The *Plasmodium berghei* serine protease PbSUB1 plays an important role in male gamete egress

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**Abstract**
The *Plasmodium* subtilisin-like serine protease SUB1 is expressed in hepatic and both asexual and sexual blood parasite stages. SUB1 is required for egress of invasive forms of the parasite from both erythrocytes and hepatocytes, but its subcellular localisation, function, and potential substrates in the sexual stages are unknown. Here, we have characterised the expression profile and subcellular localisation of SUB1 in *Plasmodium berghei* sexual stages. We show that the protease is selectively expressed in mature male gametocytes and localises to secretory organelles known to be involved in gamete egress, called male osmiophilic bodies. We have investigated PbSUB1 function in the sexual stages by generating *P. berghei* transgenic lines deficient in PbSUB1 expression or enzyme activity in gametocytes. Our results demonstrate that PbSUB1 plays a role in male gamete egress. We also show for the first time that the PbSUB1 substrate PbSERA3 is expressed in gametocytes and processed by PbSUB1 upon gametocyte activation. Taken together, our results strongly suggest that PbSUB1 is not only a promising drug target for asexual stages but could also be an attractive malaria transmission-blocking target.

**KEYWORDS**
egress, gametocytes, malaria, Plasmodium berghei, SERA3, SUB1

**1 | INTRODUCTION**

Malaria is a devastating disease with 216 million cases globally and 445,000 lethal outcomes in 2016 (Organization WH, 2017), mainly in children under the age of 5. The disease is caused by intracellular parasites of the *Plasmodium* genus, transmitted by the bite of anopheline mosquitoes. After an initial reproductive cycle in the liver, malaria parasites invade erythrocytes, where they either multiply asexually by forming schizonts or differentiate into gamete precursors called gametocytes. Asexual blood stages are responsible for the clinical manifestations of the disease, whereas gametocytes are essential for parasite transmission. When mature gametocytes are ingested by mosquitoes during a blood meal, they differentiate into gametes and egress from the host red blood cell by an inside-out mode of egress (Sologub et al., 2011). Each female gametocyte forms a single macrogamete, whereas male gametocytes undergo a drastic transformation, known as exflagellation, by forming eight flagellar microgametes. Male gametes contact surrounding erythrocytes, forming so-called “exflagellation centres,” in which fertilisation of the female takes place. Gamete egress follows a fixed programme, requiring discharge of
secretory organelles called osmiophilic bodies (OBs) followed by inside-out disruption of the parasitophorous vacuole membrane (PVM) and erythrocyte membrane (Ponzi et al., 2009; Sologub et al., 2011; Talman et al., 2011). This entire process is crucial for fertilisation and zygote formation and represents a major bottleneck in the Plasmodium life cycle, involving an approximate 300-fold loss of parasite abundance (Kuehn & Pradel, 2010).

In male Plasmodium berghei gametocytes, OBs are club-shaped, electron-dense vesicles that are smaller than the oval-shaped female OBs (Olivieri et al., 2015). Because of their gender-specific morphological features, these vesicles are named male osmiophilic bodies (MOBs; Olivieri et al., 2015). Upon gametocyte activation, OBs accumulate at the gametocyte plasma membrane in multiple foci prior to secreting their contents into the parasitophorous vacuole (PV). In contrast, MOBs cluster together to form larger structures and then release their contents at only a few focal points (Olivieri et al., 2015). These different behaviours likely reflect specific mechanisms of vesicle discharge. A specific MOB marker was recently described in Plasmodium yoelii (Tachibana, Ishino, Takashima, Tsuboi, & Torii, 2018).

The threat of emerging malaria strains resistant to currently available drugs has made the search for novel drug targets compelling. In particular, the scientific community has recently acknowledged an urgent need for transmission-blocking drugs targeting the sexual stages. Drugs effective against gametocytes/gametes, used in combination with drugs against asexual stages, would have the dual advantage of lowering parasite transmission and preventing the spread of drug-resistant parasites. The subtilisin-like protease SUB1 is a Plasmodium serine protease that is expressed both in the liver and blood schizonts and then secreted into the PV from vesicular organelles called exonemes just prior to schizont rupture (Yeoh et al., 2007). The proteolytic activity of SUB1 has been shown to be important for maturation and egress of invasive merozoites from both infected erythrocytes (Thomas et al., 2018; Yeoh et al., 2007) and hepatocytes (Tawk et al., 2013). During Plasmodium falciparum merozoite egress, PfSUB1 carries out the maturation of a family of papain-like proteins called the serine repeat antigens (SERAs), whereas for merozoite maturation, the role of SUB1 is to undertake the processing of several merozoite surface proteins (Yeoh et al., 2007). Orthologs of sub1 are found in every Plasmodium species examined, suggesting a conserved functional role.

