A 39-Amino-Acid C-Terminal Truncation of GDV1 Disrupts Sexual Commitment in *Plasmodium falciparum*

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ABSTRACT  Malaria is a mosquito-borne disease caused by apicomplexan parasites of the genus *Plasmodium*. Completion of the parasite’s life cycle depends on the transmission of sexual stages, the gametocytes, from an infected human host to the mosquito vector. Sexual commitment occurs in only a small fraction of asexual blood-stage parasites and is initiated by external cues. The gametocyte development protein 1 (GDV1) has been described as a key facilitator to trigger sexual commitment. GDV1 interacts with the silencing factor heterochromatin protein 1 (HP1), leading to its dissociation from heterochromatic DNA at the genomic locus encoding AP2-G, the master transcription factor of gametocytogenesis. How this process is regulated is not known. In this study, we have addressed the role of protein kinases implicated in gametocyte development. From a pool of available protein kinase knockout (KO) lines, we identified two kinase knockout lines which fail to produce gametocytes. However, independent genetic verification revealed that both kinases are not required for gametocytogenesis but that both lines harbor the same mutation that leads to a truncation in the extreme C terminus of GDV1. Introduction of the identified nonsense mutation into the genome of wild-type parasite lines replicates the observed phenotype. Using a GDV1 overexpression line, we show that the truncation in the GDV1 C terminus does not interfere with the nuclear import of GDV1 or its interaction with HP1 in vitro but appears to be important to sustain GDV1 protein levels and thereby sexual commitment.

IMPORTANCE  Transmission of malaria-causing *Plasmodium* species by mosquitoes requires the parasite to change from a continuously growing asexual parasite form growing in the blood to a sexually differentiated form, the gametocyte. Only a small subset of asexual parasites differentiates into gametocytes that are taken up by the mosquito. Transmission represents a bottleneck in the life cycle of the parasite, so a molecular understanding of the events that lead to stage conversion may identify novel intervention points. Here, we screened a subset of kinases implicated in gametocyte development. From a pool of available protein kinase knockout (KO) lines, we identified two kinase knockout lines which fail to produce gametocytes. However, independent genetic verification revealed that both kinases are not required for gametocytogenesis but that both lines harbor the same mutation that leads to a truncation in the extreme C terminus of GDV1. Introduction of the identified nonsense mutation into the genome of wild-type parasite lines replicates the observed phenotype. Using an GDV1 overexpression line, we show that the truncation in the GDV1 C terminus does not interfere with the nuclear import of GDV1 or its interaction with HP1 in vitro but appears to be important to sustain GDV1 protein levels and thereby sexual commitment.

KEYWORDS  GDV1, gametocytes, *Plasmodium falciparum*, transmission, kinases

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Malaria is a devastating disease caused by parasites of the genus *Plasmodium*, leading to ~405,000 deaths per year (1). *Plasmodium falciparum* causes the most severe and life-threatening form of human malaria. The complex life cycle involves interactions with multiple tissues in two different organisms, the human host and the mosquito vector. Inside the human host, *P. falciparum* predominantly infects red blood cells (RBC) where it asexually replicates. A small fraction (0 to 20%) of parasites commits to sexual development (gametocytogenesis) (2). Gametocytogenesis occurs preferentially in the extravascular compartment in the bone marrow and spleen (3–9). After 10 to 12 days, mature stage V gametocytes are released into the peripheral circulation to allow transmission to mosquitoes.

Sexual commitment can be initiated by metabolic cues in the human host. Specifically, it has been described that depletion of lysocephatidylcholine (LysoPC), a common component of human serum, leads to increased rates of gamocyte production and therefore represents the first molecularly defined factor known to inhibit or trigger sexual conversion (10). Sexual commitment depends on upregulation of the *ap2-g* gene (2, 11), which requires removal of heterochromatin protein 1 (HP1) from chromatin. HP1 interacts directly with the gametocyte development 1 protein (GDV1), which causes HP1 to dissociate (12). HP1 is responsible for repression of a range of genes (13), while GDV1 specifically acts on the *ap2-g* locus. How this specificity is achieved is not known. Furthermore, how a drop in LysoPC levels is sensed and transduced into GDV1-mediated HP1 removal is not understood.

