The cyclic nucleotides 3',5'-cyclic adenosine monophosphate (cAMP) and 3',5'-cyclic guanosine monophosphate (cGMP) are intracellular messengers found in most animal cell types. They usually mediate an extracellular stimulus to drive a change in cell function through activation of their respective cyclic nucleotide-dependent protein kinases, PKA and PKG. The enzymatic components of the malaria parasite cyclic nucleotide signalling pathways have been identified, and the genetic and biochemical studies of these enzymes carried out to date are reviewed herein. What has become very clear is that cyclic nucleotides play vital roles in controlling every stage of the complex malaria parasite life cycle. Our understanding of the involvement of cyclic nucleotide signalling in orchestrating the complex biology of malaria parasites is still in its infancy, but the recent advances in our genetic tools and the increasing interest in signalling will deliver more rapid progress in the coming years.

1. Introduction

Malaria parasites are fascinating organisms to study; research efforts, however, are primarily motivated by their devastating impact on humanity each year. Although the numbers of deaths and clinical cases have fallen dramatically over the last 15 years, mainly due to successful mosquito control programmes and effective drug treatments, there are still around 200 million cases of malaria resulting in over 400,000 deaths each year. Around 70% of these deaths are in children under 5 years old in Africa [1] infected with Plasmodium falciparum who have not built up sufficient natural immunity. Worryingly, the spread of insecticide-resistant mosquito vectors and drug-resistant parasites threatens to reverse the promising recent trend. The rate at which these organisms develop resistance vastly exceeds our ability to develop and bring new interventions to the field. There are a small number of very promising new antimalarial drugs that are undergoing clinical testing, some of which have been identified by phenotypic screening of compound libraries [2]. However, the attrition rate in drug discovery is very high and so the pipeline needs to remain productive. Trends in Pharma drug discovery seem to be shifting back towards target-based approaches, which emphasizes the importance of understanding the biology of malaria parasites at the molecular level and the identification of essential biochemical pathways that can be targeted with novel, small molecule inhibitors.

The malaria parasite life cycle is complex, with extended periods in the human host and also in the Anopheles mosquito vector which transmits disease between people (figure 1). When an infected female mosquito bites a person, sporozoites are injected into the bloodstream which travel to the liver where they invade hepatocytes. Here they divide asexually and eventually the hepatocyte ruptures to liberate thousands of merozoites from each infected cell. These merozoites must then invade red blood cells where they again replicate...
asexually by a process called schizogony whereby multiple daughter cells develop within a vacuole inside the infected erythrocyte. This blood stage cycle is responsible for all the pathology associated with malaria and, in the case of \textit{P. falciparum}, takes 48 h. The synchronous rupture of mature schizonts to release the invasive merozoites causes the characteristic regularly spaced fevers associated with this infection. A subpopulation of the asexual blood stage parasites develops into male and female sexual precursors, called gametocytes, which mature in around 10 days. To continue the life cycle, gametocytes must then be taken up by a feeding mosquito where, following activation, they round up and gametes emerge from their host red blood cells. Male gametogenesis involves a visually and metabolically spectacular process known as exflagellation. Here, eight highly motile, flagellated gametes are generated from a single haploid male gametocyte within around 10 min of activation, allowing them to fertilize female gametes in the insect midgut. The resulting zygote develops into a motile ookinete which burrows through the mosquito midgut epithelium to form an oocyst. Asexual replication produces thousands of sporozoites, which when mature are released from the oocyst and migrate to the mosquito salivary glands to continue the cycle. It is clear that this complex life cycle must be tightly regulated to allow appropriate development and survival of all these highly specialized parasite forms in the varied environments they encounter. This has led researchers to investigate the signalling pathways that might be responsible for coordinating the timely progression of this intricate life cycle. This review will focus on current knowledge of the role of cyclic nucleotide signalling in regulating malaria parasite development.