In P. falciparum, maturation of PfSUB1 has been described in detail. The protease is synthesised as a preprotein that undergoes maturation by proteolytic processing in two consecutive steps: first, in an autocatalytic conversion, to a 54-kDa form called p54, by the release of a prodomain (Sajid, Withers-Martinez, & Blackman, 2000), and finally, by means of the P. falciparum protease plasmaspin X, to a terminal 47-kDa species called p47 (Nasamu et al., 2017; Pino et al., 2017). In P. berghei, a similar processing pattern has been shown, with a ~70-kDa fragment likely corresponding to full-length PbSUB1 and a doublet at 44–47 kDa likely corresponding to the two processed forms of PbSUB1 (Suarez, Volkmann, Gomes, Billker, & Blackman, 2013). The subtilisin prodomain functions as an intramolecular chaperone, being essential for correct folding of the catalytic domain and for enzyme maturation (Jean, Hackett, Martin, & Blackman, 2003). Like many subtilisin prodomains, the SUB1 prodomain is a potent and highly selective inhibitor of SUB1 protease activity: it possesses inhibition constants in the nanomolar range and does not inhibit other subtilisin-like serine proteases (Jean et al., 2003).

It was recently shown that PbSUB1 is also expressed in P. berghei sexual stages (Pino et al., 2017), but the function of the enzyme in gametocytes/gametes and its potential substrates in these stages are still unknown. Here, we show that PbSUB1 is selectively expressed in male P. berghei gametocytes and that it localises to MOBs (Olivieri et al., 2015). Using conditional depletion of PbSUB1 expression or enzyme activity, we show that the protease plays a role in male gamete egress. Our results strongly suggest that PbSUB1 is not only a promising drug target for asexual stages but could also be an attractive malaria transmission-blocking target.

## RESULTS

### 2.1 PbSUB1 is expressed in both asexual and sexual stage parasites and localises to MOBs in P. berghei gametocytes

To investigate the regulation of sub1 gene expression in gametocytes, we generated P. berghei transgenic lines in which DNA sequences upstream of the psub1 start codon were cloned upstream of a reporter gene (green fluorescent protein [GFP]) and integrated into a silent locus (Figure S1), previously successfully used to integrate constructs into the P. berghei genome (Kooij, Rauch, & Matuschewski, 2012). The first two constructs, named SUB1/prom1 and SUB1/prom2, included the DNA sequence spanning the region between 1,279 bp and either 25 or 450 bp upstream of the psub1 ATG, respectively, including or excluding an intron present in the psub1 5’ UTR (Figure 1a). The resulting transgenic parasite lines were both found to specifically display reporter gene expression in the cytoplasm of gametocytes, with no detectable expression in asexual blood stage parasites (SUB1/prom2 expression is shown in Figure 1b, as an example). We then generated a third construct, where a larger DNA segment, extending from 2,385 to 450 bp upstream of the psub1 ATG was fused to gfp. In the obtained transgenic line, named SUB1/prom3, the reporter gene was expressed both in gametocytes and schizonts (Figure 1b). These results suggested that the psub1 gene includes two distinct regulatory regions: one, closest to the ATG, that specifically drives gene expression in the sexual stages and another, further upstream, necessary to drive expression also in asexual stages. Interestingly, in all the SUB1/prom lines, the GFP fluorescence signal significantly increased in activated male gametocytes (Figure 1b), indicating up-regulation of psub1 gene expression during male gamete formation.