Kinases are key transducers of signals in cellular processes in various stages of the *Plasmodium* life cycle (14, 15) and are likely candidates to play important roles in gametocyte commitment and development. A study by Solyakov et al. (14) has identified a panel of likely and confirmed nonessential protein kinases, some of which are transcribed during sexual development (PlasmoDB) or in gametocytes (16–19). Aiming to identify protein kinases involved in sexual development, we screened eight knockout (KO) lines for phenotypes in gametocyte induction and/or maturation. Two lines made no gametocytes, but subsequent validation showed that their gametocytogenesis defect was not due to the absence of these kinases. Instead, we found that both lines shared the same truncation in the C-terminal end of GDV1, which caused the loss of gametocyte development. Here, we address the importance and role of the GDV1 C-terminal for sexual commitment and interaction with HP1. We show that the loss of the C-terminal 39 amino acids of GDV1 does not interfere with nuclear import and interaction with HP1 in vitro but prevents GDV1 from triggering efficient sexual commitment.

### RESULTS

**Identification and characterization of *Plasmodium falciparum* kinase KO lines with a gametocytogenesis phenotype.** It has been shown previously in *P. berghei* that protein kinases that are nonessential during the asexual blood stages are essential in other life cycle stages, for example, during parasite transmission in the mosquito (15). To identify kinases important for gametocytogenesis, we investigated the role of a group of likely nonessential kinases (14) during asexual blood stages development. Using the lines described by Solyakov et al. (14), which have been generated by single crossover gene disruption, we induced sexual development using conditioned medium (20) and followed progression through stages I to V of gametocytogenesis (Fig. 1A). Six of the eight KO lines displayed normal gametocyte development, while two, TKL2 (PF3D7_1121300) and eIK2 (PF3D7_0107600) kinase KO lines, produced very few (<0.1%) gametocytes (Fig. 1B). Of these, one has a disrupted tyrosine kinase-like 2 (*tkl2*) locus, which has been characterized as a protein kinase secreted outside the red blood cell (17). Gene loss and accumulation of mutations are frequently observed in parasite lines kept in continuous in vitro culture over time, and the loss of the ability to form gametocytes is not uncommon (21). To exclude mutations in the *ap2-g* gene, which was identified previously through a loss-of-function mutant (2), we sequenced the *ap2-g* locus in the 3D7/TKL2 KO parasite line. The sequencing results confirmed that the phenotype observed was not associated with mutations in *ap2-g*, leading us
to conclude that the deletion of TKL2 was possibly the cause for the observed phenotype. In order to verify the role of TKL2 in gametocyte induction, we generated a DiCre-mediated TKL2 conditional KO line in NF54 parasites (NF54/TKL2:loxPint). We used CRISPR/Cas9 to simultaneously introduce a DiCre cassette into the pfs47 locus, as previously described (22, 23), and to flank the kinase domain of tkl2 with two loxPints (Fig. 1C and Fig. S1A and B). To address the role of TKL2 in gametocyte development, we treated the NF54/TKL2:loxPint line with dimethyl sulfoxide (DMSO; control) or rapamycin (KO) (Fig. S1B). We then induced sexual commitment using conditioned medium (20) and monitored gametocyte development. No difference in commitment or development between the control and the rapamycin-induced NF54/TKL2:loxPint parasites (Fig. 1D) was observed. These results show that TKL2 is not involved in sexual commitment or gametocyte development/maturation and that another mutation is likely the cause for the observed phenotype.

A common GDV1 truncation is found in both kinase KO lines deficient in gametocyte formation. The second kinase KO where a gametocytogenesis defect was identified was the eukaryotic initiation factor serine/threonine kinase 2 (eIK2) KO line.
(3D7/eIK2 KO) (Fig. S2A). eIK2 has previously been characterized as nonessential during sexual development in *P. falciparum* and *P. berghei*, and eIK2 KO lines appeared to undergo normal gametocyte development in rodent *Plasmodium* species (24). This indicated that, as does 3D7/TKL2 KO, the 3D7/eIK2 KO line also harbors a mutation preventing efficient gametocyte development. Sequencing of the *ap2-g* locus in this parasite line as described above showed no mutations in the coding region of *ap2-g*. Therefore, a potentially unknown mutation underlies the loss of gametocytes in these parasite lines.

To understand the nature of the block in sexual development, we analyzed the transcriptome of induced wild-type 3D7 parasites and two eIK2 KO clones (clones C3 and F12) using transcriptome sequencing (RNA-seq). Samples were collected for RNA extraction between 28 and 32 h postinvasion (hpi) after induction with conditioned medium (Fig. 2A). The RNA-seq analysis revealed a significant downregulation in 3D7/eIK2 KO parasites of genes known to be upregulated during gametocytogenesis, including genes that have been shown to be AP2-G-dependent (2, 10, 12, 25–27) (Fig. 2B and Table S1). We found *ap2-g* itself to be downregulated in 3D7/eIK2 KO parasites, but this reached significance only in one of the clones. Together with the lack of mutations in *ap2-g* itself, these results suggested that the block in gametocytogenesis was upstream of AP2-G function during sexual commitment. At that time, GDV1 was shown to be an upstream activator of AP2-G expression (12), so we sequenced the *gdv1* locus in the eIK KO clones and identified a nonsense mutation in *gdv1* that results in a premature stop codon leading to a C-terminal truncation of 39 amino acids (*GDV1 D39*) (Fig. 2C and Fig. S2B). Sequencing of the 3D7/TKL2 KO parasite clones showed the same mutation (Fig. S2B), suggesting that the deletion of the last 39 amino acids of GDV1 in both mutant lines is responsible for the gametocytogenesis phenotype observed in both kinase KO lines.