Figure 1. Stage-specific involvement of cyclic nucleotide signalling components and calcium-dependent protein kinases. A schematic depicting all the key stages of the complex \textit{P. falciparum} life cycle. The inner circles depict which life-cycle stages have been associated with specific cyclic nucleotide signalling components and calcium-dependent protein kinases.
Cyclic nucleotides are intracellular messenger molecules. Cyclic AMP (cAMP) and cyclic GMP (cGMP) are synthesized by adenylyl and guanylyl cyclase from ATP and GTP, respectively. Cyclic nucleotide signalling plays diverse functions across the phylogenetic tree. In mammals, for example, cAMP has multiple roles ranging from auditory function to mediating hormone action. cGMP is a central player in processes such as cardiac function and light detection in the eye. In single-celled eukaryotes, again cyclic nucleotides have myriad roles ranging from differentiation in *Dictyostelium* [3] to motility in ciliates [4]. cAMP and cGMP are hydrolysed by cyclic nucleotide phosphodiesterases (PDEs), which inactivate the messenger molecules. When cellular cyclic nucleotide concentrations reach threshold levels, following activation of a cyclase/inactivation of a PDE, they stimulate cAMP-dependent protein kinase (PKA) or cGMP-dependent protein kinase (PKG). The only other apparent cyclic nucleotide effector in malaria parasites is a molecule with putative cAMP-binding sites (PF3D7_1417400), designated PfEpac [5,6]. Prior to the availability of the full *P. falciparum* genome sequence, identification of some of these key enzymatic components in malaria parasites was a laborious process, but bioinformatic approaches have facilitated the task. Subsequent steps involved demonstration of biochemical activity and also functional analysis, which is still under way using genetic approaches (figure 2 and table 1). Although genetic manipulation of *P. falciparum* remains a significant challenge, the tools available have improved greatly in recent years and so advances in our understanding of these parasite signalling pathways will accelerate in the coming years. Functional studies of the key enzymatic components also facilitate our interpretation of pharmacological studies and in some cases necessitate a re-evaluation of the data.

2. Cyclase enzymes are distinct from those of other organisms

It has been reported that the properties of adenylyl cyclase activity derived from *P. falciparum* and the infected human red blood cell are distinct; for example, the host enzyme is activated by forskolin, whereas that of the blood stage parasite is not [23]. Malaria parasites have two adenylyl cyclases and two guanylyl cyclases that are topologically different to those of other organisms outside of the alveolate superphylum [24]. Although it is a predicted integral membrane protein, adenylyl cyclase alpha (ACα) is quite distinct from the membrane-associated mammalian G protein-dependent adenylyl cyclases. It has an apparently bifunctional structure with a C-terminal catalytic domain containing all the
Table 1. Phenotypes of mutants corresponding to pathway components. Details of the mutants that have been generated to date for the key players in malaria parasite cyclic nucleotide signalling (n.a., not applicable; n.d., not determined).