We next investigated the subcellular localisation of PbSUB1 in mature P. berghei gametocytes by immunofluorescence (IFA) using an anti-PbSUB1 antibody (Suarez et al., 2013). Antibody specificity was
confirmed in gametocytes by double IFA with an anti-HA monoclonal antibody on a $P. berghei$ line in which the endogenous $psub1$ gene had been modified with a triple HA tag at the C-terminal end (Suarez et al., 2013). The two signals colocalised in gametocytes, producing a punctate pattern (Figure S2). Double IFA with antibodies to the nuclear protein SE translocation proto-oncogene like, which is highly abundant in male gametocytes (Pace et al., 2006) and with the female gametocyte marker G377 (Olivieri et al., 2015), showed that $PbSUB1$ expression in gametocytes is male specific (Figure 2a). Of 100 $SUB1$-positive gametocytes examined, none showed a fluorescence signal with the G377 antibody. Colocalisation experiments with an antibody against the male development protein 1 (MDV1), a gametocyte-specific protein associated with OBs in both $P. berghei$ male and female gametocytes (Ponzi et al., 2009), showed that $PbSUB1$ specifically localises to these secretory organelles in males (MOBs; Figure 2b) with a Pearson’s correlation coefficient of .94 ($P < .001$). Further examination showed that $PbSUB1$ follows the typical MOB pattern, being released upon gametocyte activation and not detectable in mature male gametes (Figure 2b).

2.2 | $PbSUB1$ enzyme activity is required for male gamete egress

To investigate whether the proteolytic activity of $PbSUB1$ is essential for gamete egress, we took advantage of the capacity of the $SUB1$ prodomain to inhibit with high specificity the proteolytic activity of its cognate catalytic domain. To do this, we produced a transgenic $P. berghei$ line, named $SUB1/prod$, expressing an HA-tagged extra copy of the prodomain fused to the 5’ flanking region and the first 90 residues of the $mdv1$ gene, in order to express the chimera in gametocytes and target the prodomain to OBs (Figure S3a). After confirming the integration event in clones of the $SUB1/prod$ line (Figure S3b), the expression profile of the HA-tagged $PbSUB1$ prodomain chimera was analysed by IFA and western blot (Figure S4). This showed that the transgene was specifically expressed in gametocytes, but the HA-tagged prodomain did not colocalise with $PbSUB1$, suggesting that additional signals are required to traffic the chimera to MOBs (Figure 3). In male gametocytes, the transgenic prodomain showed a peripheral distribution with thickening of the signal in some areas around the parasite (Figure S4).

To assess whether the endogenous $PbSUB1$ proteolytic activity was inhibited as anticipated in the $SUB1/prod$ line, we preliminarily investigated whether proteolytic processing of the PV protein $PbSERA3$, known to be a substrate of $PbSUB1$ during egress of merozoites from $P. berghei$ liver schizonts (Tawk et al., 2013), also occurred in the sexual stages. The expression profile and subcellular localisation of $PbSERA3$ has not previously been investigated in $P. berghei$ blood stages, although its orthologue in $P. falciparum$, PfSERA6, undergoes maturation by PfSUB1 in asexual blood stages (Ruecker et al., 2012; Thomas et al., 2018). As shown in Figure 3b, SERA3 is expressed in $P. berghei$ blood schizonts, showing a peripheral localisation. It was also found to be expressed in gametocytes, displaying a diffuse signal, stronger in the cell periphery in both sexes; in addition, female gametocytes also showed a punctate pattern in the cytoplasm.