**The carboxy-terminal 39 amino acids of GDV1 are important for its function.** To verify genetically the identified mutation in *gdv1*, we generated a 3× hemagglutinin (3×HA)-tagged version of GDV1Δ39 and introduced it in the endogenous *gdv1* locus in the NF54 parasite line (NF54/GDV1Δ39:HA) (Fig. 3A and Fig. S3A and B). GDV1Δ39:HA parasites lost the ability to form gametocytes (Fig. 3B), suggesting that the GDV1 C terminus plays an essential role during sexual commitment or development. Determination of the localization or expression levels of GDV1Δ39:HA was not possible, as we could not confidently distinguish true signal from background fluorescence. We repeatedly failed to obtain parasites expressing 3×HA-tagged full-length GDV1 from the endogenous locus to compare its expression levels and the localization to those of the truncated GDV1 version. Notably, direct C-terminal tagging of GDV1 at the endogenous locus was also not successful in other studies, unless when in combination with a destabilization domain (12, 28).

Therefore, we resorted to a system that allows robust testing of GDV1-dependent gametocyte induction by conditional overexpression of GDV1-green fluorescent protein (GFP) and quantifying sexual conversion (29). To do that, we first introduced an ectopic *gdv1-gfp* fusion gene under the control of the calmodulin promoter and a *glmS* ribozyme in the 3’ untranslated region into the *cg6* (*glp3, PF3D7_0709200*) locus in NF54 parasites. In the presence of glucosamine, the *glmS* ribozyme destabilizes the mRNA preventing GDV1:GFP expression, while in the absence of glucosamine GDV1:GFP is overexpressed, leading to gametocyte induction (30). For simplicity, the NF54/iGP2 line described by Boltryk and colleagues (29) has been renamed NF54/GDV1:GFP_cOE in this study (cOE stands for conditional overexpression).

To test GDV1Δ39 function, we introduced a *gdv1Δ39-gfp-glmS* cassette into the *cg6* locus, generating a conditional GDV1Δ39:GFP overexpression parasite line (NF54/GDV1Δ39:GFP_cOE) (Fig. 3D and Fig. S3C and D). We then compared the sexual conversion rates in the NF54/GDV1:GFP_cOE and NF54/GDV1Δ39:GFP_cOE parasites in the presence and absence of glucosamine. In contrast to that of GDV1:GFP, overexpression of GDV1Δ39:GFP failed to trigger a significant increase of sexual commitment (Fig. 3D).
These results suggest that the full integrity of the GDV1 C terminus is important for sexual commitment.

**GDV1Δ39 is imported into the nucleus and retains the ability to interact with HP1.** GDV1 is a nuclear protein, and we hypothesized that the deletion of a predicted C-terminal nuclear bipartite localization sequence (nLS mapper, http://nls-mapper.iab.keio.ac.jp/cgi-bin/NLS_Mapper_form.cgi) may interfere with GDV1 nuclear localization and hence its ability to interact with HP1 at heterochromatic loci. Therefore, we localized induced GDV1:GFP and GDV1Δ39:GFP by immunofluorescence at 28 to 32 hpi (see Fig. 4A for reference). As expected, control GDV1:GFP_cOE parasites showed a clear punctate and nuclear GDV1:GFP signal (Fig. S3E) (12). GDV1Δ39:GFP_cOE also showed localized GDV1Δ39:GFP signal in the nucleus, but the signal was weaker and more diffuse compared with that of GDV1:GFP (Fig. S3F). In order to quantify and compare GFP levels in GDV1:GFP_cOE and GDV1Δ39:GFP_cOE parasites, we performed a whole-cell protein extraction for Western blotting using GFP-specific antibodies. HSP70-specific antibodies were used as a loading control (Fig. 4B and C). The Western blot (WB) showed a clear reduction of GDV1Δ39:GFP compared to GDV1:GFP (Fig. 4C).