| protein description | type of mutant | blood stage phenotype | sexual stage phenotype | sporozoite/liver stage phenotype | references |
|--------------------|----------------|-----------------------|------------------------|-------------------------------|------------|
| adenyl cyclase α    |                |                       |                        |                               |            |
| *P. falciparum* (PF3D7_1404600) | no mutant | n.a.                  | n.a.                   |                               |            |
| *P. berghei* (PBANKA_1037500) | gene deletion | none                  | none                   | dramatically reduced apical exocytosis in response to liver cell transit | [9]        |
| adenyl cyclase β    |                |                       |                        |                               |            |
| *P. falciparum* (PF3D7_0802600) | no mutant | n.a.                  | n.a.                   |                               |            |
| *P. berghei* (PBANKA_1227600) | gene deletion | essential *           | n.a.                   |                               | [10]       |
| guanylyl cyclase α  |                |                       |                        |                               |            |
| *P. falciparum* (PF3D7_1138400) | no mutant | n.a.                  | n.a.                   |                               |            |
| *P. berghei* (PBANKA_0910300) | gene deletion | slow *                |                        |                               | [10,11]    |
| guanylyl cyclase β  |                |                       |                        |                               |            |
| *P. falciparum* (PF3D7_1360500) | gene deletion | none                  | none                   | n.d.                          | [12]       |
| *P. berghei* (PBANKA_1136700) | gene deletion | slow *                |                        | ookinete motility and mosquito midgut invasion defect | [6,11,13] |
| phosphodiesterase α |                |                       |                        |                               |            |
| *P. falciparum* (PF3D7_1209500) | gene deletion | none                  | n.d.                   | n.d.                          | [14]       |
| *P. berghei* (PBANKA_1019350) | gene deletion | none                  |                        |                               | [10]       |
| phosphodiesterase β |                |                       |                        |                               |            |
| *P. falciparum* (PF3D7_1321500) | no mutant | n.a.                  | n.a.                   | n.a.                          |            |
| *P. berghei* (PBANKA_1419800) | gene deletion | essential *           |                        |                               | [10]       |
| phosphodiesterase γ |                |                       |                        |                               |            |
| *P. falciparum* (PF3D7_1321600) | gene deletion | none                  | none                   | n.d.                          | [12]       |
| *P. berghei* (PBANKA_1419900) | no mutant | n.a.                  | n.a.                   | n.a.                          |            |
| *P. yoelii* (PY17X_1421600) | gene deletion | peak blood stage parasitaemia reduced by 50% | none | reduction in salivary gland sporozoites (55%) and substrate-dependent gliding defect | [15]       |
| phosphodiesterase δ |                |                       |                        |                               |            |
| *P. falciparum* (PF3D7_1470500) | gene deletion | none                  | gamete emergence        | n.a.                          | [12]       |

(Continued.)
canonical residues required for cAMP synthesis and six predicted transmembrane domains, which have significant sequence identity to voltage-gated K⁺ channels. Expression in Dictyostelium and Xenopus confirmed the predicted adenylyl cyclase activity [25], while activity of the predicted voltage-gated K⁺ channel has so far not been confirmed [26]. However, in the related ciliate Paramecium, which has a gene with the same configuration, it was demonstrated prior to gene sequencing that a purified native adenylyl cyclase has the properties of a voltage-gated K⁺ channel [27]. It will be important to understand the physiological significance of a malaria parasite adenylyl cyclase enzyme which might transport K⁺ ions to regulate cAMP synthesis.

ACα (PBANKA_1037500) has been deleted in the P. berghei rodent malaria parasite [9]. Blood stage parasite growth and gametocyte production was not affected by gene deletion, nor was subsequent development in the mosquito, up to and including sporozoite growth and gliding. However, mutant sporozoites lacking this gene were unable to undergo apical regulated exocytosis in response to transit of sporozoites through hepatocytes (a prerequisite for infection to be established). Mutant sporozoites also showed a 50% reduction in invasion of a mouse hepatoma cell line [9]. The study also reported that pharmacological elevation of cAMP levels and inhibition of PKA activity, respectively, stimulated or reduced apical regulated exocytosis in both P. yoelii and P. falciparum sporozoites. Interestingly, in light of the predicted voltage-gated K⁺ channel domain of ACα, this study found that extracellular K⁺ is also required for sporozoite exocytosis. Together these results point to a role for cAMP signalling in apical regulated exocytosis in sporozoites. The involvement of intracellular messengers in

Table 1. (Continued.)