To investigate proteolytic processing of $PfSERA3$ by $PbSUB1$, noninduced gametocytes and supernatants from purified activated gametocytes were collected from the transgenic $SUB1/prod$ and its
parental parasite line and analysed by western blot (Figure 3c). Nonactivated gametocytes from both the wt and transgenic lines showed the same processing pattern, consisting of the PbSERA3-130 precursor and the intermediate maturation products PbSERA3-110, PbSERA3-100, PbSERA3-72, and PbSERA3-55 (Ruecker et al., 2012; Schmidt-Christensen, Sturm, Horstmann, & Heussler, 2008). In supernatants of exflagellating wt gametocytes, two further processing products were visible, at around 63 and 48 kDa, respectively, the latter being the final maturation product (Ruecker et al., 2012). The 63-kDa band, not previously described, could be a processing product specific to sexual stages. The 55-kDa band, unexpectedly disappearing from the SUB1/prod exflagellation supernatants, could remain partially associated to PVM, as previously reported for the SERA3 orthologue in *P. falciparum*, PfSERA6 (Ruecker et al., 2012). Both the 63- and 48-kDa bands were prominent in supernatants from the wt line but were barely detectable in those of the activated SUB1/prod line; in contrast, the full-length protein was undetectable in supernatants from the wt parasites, whereas it was conspicuous in supernatants of the SUB1/prod line. This suggested that (a) the two terminal PbSERA3 proteolytic processing events occur during gamete egress; (b) these processing events are PbSUB1 dependent in *P. berghei* sexual stages; and (c) PbSERA3 processing is significantly reduced in SUB1/prod gametocytes, demonstrating that in this line, PbSUB1 activity is strongly impaired. Although PbSUB1 is only expressed in male gametocytes, its substrate PbSERA3 is expressed in both genders. There may be several explanations for this, including the fact that expression of the protein in female gametocytes may be redundant or that its function in females does not require proteolytic maturation. Having confirmed inhibition of endogenous PbSUB1 activity in the SUB1/prod line, the efficiency of male gamete formation was evaluated in this line by counting exflagellation centres in vivo. As shown in Figure 3d, male gametes of the SUB1/prod line showed a significant reduction in exflagellation rates, compared with wt gametes, indicating that PbSUB1 enzymatic activity plays an important role in male gamete formation.

To confirm PbSUB1 function in gametocytes, we generated a second transgenic *P. berghei* line, named SUB1/asex, in which PbSUB1
expression was selectively abolished in gametocytes. To do this, the endogenous pbsub1 promoter was substituted with the promoter of the apical membrane antigen 1 (ama1; Figure S5a). The transgenic line was cloned and positive clones were identified by diagnostic PCR analysis (Figure S5b). AMA1 is expressed at approximately the same developmental stage as PbSUB1 in mature asexual stages but is not expressed in gametocytes (Kocken et al., 1998). The SUB1/asex line was characterised by IFA and western blot analysis. In both cases, PbSUB1 was found to be expressed in transgenic schizonts in a similar manner to wt parasites, whereas it was not detected in SUB1/asex gametocytes (Figure 4a,b). In addition, the proportion of nonreleased male gametes still possessing an intact PVM was significantly higher in the SUB1/asex line than in the wt parasites (Figure 4e). Taken together, these results indicate a defect in PVM rupture of SUB1/asex male gametes and are consistent with the previously described role of

FIGURE 3 Transgenic expression of the PbSUB1 prodomain reduces endogenous PbSUB1 protease activity and inhibits exflagellation. (a) IFA of the transgenic line SUB1/prod with an anti-HA antibody. Anti-PbSUB1 was used as an OB marker. Nuclei are stained with DAPI. Scale bar 5 μm. (b) IFA of Plasmodium berghei blood stages with anti-SERA3. Anti-Sep1 antibody was used as a PVM marker; anti-SET antibody was used as a gender marker; nuclei are stained with DAPI. Scale bar 5 μm. (c) Western blot analysis with anti-PbSERA3 of extracts from 5 × 10⁶ wt gametocytes (gams) and 5 × 10⁶ SUB1/prod gametocytes, wt exflagellation supernatants (exfl. sup.) from 5 × 10⁶ gametocytes, and SUB1/prod exflagellation supernatants from 5 × 10⁶ gametocytes. Anti-SUB1 was used as a loading control. Asterisks highlight the seven main processing products: PbSERA3-130, PbSERA3-110, PbSERA3-100, PbSERA3-72, PbSERA3-55, PbSERA3-65, and PbSERA3-48. Red asterisks highlight the full-length protein and the terminal processing product PbSERA3-48. This experiment was conducted in two biological replicates, both showing the same processing pattern in gametocytes and exflagellation supernatants. (d) Exflagellation rates of the SUB1/prod line in two independent experiments, each performed in triplicate, at 15 min postinduction, expressed as a percentage of exflagellation rates in the wt line (Student’s t test: pexp1 < 0.005 and pexp2 < 1.02 × 10⁻⁵; error bars represent +/− standard deviation of the mean value)
SUB1 in schizont PVM rupture (Thomas et al., 2018). Because the AMA1 promoter is active in the mosquito stages (Schmidt‐Christensen et al., 2008), phenotypic analysis of the SUB1/asex transgenic line could not be taken further.