To quantify the localization of GDV1Δ39:GFP in the cytoplasm compared to that in the...
nucleus, we prepared cytosolic and nuclear protein extracts using subcellular fractionation (Fig. 4B and D). We determined the cytoplasmic fraction using anti-aldolase antibodies (31), and anti-histone 3 antibodies were used to determine the nuclear fraction (32). GDV1\textsuperscript{D39}:GFP was only detected in the nuclear fraction, further supporting that its nuclear localization is not affected by the C-terminal truncation (Fig. 4D). To test if the GDV1\textsuperscript{D39} deletion affects its interaction with HP1, we performed an in vitro assay where 6×His-tagged GDV1 (His-GDV1) and GDV1\textsuperscript{D39} (His-GDV1\textsuperscript{D39}) versions were coexpressed with Strep-tagged HP1 in Escherichia coli bacteria. Interaction between His-GDV1 and Strep-HP1 is detected by affinity purification of His:GDV1 and analysis of coeluted proteins by Coomassie staining (12). His-tagged SIP2 does not interact with Strep-HP1 and was used as a negative control (Fig. 4E). As previously shown, His-GDV1 pulled down HP1, which was not observed when SIP2 was used as a bait (12). Interestingly, His-GDV1\textsuperscript{D39} also pulled down HP1, showing that the GDV1 C terminus is not essential for the interaction in E. coli (Fig. 4E). This observation indicates that the
FIG 4  GDV1Δ39 expression, localization, and interaction with HP1. (A) Representation of the protocol used to collect the samples used to characterize expression and localization of GDV1Δ39:GFP. (B) Illustration of the subcellular fractionation workflow. (C) Western blot (Continued on next page)
interaction of GDV1Δ39:GFP and HP1 could still occur in the parasite but that it is insufficient to trigger gametocytogenesis.

An explanation for the lack of induction despite the apparent correct localization could be that GDV1Δ39:GFP levels do not reach the threshold required for efficient gametocyte induction. To examine expression of the GDV1Δ39:GFP, we analyzed protein levels at the single-cell level using flow cytometry in uninduced and induced NF54/GDV1:GFP_cOE and NF54/GDV1Δ39:GFP_cOE parasites (Fig. 4F and G and Fig. S4). As expected, NF54/GDV1:GFP_cOE parasites show a robust increase of GDV1:GFP expression through glucosamine removal. A measurable increase of the mean fluorescence of GDV1Δ39:GFP was also observed upon induction in most NF54/GDV1Δ39:GFP_cOE parasites but at levels well below the GDV1:GFP levels observed for NF54/GDV1:GFP_cOE parasites. The reduction on the protein levels of GDV1Δ39:GFP compared to those of GDV1:GFP is not explained by differences in transcript level. Quantitative reverse transcription-PCR (RT-qPCR) shows ~1.5-fold higher transcript levels of gdvΔ39:gfp compared to those of gdv1:gf, while both lines show an equal increase upon induction (~2-fold) (Fig. S4D and E). The increase of transcript levels for gdv1Δ39:gf is likely explained by an additional copy of the plasmid that integrated into the genome: qPCR analysis of genomic DNA from NF54/GDV1Δ39:GFP_cOE parasites using primers specific for GFP showed an ~1.8-fold increase of DNA content compared to that of genomic DNA (gDNA) from NF54/GDV1:GFP_cOE parasites (data not shown). Interestingly, a small proportion of NF54/GDV1Δ39:GFP_cOE parasites displayed GDV1Δ39:GFP fluorescence at a level similar to that of GDV1:GFP in the NF54/GDV1:GFP_cOE control line. In line with its nuclear localization, GDV1Δ39:GFP may therefore contribute to the formation of gametocytes in these parasites.

DISCUSSION

The aim of this study was to identify nonessential kinases as regulators of gametocyte commitment/development in P. falciparum. While several parasite lines of the kinase knockout collection (14) were able to form gametocytes, two kinase KO lines showed a gametocytogenesis phenotype that led to the identification of a nonsense mutation in gdv1 that results in a 39-amino-acid (aa) truncation of the GDV1 C terminus. This mutation may have been acquired by the common parental line prior to generation of the original transgenic lines, although several other clones from the Solyakov study that we tested here are able to form gametocytes, possibly reflecting that only a proportion of the parasite population in the parental line carried the mutation. Alternatively, it cannot be excluded that the mutation arose independently in these two lines. Regardless of the origin of the mutations, our results show that the premature stop codon mutation in gdv1 resulting in a 39-amino-acid C-terminal truncation in the tk2 and elk2 KO lines is sufficient to abolish sexual commitment. Loss of gametocytogenesis is a common occurrence in culture-adapted strains (2), and spontaneous loss-of-function mutations for gdv1 leading to loss of gametocytogenesis were previously identified (33, 34). Our results extend previous studies of GDV1 truncations: we observed that the truncated GDV1Δ39:GFP protein was present at substantially reduced levels compared to those of full-length GDV1:GFP.
We propose that the GDV1Δ39:GFP truncation leads to reduced GDV1 protein stability, which is most likely the underlying cause for the lack of gametocytes in the GDV1Δ39 mutants. However, the truncation of GDV1 results neither in a strong nuclear localization defect when overexpressed as a GFP fusion protein nor in a failure to interact with HP1 expressed in bacteria. It will be important to show in the future whether the few NF54/GDV1,Δ39:GFP_cOE parasites, which show similar levels of GDV1,Δ39:GFP to those of GDV1:GFP in NF54/GDV1:GFP_cOE parasites, are able to induce gametocytogenesis. If they fail to do so, it could point toward additional functions of the GDV1 C terminus, potentially contributing to bringing GDV1 to the ap2-g locus through interactions with a yet unknown protein.