| protein description | type of mutant | blood stage phenotype | sexual stage phenotype | sporozoite/liver stage phenotype | references |
|---------------------|----------------|----------------------|-----------------------|---------------------------------|------------|
| P. berghei (PBANKA_0835600) | gene deletion | none | ookinete motility and mosquito midgut invasion defect | n.a. | [6] |
| **cAMP-dependent protein kinase catalytic subunit** | | | | | |
| P. falciparum (PF3D7_0934800) | no mutant | n.a. | n.a. | n.a. | |
| P. berghei (PBANKA_0835600) | gene deletion | essential * | | | [10] |
| **cAMP-dependent protein kinase regulatory subunit** | | | | | |
| P. falciparum (PF3D7_1223100) | over-expression | growth rate reduced by 78% | reduced stiffness of immature gametocytes | | |
| P. berghei (PBANKA_1438000) | no mutant | n.a. | n.a. | n.a. | |
| **cAMP-dependent protein kinase** | | | | | |
| P. falciparum (PF3D7_1436600) | inhibitor-resistant mutant ** | block in schizont rupture | block in gamete egress | n.d. | [17,18] |
| P. falciparum (PF3D7_1436600) | FKBP DD knockdown | block in schizont rupture | | n.d. | [19] |
| P. berghei (PBANKA_1008200) | inhibitor-resistant mutant ** | block in schizont rupture | block in gamete egress, ookinete motility defect | n.d. | [20] |
| P. berghei (PBANKA_1008200) | conditional gene excision | n.a. | n.a. | block in late liver stage development | [21] |
| **Epac-like protein** | | | | | |
| P. falciparum (PF3D7_1417400) | truncation | none | n.a. | n.a. | [22] |
| P. berghei (PBANKA_1025300) | no mutant | n.a. | n.a. | n.a. | |

*PlasmoGem phenotype
**in conjunction with specific kinase inhibitor
albumin-stimulated sporozoite gliding was examined using pharmacological agents in a subsequent study. It was concluded that cAMP and PKA activity stimulated gliding and that elevation of cAMP levels could bypass the need for addition of albumin [28]. Using the PDE inhibitors IBMX and Zaprinast, the authors found that only the former stimulated sporozoite gliding and concluded that cAMP rather than cGMP signalling is involved in sporozoite motility. Curiously, it has previously been shown that IBMX does not inhibit PDE activity in P. falciparum blood stage parasites [16,29], but it is possible that it can inhibit PDE activity in sporozoites from rodent malaria parasites. The negative result obtained with Zaprinast (which is expected to raise cGMP levels) is seemingly at odds with the more recent observation that PKG activity is required for sporozoite gliding [30], but it is possible that the PDE activity in sporozoites is insensitive to Zaprinast. The authors next measured the effects of agents that influence mammalian heterotrimeric G proteins, which in mammals regulate transmembrane adenylyl cyclases (cholera toxin: stimulatory and mastoparan: inhibitory), to investigate whether this could be the mechanism that regulates cAMP synthesis during sporozoite gliding. Although the use of both of these agents suggested a role for heterotrimeric G proteins in sporozoite gliding, the apparent absence of the encoding genes in the parasite genome makes the results difficult to interpret. A more recent study investigated the effects of cholera toxin and mastoparan on P. falciparum sexual commitment. It was surprisingly found that mastoparan led to elevated CAMP levels and that cholera toxin had no effect, and it was concluded that the effects of both agents were unlikely to be via interaction with heterotrimeric G proteins [31].