3 | DISCUSSION

In this study, we analysed the expression profile and function of the subtilisin‐like serine protease PbSUB1 in P. berghei sexual stages. We first examined sub1 gene expression and showed that the sub1 promoter is bipartite, with one region driving gene expression in the sexual stages, with a peak in activated male gametocytes, and another segment necessary to drive expression also in asexual stages. We then investigated the subcellular localisation of the protease in the sexual stages, showing that PbSUB1 is expressed specifically in male gametocytes and localises to specialised secretory organelles called MOBs, previously shown to be involved in the exflagellation process (Olivieri et al., 2015; Tachibana et al., 2018). We have also demonstrated here that the enzymatic activity of PbSUB1 is important for male gamete egress from the host cell. In P. berghei, three proteins localising to OBs have been previously reported to play a role in gamete emergence: P. berghei gametocyte 377 protein (PbG377; de Koning‐Ward et al., 2008; Olivieri et al., 2015), gamete egress and sporozoite traversal protein (PbGEST; Talman et al., 2011), and MDV1 (Ponzi et al., 2009). PbG377 is expressed only in female gametocytes and plays a role in OB biogenesis (Alano et al., 1995; Severini et al., 1999): its absence causes a dramatic reduction in OB size, as well as a slight delay in female gametocyte egress (de Koning‐Ward et al., 2008; Olivieri et al., 2015). PbGEST and MDV1 are expressed in both genders and localise to OBs/MOBs (Ponzi et al., 2009; Talman et al., 2011). Genetic deletion of PbGEST and MDV1 strongly affects gamete egress from the host cell. In these parasites, gametocytes of both genders fully develop inside RBCs but remain entrapped within the PVM after activation of gametogenesis. In parasites lacking PbSUB1 in gametocytes (SUB1/asex transgenic line), we observed that at 20 min postactivation, around 40% of SUB1/asex male gametes still had an intact PVM, compared with only 2% in the wt line. This is
consistent both with the described function of SUB1 in asexual stages, being a protease essential for schizont PVM rupture (Thomas et al., 2018), and with the role of OB/MOBs in gamete egress. Interestingly, PbSUB1 is the only OB/MOB resident protein described to be expressed also in asexual stages, where it localises to exosomes, secretory vesicles involved in parasite egress (Yeoh et al., 2007). This suggests that MOBs and exosomes may belong to the same category of vesicles and possibly display some overlap in their function and protein content and share the same localisation signals. In conclusion, we described a novel specific marker of MOBs and showed that it plays a role in gamete egress.

We also showed here for the first time that the PbSUB1 substrate PbSERA3 is expressed in gametocytes and processed by PbSUB1 upon gametocyte activation. SERA3 shows a localisation compatible with the PV both in schizonts and male gametocytes, consistent with the localisation of its P. falciparum orthologue (Ruecker et al., 2012; Suarez et al., 2013; Tawk et al., 2013), and a punctate cytoplasmic pattern in female gametocytes. In nonactivated gametocytes, SERA3 is mostly present in the unprocessed form of 130 kDa and in the partial maturation product of 55 kDa, whereas in supernatants from activated gametocytes, the most abundant band is the final maturation product of 48 kDa, thus demonstrating that PbSUB1 only processes its substrate PbSERA3 when it is discharged into the PV.

SUB1 is already considered a promising drug target for malaria chemotherapy, being essential in both hepatic and asexual blood stages of the Plasmodium life cycle. Several P. falciparum SUB1-specific inhibitors have been developed (Gemma et al., 2012; Giovani et al., 2014; Kher et al., 2014) and in some cases were shown to be effective in inhibiting parasite emergence from erythrocytes, indicating that SUB1 is a druggable target. In addition, the P. falciparum PfSUB1 and Plasmodium vivax PvSUB1 x-ray crystal structures were recently made available (Giganti et al., 2014; Withers-Martinez et al., 2014). Because no structurally similar proteases have been identified in the human genome, the development of highly selective SUB1 inhibitors appears feasible. Here, we have demonstrated that PbSUB1 plays an important role in disease transmission, making the protease an even more interesting therapeutic candidate, given the current focus on malarial molecular targets suitable for transmission-blocking drug development. Exploring the possible presence of this protease in the sexual stages of P. falciparum and investigating its role in these stages would be therefore a very promising research topic.