MATERIALS AND METHODS

Plasmid construction and transfection. The construction of each of the ePK knockout plasmids here characterized has been described in reference 14. The pmK-RQ-tkl2-loxPint donor plasmid (synthesized by Geneart) contains a recodonized (rc) version of sequence containing the glycine-rich loop in the kinase domain of tkl2 (rc. Gly loop) flanked by two loxPins and homology regions for homology-directed repair. The pDC2-Cas9-hDHFRyFCU guideRNA plasmid targeting tkl2 locus (pDC2_TKL2_gRNA) was generated using the primer pairs pDC2_TKL2_gRNA1_FOR/pDC2_TKL2_gRNA1_REV. Because we did not have a 3D7:DiCre line, we generated the 3D7:TKL2:loxPint conditional KO line by doing, for the first time, a double transfection with the pmK-RQ-tkl2-loxPint and pDC2_TKL2_gRNA, together with the pB5P::47DiCre (containing the DiCre cassette) and the CRISPR/Cas9 plasmid pDC287 containing the guide RNA targeting the Pf47 locus, as previously described (23). The plasmids were suspended in 100 μl of P3 primary cell solution, 40 μg of each rescue plasmid, and 20 μg of pDC2-Cas9-hDHFRyFCU guide RNA for each respective rescue plasmid and transfected into the 3D7 parasites. Briefly, purified P. falciparum 3D7 schizont stages were electroporated using Amaxa 4D-Nucleofector (Lonza), program FP158 (35). Transfected parasites were selected using 5 nM WR99210 (Jacobus Pharmaceutical), and after a first round of selection, parasites were cloned.

To generate the pmK-RQ-gdv1Δ39-HA plasmid, which upon integration into the endogenous gdv1 locus mimics the mutation found in the kinase KO lines, the gdv1 (PF3D7_0935400) 3′ homology region was PCR amplified from NF54 genomic DNA with primers 268/269 (Table S2). The amplified PCR fragment was Gibson-cloned into an AfIII-digested plasmid synthesized by Geneart that contains a recodonized truncated gdv1Δ39 version and the sequence encoding the 3′-HA tag (Table S2). To generate the pD_cgdv1Δ39-gfp-glmS plasmid, we amplified the gdv1Δ39 sequence from the pmK-RQ-gdv1Δ39-HA plasmid using primers 383/384 (Table S2) and introduced the PCR fragment using Gibson assembly into the donor plasmid pD_cgdv1Δ39-gfp-glmS (29) digested with Eagl and BsaBI. The guideRNA cassette to mutate endogenous gdv1 was generated using the primer pairs pDC2_GDV1Δ39_gRNA1_FOR/pDC2_GDV1Δ39_gRNA1_REV and cloned into the pDC2-Cas9-hDHFRyFCU plasmid as previously described (22). The rescue plasmid pmK-RQ-gdv1Δ39-HA and the CRISPR/Cas9 plasmid pDC2-Cas9-hDHFRyFCU were suspended in 100 μl of P3 primary cell solution and 40 μg and 20 μg DNA, respectively, and transfected using Amaxa 4D-Nucleofector (Lonza). Briefly, purified P. falciparum NF54:DiCre schizont stages were electroporated using program FP158 (35). Selection of parasites transfected was done using 5 nM WR99210 (Jacobus Pharmaceutical), and after a first round of selection, parasites were cloned. Transfection of NF54 parasites using the CRISPR/Cas9 pH_HCgc6 suicide plasmid (29) and the pD_cgdv1Δ39-gfp-glmS donor construct was performed as described previously (12). Fifty μg each of the suicide plasmid and donor plasmid was transfected and parasites were cultured in the presence of glucosamine to block NF54/GDV1,Δ39:GFP_cOE protein overexpression. Twenty-four hours after transfection and for six subsequent days in total, the transfected populations were treated with 4 nM WR99210 and then cultured in the absence of drug selection until a stably propagating transgenic population was obtained. All primers, guide RNAs, and fragments used in the construction and integration of the constructs as well as confirmation of rapamycin-mediated excision are described in Table S2.

