: Increasing incidence of Plasmodium knowlesi. Malar J. 2014;13:390.
4. William T, Rahman HA, Jelip J, Ibrahim MY, Menon J, Grigg MJ, et al. Increasing incidence of Plasmodium knowlesi Malaria following control of Plasmodium falciparum and P. vivax Malaria in Sabah, Malaysia. PLoS Negl Trop Dis. 2013;7:e2026.
5. WHO. World malaria report 2017. Geneva: World Health Organization; 2017. http://apps.who.int/iris/bitstream/10665/259492/1/9789241565523-en.pdf?ua=1.
6. Trager W, Jensen JB. Human malaria parasites in continuous culture. Science. 1976;193:673–5.
7. Butcher GA, Mitchell GH. The role of Plasmodium knowlesi in the history of malaria research. Parasitology. 2016;145:6–17.
8. Kocken CHM, Dzwaya H, Van der Wel A, Beetsma AL, Mwenda JM, Thomas AW. Plasmodium knowlesi provides a rapid in vitro and in vivo transfection system that enables double-crossover gene knockout studies. Infect Immun. 2002;70:655–60.
9. Moon RW, Hall J, Rangkuti F, Ho YS, Almond N, Mitchell GH, et al. Adaptation of the genetically tractable malaria pathogen Plasmodium knowlesi to continuous culture in human erythrocytes. Proc Natl Acad Sci. 2013;110:531–6.
10. Moraes Barros RR, Gibson TJ, Kite WA, Sá JM, Wellemes TE. Comparison of two methods for transformation of Plasmodium knowlesi: Direct schizont electroperoration and spontaneous plasmid uptake from plasmid-loaded red blood cells. Mol Biochem Parasitol. 2017;218:16–22.
11. Armistead JS, Moraes Barros RR, Gibson TJ, Kite WA, Messhron JP, Lambert LE, et al. Infection of mosquitoes from in vitro cultivated Plasmodium knowlesi H strain. Int J Parasitol. 2018;48:601–10.
12. Hawking F, Worms MJ, Gammage K. Host temperature and control of 24-hour and 48-hour cycles in malaria parasites. Lancet. 1968;291:506–9.
13. Rijo-Ferreira F, Acosta-Rodriguez VA, Abel JH, Kornblum I, Benito L, Kilaru G, et al. The malaria parasite has an intrinsic clock. Science. 2020;368:746–53.
14. Adams JH, Sim BK, Dolan SA, Fang X, Kaslow DC, Miller LH. A family of erythrocyte binding proteins of malaria parasites. Proc Natl Acad Sci USA. 1992;89:7085–9.
15. Hayton K, Gaur D, Liu A, Takahashi J, Henschen S, Singh S, et al. Erythrocyte binding protein PRRHS polymorphisms determine species-specific pathways of Plasmodium falciparum infection. Cell Host Microbe. 2008;4:40–51.
16. Cowman AF, Tonkin CJ, Tham W-H, Duraisingham MT. The Molecular basis of erythrocyte invasion by malaria parasites. Cell Host Microbe. 2017;22:232–45.
17. Gautret P, Motard A. Periodic infectivity of Plasmodium gametocytes to the vector. A review. Parasite. 1999;6:103–11.
18. Chotivanich K, Udornsangphet R, Chierakul W, Newton PN, Ruangveerayuth P, Prukittayakamee S, et al. In vitro efficacy of antimalarial drugs against Plasmodium vivax on the western border of Thailand. Am J Trop Med Hyg. 2004;70:395–7.
19. Nomura T, Carlson JM, Baird JK, del Portillo HA, Fryauff DJ, Rathore D, et al. Evidence for different mechanisms of chloroquine resistance in Plasmodium species that cause human malaria. J Infect Dis. 2001;183:1653–61.
20. Amino R, Thiberge S, Martin B, Celli S, Shorte S, Frischknecht F, et al. Quantitative imaging of Plasmodium transmission from mosquito to mammal. Nat Med. 2006;12:220–4.
21. Deitsch KW, Calderwood WS, Wellemes TE. Malaria. Cooperative silencing elements in var genes. Nature. 2001;412:875–6.
22. Azavedo MF, Del Portillo HA. Promoter regions of Plasmodium vivax are poorly or not recognized by Plasmodium falciparum. Malar J. 2007;6:20.
23. Calderwood WS, Gannoun-Zaki L, Wellemes TE, Deitsch KW. Plasmodium falciparum var Genes are regulated by two regions with separate promoters, one upstream of the coding region and a second within the intron. J Biol Chem. 2003;278:34125–32.
24. De Koning-Ward TF, Speranca MA, Waters AP, Janse CJ. Analysis of stage specificity of promoters in Plasmodium berghei using luciferase as a reporter. Mol Biochem Parasitol. 1999;100:141–6.
25. De Niz M, Helm S, Horstmann S, Annoura T, del Portillo HA, Khan SM, et al. In vivo and in vitro characterization of a Plasmodium liver stage-specific promoter. PLoS ONE. 2015;10:e0124734.
26. Hasenkamp S, Sidaway A, Devine O, Roye R, Horrocks P. Evaluation of bioluminescence-based assays of anti-malarial drug activity. Malar J. 2013;12:58.
27. Swann J, Corey V, Scherer CA, Kato N, Comer E, Maetani M, et al. High-throughput luciferase-based assay for the discovery of therapeutics that prevent malaria. ACS Infect Dis. 2016;2:281–93.
28. Lin JW, Sajid M, Ramesar J, Khan SM, Janse CJ, Franke-Fayard B. Screening inhibitors of P. berghei blood stages using bioluminescent reporter parasites. Methods Mol Biol. 2013;923:507–22.
29. Calit J, Dobrescu I, Gaitán XA, Borges MH, Ramos MS, Eastman RT, et al. Screening the Pathogen Box against Plasmodium sexual stages using a new nanoluciferase based transgenic line of P. berghei identifies transmission-blocking compounds. Antimicrob Agents Chemother. 2018;62:e01053–18.
30. Walliker D, Quakyi IA, Wellems TE, McCutchan TF, Szarfman A, London WT, et al. Genetic analysis of the human malaria parasite Plasmodium falciparum. Science. 1987;236:1661–6.