4 EXPERIMENTAL PROCEDURES

4.1 DNA construct generation

To produce the SUB1/prom constructs, the putative psub1 3' UTR, spanning from -36 to +766 bp from the stop codon, was amplified by genomic DNA with the primer pair SUB1-3'UTR_for and SUB1-3'UTR_rev (Table S1), digested with Ncol/SacII and cloned in the plasmid pBAT-GFP (Kooij et al., 2012), kindly provided by Taco Kooij, digested with the same enzymes. Two putative psub1 promoter regions, respectively named SUB1-prom1 and SUB1-prom2, respectively, of 830 bp (from -1,279 to -450 bp from ATG) and 1,956 bp (from -2,385 to -450 bp from ATG, partially overlapping the EST of gene located upstream of sub1) were amplified by genomic DNA. The amplification products were respectively digested with SpeI/Ndel and SacII/NotI and cloned into the pBAT-GFP, containing the putative psub1 3'UTR, digested with the same enzymes. The obtained constructs were respectively named pSUB1-prom1 and pSUB1-prom2.

In order to produce the Sub1/prod construct, MDV1 N-terminal 90 aa were amplified by genomic DNA with the primer pairs MDV1 and digested with EcoRI/Narl, and PbSUB1 prodomain was amplified with the primer pair SUB1-prod and digested with Narl/BssHII. Both PCR fragments were simultaneously cloned into a plasmid derived from the plasmid pDEF-SSU-hDHFR alias pL0008, a plasmid containing sequences for integration in the ribosomal DNA (kindly provided by Prof. C.J. Janse) digested with EcoRI/BssHII. The coding regions were sequenced with the primers prod_seq1 to prod_seq5 (Table S1).

MDV1 promoter region, spanning from -923 to -133 bp from the ATG, was then amplified with the primer pair MDV1-pr, digested with EcoRI/PstI, and cloned into the resulting plasmid, also digested with the same enzymes.

To produce the Sub1/asex construct, a gene targeting construct was made by modifying the plasmid pL1313 designed for targeted gene disruption by double crossover homologous recombination (Laurentino et al., 2011). A 956 bp 5'-targeting region and a 1,006 bp 3' targeting region were amplified from P. berghei genomic DNA using primer pairs sub1 swap and sub1 swap R (Table S1) and cloned into the pL1313 plasmid digested with KpnI/HindIII and NcoI/NotI, respectively. The 5'-targeting region includes a DNA sequence upstream of psub1 start codon (-1,420 bp to -477), whereas the 3' targeting region includes part of the coding sequence (-107 to +881 bp from psub1 start codon). The coding region was sequenced with the primers sub1 seq1 and sub1 seq2 (Table S1) to confirm that no mutations were introduced during the amplification process.

The AMA1 promoter, amplified by genomic DNA with the primer pairs sub1 swap prAMA1, was then cloned into the resulting plasmid digested with EcoRV/BamHI. This promoter region had previously been used and characterised in P. falciparum (Olivieri et al., 2011).

4.2 Transgenic parasite production and cloning

P. berghei ANKA high gametocyte producer cloned line (8417HP) was maintained in Swiss mice. Synchronous infections were established in CD1 mice by intravenous injection of purified infective schizonts (Janse & Waters, 1995). Blood was collected by heart puncture under anaesthesia and leukocytes were removed using Plasmodipur leukocyte filters (Euro-Dagnostica). Schizont- or gametocyte-infected erythrocytes were separated from uninfected cells through 14.3% Nycodenz density gradient centrifugation (Janse & Waters, 1995).

Transfection and selection of transformed parasites was performed using standard genetic modification technologies for P. berghei (Janse
et al., 2006) using P. berghei HP as the parent parasite line. Cloned parasite lines were obtained by limiting dilution.

Plasmid integration in the transgenic lines was confirmed by diagnostic PCR reactions with the primers prAMA1-int-for and SUB1_seq2 for the SUB1/asex line and diag-mdv-for diag-mdv-rev (Table S1).