Plasmodium falciparum in vitro culture of asexual and sexual blood stages. Plasmodium falciparum parasite lines used in this study were all derived from the NF54 strain (originally isolated from an imported malaria case in the Netherlands in the 1980s [BEI Resources, catalog number MRA-1000]) (36). Asexual parasites were cultured in human blood (UK National Blood Transfusion Service) and RPMI 1640 medium containing 0.5% wt/vol AlbumaxII (Invitrogen) at 37°C, as previously described (37). Asexual parasites were used to produce gametocytes by seeding asexual rings at 1% or 3% parasitemia and 40 μg DNA, respectively, and transfected using Amaxa 4D-Nucleofector (Lonza). Briefly, purified P. falciparum 3D7 schizont stages were electroporated using program FP158 (35). Selection of parasites transfected was done using 5 nM WR99210 (Jacobus Pharmaceutical), and after a first round of selection, parasites were cloned. Transfection of NF54 parasites using the CRISPR/Cas9 pH_HCgc6 suicide plasmid (29) and the pD_cgdv1Δ39-gfp-glmS donor construct was performed as described previously (12). Fifty μg each of the suicide plasmid and donor plasmid was transfected and parasites were cultured in the presence of glucosamine to block NF54/GDV1,Δ39:GFP_cOE protein overexpression. Twenty-four hours after transfection and for six subsequent days in total, the transfected populations were treated with 4 nM WR99210 and then cultured in the absence of drug selection until a stably propagating transgenic population was obtained. All primers, guide RNAs, and fragments used in the construction and integration of the constructs as well as confirmation of rapamycin-mediated excision are described in Table S2.

Plasmodium falciparum sexual induction. An asexual ring stage culture (3%) was induced for sexual conversion using 50% spent medium, expecting the sexually committed merozoites to invade and develop during the next cycle (20, 37). The overexpressing NF54/GDV1:GFP_cOE and NF54/GDV1,Δ39:GFP_cOE parasite lines were kept in the constant presence of 2.5 mM glucosamine to block ectopic GDV1 expression and therefore sexual induction, while sexual induction was achieved by culturing the parasites in the absence of glucosamine, as previously described (29).
Time course of gametocyte induction, RNA extraction, and RNA-seq library preparation. The samples were collected during the asexual cycle at 28 to 32 hpi and in the matching cycle at 28 to 32 hpi after induction of sexual commitment. The infected RBCs pellets were collected at the respective time point, centrifuged and solubilized in 10 volumes of TRIzol (Ambion) premayed to 37°C. lysed for 5 min by mixing vigorously at 37°C, and immediately frozen at −80°C until extraction. Complete RNA was isolated from the samples using TRIzol/chloroform extraction followed by isopropanol precipitation, and its concentration and integrity was verified using Agilent Bioanalyzer (RNA 6000 Nano kit) and NanoDrop 1000 spectrophotometer. One to two Illumina HiSeq2500 system (100-bp paired-end reads) according to the manufacturer’s manual. All samples were generated in duplicates or triplicates, and uninduced controls were always generated and processed in parallel. Raw data are available through GEO database repository (study GSE158689).

**RNA-seq data analysis.** The generation of raw data in the form of *.fastq files quality control and adapter trimming was performed using the default analysis pipelines of the Sanger Institute. The raw data were transformed into paired *.fastq files using Samtools software (version 1.3.1). The generated reads were realigned to Plasmodium falciparum genome (PlasmoDB-38 release) in a splice aware manner with HISAT2 (40) using --known-splice-site-infile option within the splicing sites file generated based on the current genome annotation. Resulting *.bam files were sorted and indexed using Samtools and inspected visually using Integrated Genome Viewer (version 2.3.91). High-throughput sequencing (HT-seq) python library (40) was used to generate reads counts for all genes for further processing. Raw counts were normalized to median-ratio and then tested against linear models of time nested in line and line nested within time using a negative binomial model for the unnormalized counts using DESeq2, differential genes being selected for a false-discovery rate of <0.1 (41).