The second adenylyl cyclase, ACβ, is an orthologue of the bicarbonate-sensitive soluble adenylyl cyclases [25]. So far, this gene has proved refractory to deletion in both P. berghei and P. falciparum, suggesting that it may be essential in the asexual blood stage. This is consistent with the fact that high levels of ACβ mRNA were measured in P. falciparum late blood stages [32]. Also, an antibody raised to P. falciparum PfACβ (PF3D7_0802600) reacted with both late schizonts and free merozoites [5]. Two distinct classes of soluble AC inhibitors (KH7 and 2CE) were found to kill P. falciparum blood stage parasites with LD₅₀ values of 8.5 μM and 60 μM, respectively [33], further suggesting that PfACβ is essential in blood stages. Experiments using synchronized cultures showed that parasites died when KH7 was added during the first 24 h of the cycle, but were most sensitive between 24 and 31 h post-invasion. Expression of the N-terminal 785 amino acids of PfACβ (containing the predicted catalytic domains) in insect cells did not allow investigation of bicarbonate sensitivity of the enzyme, but it did reveal a distinct pH sensitivity (optimum 7.5), which led the authors to hypothesize that PfACβ acts as a pH sensor during blood stage development. Exposure of merozoites to a buffer containing low levels of K⁺, as would be encountered on entering the plasma post egress, triggered calcium release and discharge of AMA1 (PF3D7_1133400) from the micronemes. The buffer also triggered bicarbonate ion-sensitive cAMP synthesis that was reduced by addition of KH7 suggesting a key role for cAMP synthesis by PfACβ in these events [5,34]. It has been shown in Toxoplasma that ACβ is located on the outer face of the rhoptries facing the cytosol [7] and its rhoptry association is dependent on the presence of an interacting partner (AIP) of the armadillo repeats only protein (ARO). The latter is also known to be targeted to the rhoptries in P. falciparum, where it is attached to the cytosoic face of the rhoptry membrane by dual acylation [8]. It has also been observed that the P. falciparum PKA regulatory subunit has a similar location to PfARO (PF3D7_0414900) and is also probably targeted to the rhoptries [8]. Interestingly, PfARO is phosphorylated in a PKG-dependent manner at Ser₃₃ and Ser₃₆ [35], suggesting this may be a focal point for crosstalk between these two pathways during merozoite invasion.

The two malaria parasite guanylyl cyclases, GCα and GCβ, have unusual bifunctional structures [36], which are only shared by apicomplexans and ciliate relatives such as Paramecium and Tetrahymena [4]. They comprise a pair of C-terminal cyclase catalytic domains and two sets of six predicted transmembrane helices, reminiscent of the topology of mammalian G protein-dependent adenylyl cyclases. Key purine-binding residues in their catalytic domains, however, are clearly characteristic of guanylyl cyclases, and heterologous expression of the P. falciparum GCβ (PF3D7_1360500) catalytic domain [36] and the whole Paramecium enzyme in insect cells [4] have demonstrated that they are functional guanylyl cyclases. Curiously, amino acid motifs characteristic of the mammalian C1 AC catalytic domain are found in the C2 domain of the alveolate enzymes and vice versa. Evidence for membrane association was obtained using immunoelectron microscopy in both of these studies. The N-terminal region of all the alveolate GCs has sequence and structural homology with type IV P-type ATPases [4,11,36], but no functional activity has so far been demonstrated for this domain. Interestingly, proteolytic processing of the region linking the two functional domains of the Paramecium GC was observed in Sf9 cells, but whether this happens in vivo is not known. It will be intriguing to uncover the significance of the genetic linkage of these two distinct enzyme activities in apicomplexan parasites and their free-living relatives.

Early pharmacological studies [37–39] suggested cGMP as a possible regulator of exflagellation. Xanthurenic acid, a product of tryptophan catabolism and present in the mosquito midgut, is known to stimulate exflagellation [40,41]. Later it was shown that addition of xanthurenic acid to crude membrane preparations from P. falciparum gametocytes led to increased levels of cGMP, suggesting that xanthurenic acid may stimulate guanylyl cyclase activity (directly or indirectly) [42], again linking this pathway with exflagellation.

Transcriptomic studies have shown that the patterns of mRNA expression for PfGCα (PF3D7_1138400) and PfGCβ are very different (http://plasmodb.org), suggesting they each have roles in distinct life-cycle stages. Two studies have reported deletion of GCβ (PBANKA_1136700) in P. berghei which revealed a role in ookinete motility and mosquito midgut invasion [6,13]. Exflagellation occurred at a similar rate in mutant and wild-type lines, indicating that GCβ is not essential for this process in P. berghei. Similar findings were obtained following disruption of the P. falciparum GCβ in both the cyclase domain and the P-type ATPase domain. Gametogenesis was able to proceed normally despite the fact significantly reduced guanylyl cyclase activity was measured in mutant gametocyte membrane preparations [12]. It was deduced that the residual GC activity was due to the presence of GCα, which may also
be expressed in gametocytes, and that this was sufficient for
gametogenesis to occur normally. GCox has proved refractory
to deletion in both *P. berghei* and *P. falciparum*, suggesting
an essential role in blood stage development [6,11].