31. Murphy JR, Weiss WR, Fyrauff D, Dowler M, Savransky T, Stoyanov C, et al. Using infective mosquitoes to challenge monkeys with Plasmodium knowlesi in malaria vaccine studies. Malar J. 2014;13:215.

32. Zeeman AM, Voorberg-Van Der Wel A, Kocken CHM. Ex vivo culture of Plasmodium vivax and Plasmodium cynomolgi and in vitro culture of plasmodium knowlesi blood stages. Methods Mol Biol. 2013;923:35–49.

33. Sá JM, Kaslow SR, Moraes Barros RR, Brazeau NF, Parobek CM, Tao D, et al. Plasmodium vivax chloroquine resistance links to pvcrt transcription in a genetic cross. Nat Commun. 2019;10:4300.

34. Barros RRM, Stainer J, Sa JM, Salzman RE, Melendez-Muniz VA, Mu J, et al. Editing the Plasmodium vivax genome, using zinc-finger nucleases. J Infect Dis. 2015;211:125–9.

35. Thawnashom K, Kaneko M, Xangsayarath P, Chaiyawong N, Yahata K, Asada M, et al. Validation of Plasmodium vivax centromere and promoter activities using Plasmodium yoelii. PLoS ONE. 2019;14:e0226884.

36. Deitsch K, Driskill C, Wellems T. Transformation of malaria parasites by the spontaneous uptake and expression of DNA from human erythrocytes. Nucleic Acids Res. 2001;29:850–3.

37. Livak KJ, Schmittgen TD. Analysis of relative gene expression data using real-time quantitative PCR and the 2-DDCT method. Methods. 2001;25:402–8.

38. Smilkstein M, Sriwilaijaroen N, Kelly JX, Wilairat P, Riscoe M. Simple and inexpensive fluorescence-based technique for high-throughput antimalarial drug screening. Antimicrob Agents Chemother. 2007;51:1926–33.

39. Johnson JD, Dennull RA, Gerena L, Lopez-Rubio J. Genome editing in the human malaria parasite Plasmodium falciparum using engineered zinc-finger nucleases. Nat Methods. 2012;9:993–8.

40. Ribaut C, Berry A, Chevalley S, Reybier K, Morlais I, Parzy D, et al. Concentration and purification by magnetic separation of the erythrocytic stages of all human Plasmodium species. Malar J. 2008;7:45.

41. Iwanaga S, Khan SM, Kaneko I, Christodoulou Z, Newbold C, Yuda M, et al. Functional identification of the Plasmodium centromere and generation of a Plasmodium artificial chromosome. Cell Host Microbe. 2010;7:245–55.

42. Voorberg-Van der Wel A, Zeeman AM, van Amsterdam SM, van den Berg A, Klooster EJ, Iwanaga S, et al. Transgenic fluorescent Plasmodium cynomolgi liver stages enable live imaging and purification of malaria hypnozoite-forms. PLoS ONE. 2013;8:e54888.

43. Mahdi Abdel Hamid M, Remarque EJ, El Hassan IM, Hussain AA, Narum DL, Thomas AW, et al. Malaria infection by sporozoite challenge induces high functional antibody titres against blood stage antigens after a DNA prime, poxvirus boost vaccination strategy in Rhesus macaques. Malar J. 2011;10:29.

44. Williamson DH, Janse CJ, Moore PW, Waters AP, Preiser PR. Topology and replication of a nuclear episomal plasmid in the rodent malaria Plasmodium berghei. Nucleic Acids Res. 2002;30:726–31.

45. Voorberg-van der Wel AM, Zeeman AM, Nieuwenhuis IG, van der Werff NM, Klooster EJ, Kloop O, et al. A dual fluorescent Plasmodium cynomolgi reporter line reveals in vitro malaria hypnozoite reactivation. Commun Biol. 2020;3:7.

46. Straimer J, Lee MCS, Lee AH, Zeitler B, Williams AE, Pearl JR, et al. Site-specific genome editing in Plasmodium falciparum using engineered zinc-finger nucleases. Nat Methods. 2012;9:993–8.

47. Ghorbal M, Gorman M, MacPherson CR, Martins RM, Scherf A, Lopez-Rubio J. Genome editing in the human malaria parasite Plasmodium falciparum using the CRISPR-Cas9 system. Nat Biotechnol. 2014;32:819–21.

48. Mohring F, Hart MN, Rawlinson TA, Henriuci R, Charleston JA, Diez Benavente E, et al. Rapid and iterative genome editing in malaria parasite Plasmodium knowlesi provides new tools for P. vivax research. Elife. 2019;8:e45829.

49. Azevedo MF, Nie CQ, Elsworth B, Charnaud SC, Sanders PR, Crabb BS, et al. NanoLuc-based luminescence system for monitoring Plasmodium falciparum transfected with ultra bright nanoluc luciferase provides new tools for P. vivax research. Commun Biol. 2019;2:112571.

50. De Niz M, Stainer RR, Wacker R, Keller D, Heussler VT. An ultrasensitive reporter line reveals in vitro malaria hypnozoite reactivation. Commun Biol. 2019;2:112571.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.