### 4.3 In vivo evaluation of GFP fluorescent parasites

Parasites from the SUB1/prom1 and SUB1/prom2 lines were synchronised by intravenous injection of purified infective schizonts and analysed at the fluorescence microscope to investigate GFP expression profile. Nuclei were stained with Hoechst. GFP fluorescence was analysed in synchronous schizonts at different time points (20, 21, and 22 hr postinvasion) and blood smears were produced at each time point to confirm by IFA with anti-PbSUB1 that the sample included PbSUB1-expressing schizonts. In order to investigate GFP expression in the sexual stages, synchronous gametocytes at 29 hpi were collected and analysed at the fluorescence microscope. The transgenic lines SUB1/prom1 and SUB1/prom2 showed GFP fluorescence with similar intensity. Scale bar 5 μm.

### 4.4 In vivo evaluation of male gamete egress

Transgenic and wt parasites were induced to form gametes in RPMI pH 8.0 supplemented with 50-μM xanthurenic acid and kept at 20°C. Male gamete formation was measured by counting exflagellation centres at the optical microscope. Blood collected from the mouse tail was diluted 1:25 and counted in Neubauer chamber in a total volume 0.1 μl roughly corresponding to 24,000 erythrocytes. Exflagellation centre counts were normalised on the number of male gametocytes per 24,000 erythrocytes in each sample analysed, quantified on Giemsa-stained blood smears with an optical microscope.

### 4.5 IFA and western blot

For IFA, air-dried thin blood films were fixed in 4% paraformaldehyde for 30 min, permeabilised for 10 min with 0.1% Triton X100, and then blocked for 1 hr with 3% (w/v) bovine serum albumin. Samples were then probed for 1 hr with the HA-specific mAb 3F10 (Roche) diluted 1:200; and/or a rabbit polyclonal antiserum raised against PbSUB1 (Suarez et al., 2013) diluted 1:100; and/or a rabbit polyclonal antiserum raised against the PV marker Sep1 (Birago et al., 2003) diluted 1:100; and/or a mouse polyclonal antiserum raised against the gender marker SE translocation diluted 1:100; and/or a mouse polyclonal antiserum raised against the OB-marker MDV1 diluted 1:500; and/or a monoclonal mouse anti-Tubulin antibody (Sigma) diluted 1:400; and/or a mouse polyclonal antiserum raised against PbSERA3 diluted 1:50 (Suarez et al., 2013). Samples were then washed for 5 min in PBS before incubation with an FITC- or Rhodamine-conjugated antimouse and/or antirabbit IgG (Thermo Scientific) diluted 1:400. Nuclei were stained with 4′,6-diamidino-2-phenylindole and samples were mounted in Vectashield (Vector Laboratories). Pearson’s correlation coefficients were calculated using the Coloc2 plugin of Fiji (Schindelin et al., 2012).

The amount of nonreleased male gametes was calculated by dividing the amount of circular flagella surrounded by a visible membrane in bright field by the sum of circular flagella and free flagella divided by 8, the number of gametes produced by a single male gametocyte.

For western blot analysis, culture supernatants were centrifuged at 16,000 g, passed through 22-μm filters, acetone precipitated, and suspended in SDS sample buffer, whereas parasite pellets were solubilised in SDS sample buffer. All samples were then subjected to SDS-PAGE under reducing conditions, followed by transfer to nitrocellulose membrane (Sartorius). Antibodies used: rabbit polyclonal antiserum raised against PfSUB1 diluted 1:500, rabbit polyclonal antiserum raised against PbSub1 diluted 1:5,000, mouse polyclonal antiserum raised against PbSERA3 diluted 1:1,000 (Suarez et al., 2013), monoclonal mouse anti-HA diluted 1:2,000 (Roche), mouse polyclonal antiserum raised against MDV1 diluted 1:2,000, and mouse polyclonal antiserum raised against14-3-3 diluted 1:1,000 (Lalle et al., 2011). The filters were then incubated with antimouse or antirabbit HRP-conjugated antibodies (Pierce) and the immunocomplexes were visualised using chemiluminescence ECL detection system (Luminata Forte Western HRP Substrate, Millipore).

### 4.6 Ethics statement

The protocol for infecting mice with P. berghei was approved and carried out at the University of Perugia under the Italian Ministry of Health Licence 1192/2016-PR awarded in 2016, under the guidelines D.lgs. n. 26/2014 that implements the directive 86/609/EEC from the European Union.

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