**Saponin lysis and whole-cell, cytoplasmic, and nuclear protein extraction.** Ten ml of parasite culture (2 to 5% parasitemia, 4% hematocrit) was transferred to a 15-ml tube and centrifuged at 600 × g for 5 min. The supernatant was aspirated and the RBC pellet was resuspended in 5 volumes of 0.15% saponin solution (2.5 ml for 500 μl RBC). After an incubation on ice of maximum 10 min, the parasites were centrifuged at 1,503 × g for 5 min at 4°C. Subsequent steps were performed on ice in order to prevent protein degradation. The supernatant was aspirated and the pellet was resuspended in 1 ml of cold phosphate-buffered saline (PBS) and transferred to an Eppendorf tube. The parasite pellet was centrifuged at 1,503 × g for 30 sec at 4°C and washed with cold PBS until the supernatant was clear. For whole-cell protein extraction, one pellet volume (30 to 50 μl) of whole-cell protein lysis buffer (8 M urea, 5% SDS, 50 mM Tris-Cl, 1 mM EDTA, 25 mM HCl [pH 6.5]) complemented with 1× protease inhibitor cocktail (Merck) and 1 mM DTT was added to the pellet at RT in order to lyse the parasites. The tube was vortexed, heated to 94°C for 5 min, sonicated for 2 min (5 cycles of 30 sec on/30 sec off), vortexed, and heated again. Subsequently, the protein sample was centrifuged at 20,238 × g for 5 min at room temperature and the supernatant was transferred into a new tube, which was frozen at −20°C and stored until use.

For cytoplasmic and nuclear protein extraction, the parasite pellet was lysed in 300 μl of cytoplasmic lysis buffer (20 mM HEPES [pH 7.9], 10 mM KCl, 1 mM EDTA, 0.65% Igepal) complemented with 1× protease inhibitor cocktail (Merck) and 1 mM DTT (leaving the nucleus intact) and incubated on ice for 5 min (43). The lysed parasites were centrifuged at 845 × g for 3 min, and the supernatant representing the cytoplasmic protein fraction was transferred into a new tube and placed on ice. The remaining nuclear pellet was washed in 500 μl of cytoplasmic lysis buffer and centrifuged at 845 × g for 3 min. The washing was repeated until the supernatant was clear. The nuclear pellet was resuspended in 60 μl of whole-cell lysis buffer and vortexed at high speed at room temperature for 10 to 20 min. The insoluble material was centrifuged at 20,238 × g for 3 min, and the supernatant representing the nuclear protein fraction was transferred to a new tube and placed on ice. Both protein fractions were frozen at −20°C and stored until use.

**Western blotting.** Parasite extracts were solubilized in protein loading buffer, denatured at 95°C for 10 min, subjected to SDS-PAGE, and transferred onto a nitrocellulose membrane. Membranes were immunostained with mouse anti-GFP (1:250 dilution; Roche, 11811466001), rabbit anti-Aldolase-horseradish peroxidase (HRP) conjugated (1:5,000 dilution; Abcam ab38905), and rabbit anti-histone 3 (1:2,000 dilution; Abcam ab1791) primary antibodies. Antibody detection was done using chemiluminescent Western blotting using goat anti-mouse secondary antibody conjugated with HRP and the ECL Western blotting detection reagents (Amersham RPN2106) or by direct infrared fluorescence detection on the Odyssey Infrared Imaging System (Odyssey CLx, LI-COR) using IRDye 800CW goat anti-rabbit IgG (1:10,000 dilution; LI-COR) and IRDye 680CW goat anti-rabbit IgG (1:10,000 dilution; LI-COR).
Immunofluorescence assay at different parasite stages. Air-dried thin blood films of asexual parasites were fixed with 4% paraformaldehyde containing 0.0075% glutaraldehyde for 15 min and permeabilized in 0.1% (vol/vol) Triton X-100 (Sigma) for 10 min (42). Blocking was performed in 3% bovine serum albumin (BSA) for 1 h. Slides were incubated with rat anti-HA high-affinity (1:1,000 dilution; Roche, clone 3F10) at room temperature for 30 min, followed by Alexa fluor conjugated goat anti-rat IgG (1:1,000 dilution; Thermo Fisher Scientific) at room temperature for 30 min. Parasite nuclei were stained with 4',6-diamino-2-phenylindole (DAPI; Invitrogen). Slides were mounted in ProLong Gold antifade reagent (Invitrogen) and images were obtained with the inverted fluorescence microscope (Ti-E; Nikon, Japan) and processed using NIS-Elements software (Nikon, Japan).

Flow cytometry. NF54/GDV1:GFP_cOE and NF54/GDV1 Δ39:GFP_cOE parasites were grown in the presence or absence of glucosamine in order to block or allow sexual commitment, respectively. Schizonts were purified by Percoll gradient and allowed to invade fresh red blood cells for 4 h before uninvaded schizonts were removed. Flow cytometry analysis was performed at approximately 44 h post invasion, in 4 biological replicates. For one replicate, parasites were fixed for 1 h in 4% paraformaldehyde in PBS, stained with Hoechst 33342 (1:1,000 in PBS) for 10 min, and analyzed on an LSRFortessa flow cytometer (Becton, Dickinson) using FACSDivia software. For the other three replicates, live parasites were stained with Hoechst 33342 and analyzed on a BD FACSaria II flow cytometer (Becton, Dickinson) using FACSDivia software. Hoechst fluorescence was detected using a 355 nm (UV) excitation laser with a 450/50 nm bandpass filter, while GFP fluorescence was detected with a 488 nm (blue) excitation laser, a 505 nm longpass filter, and a 530/30 nm bandpass filter. At least 30,000 cells were counted for each sample. Data were analyzed using FCS Express 7 (Research Edition) software. The population was first gated on single cells based on the side and forward scatter, then on highly Hoechst-positive infected schizonts, before the median fluorescence intensity (MFI) of the GFP fluorescence was calculated for each line. An example of the gating strategy for infected cells is shown in Fig. S4. Due to the variation in fluorescence intensity between different experiments, MFI values were normalized by dividing the MFI of each infected sample by the average MFI of the uninfected samples within the same experiment (n = 4). Statistical analysis was performed using Holm-Sidak corrected multiple comparison analysis of variance (ANOVA) on samples paired within each experiment using GraphPad Prism version 8.

Reverse transcription quantitative PCR. NF54/GDV1:GFP_cOE and NF54/GDV1 Δ39:GFP_cOE parasites were synchronized to a 4-h window and grown in the presence or absence of glucosamine in order to block or allow sexual commitment, respectively. Schizonts from the 3 independent biological replicates for fluorescence-activated cell sorter (FACS) analysis were harvested at 40 to 44 hpi and used for RNA extraction. The infected RBCs pellets were solubilized in 10 volumes of TRizol (Ambion) prewarmed to 37°C, lysed for 5 min by mixing vigorously at 37°C, and immediately frozen at −80°C until extraction. Complete RNA was isolated from the samples using TRizol/chloroform extraction purified using the RNeasy Plus minikit (Qiagen). Residual gDNA was digested with TURBO DNA-free DNase I (Ambion). Three to five μg of total RNA from each sample was reverse-transcribed using the SuperScript III first-strand synthesis system (Invitrogen). qPCRs were performed using KAPA SYBR Fast ROX Low kit (Sigma-Aldrich) in a reaction volume of 20 μl. All reactions were run in technical triplicate. Cycling conditions were 95°C for 3 min, followed by 40 cycles of 95°C/3 sec and 60°C/40 sec. Product-specific amplification was ensured by melting curve analysis for each reaction. Relative transcript levels were calculated by normalizing against the housekeeping gene encoding eukaryotic translation initiation factor 2-alpha kinase (pk4, PF3D7_0628200). For gDNA generation, compound 2 arrested schizonts where harvested and genomic DNA was extracted using the Qiagen blood and tissue kit, according to the manufacturer's recommendation. All primer sequences are listed in Table S2.

In vitro protein-protein interaction experiments. In order to coexpress StreptⅢ II-tagged HP1 with a His-SUMO-tagged truncated version of GDV1, we deleted the 39 C-terminal amino acids of the coding sequence of GDV1 in the vector pStrep-HP1_H5-GDV1 (12). For this purpose, we circularized a PCR product amplified from this vector with the primers D39F and D39R using Gibson assembly. The proteins were expressed and the in vitro interaction assay was performed as previously described (14) using full-length GDV1 as the positive and SIP2 as the negative control.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

FIG S1, JGP file, 0.2 MB.
FIG S2, JGP file, 0.4 MB.
FIG S3, JGP file, 1 MB.
FIG S4, PDF file, 1.2 MB.
TABLE S1, XLSX file, 1.9 MB.
TABLE S2, DOCX file, 0.02 MB.

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M. Tibúrcio and M. Treeck conceived the study. M. Tibúrcio performed most of the parasite genetic manipulations and all the parasite line phenotyping experiments, as well as RNA-seq material collection. E. Hitz generated the NF54/GDV1Δ39:GFP_cOE parasite line. I. Niederwieser performed the in vitro protein-protein interaction experiments, and T.S. Voss supervised these experiments and provided conceptual advice and resources. Gavin Kelly performed RNA-seq analysis. RNA-seq samples were run in O. Billker’s group at the Sanger Institute. H. Davies performed the flow cytometry data analysis. C. Doerig provided the original P. falciparum kinase KO cell lines. All authors contributed to experimental design and interpretation of the results. M. Tibúrcio and M. Treeck wrote the article with contributions from all authors.

We declare that we have no competing interests.

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