### 3. The parasite phosphodiesterases are all
membrane-associated

There are four cyclic nucleotide phosphodiesterase enzymes
(PDEαs-δ) encoded in the malaria parasite genomes, but
curiously there are potentially 18 in *Toxoplasma gondii*
(TGGT1 strain; http://toxodb.org/toxo/). The malaria paras-
ite PDEs are predicted to have up to six transmembrane
domains, suggesting they are integral membrane proteins.
The mammalian PDE superfamily is much more complex
than that of *Plasmodium*. The mammalian PDEs are divided
into 11 functionally divergent but structurally similar gene
families, most containing several different genes. These genes
are frequently spliced or transcriptionally processed to give
over 100 PDE isoforms [43]. One family (PDE3) contains
a single transmembrane domain, but, unlike *Plasmodium* PDEs,
no mammalian PDE contains multiple membrane spanning
domains. Mammalian PDEs contain 15 conserved residues,
thirteen of which are conserved in all four PIPDE enzymes
[44]. A number of transcriptional studies (available via
http://plasmodb.org) have shown that the four *P. falciparum*
PDEs exhibit life cycle stage-specific patterns of expression.
PDEα (PF3D7_1209500) and PDEβ (PF3D7_1321500) are
the predominant isoforms expressed in blood stage parasites
and peak at the late trophozoite and schizont stages, while
PDEγ (PF3D7_1231600) and PDEδ (PF3D7_1470500) are
predominantly expressed in sporozoite and gametocyte
stages, respectively.

A modelling study of the malaria parasite PDEs has been
carried out using the human PDE9A crystal structure as a
template [44]. Interestingly, the model could accommodate
cGMP binding, but not cAMP binding, but as the authors
say, roles for cAMP and PKA in the parasite have been estab-
lished and so at least one of the four PIPDEs is likely to be
able to hydrolyse cAMP and perhaps be dual specific. To
date, it has been possible to measure catalytic activity for
only PDEα (PIPDE1) by expressing the C-terminal catalytic
domain in *Escherichia coli* [14,29]. This has shown that
the enzyme specifically hydrolyses cGMP. The catalytic specifici-
ties of PDEs and PDEδ for cGMP have been deduced from
phenotype analysis of null mutants (but require confirmation;
see below) while that of PDEβ remains unknown as it is the
only one refractory to deletion. Expression of PDEα also
allowed analysis of its sensitivity to some commonly used
PDE inhibitors. Importantly, it was shown that both IBMX
and theophylline, two PDE inhibitors that have been used in
a number of studies on malaria parasites and have little
or no activity against *Plasmodium* PDEs with EC_{50} >
100 μM [14,16,29]. It is important to note, in this respect,
that some studies involving parasite PDEs use preparations
in which the host red cell has not been removed and so
human PDEs will be present, whereas some studies use sapo-
nin lysis to remove the host cell material. Yuasa and
colleagues were able, however, to identify Zaprinast as a
low micromolar inhibitor of recombinant PDEα and native
PDE activity in *P. falciparum* blood stage parasites as well
as blood stage growth [29]. Zaprinast has proved an

extremely valuable tool (table 2) in a number of subsequent
studies on cyclic nucleotides in apicomplexan parasites, and
has helped provide proof of concept that malaria parasite
PDEs are valid drug targets. The effects of analogues of the
human PDE5 inhibitor, Tadalafil, on malaria parasites have
also been reported, with EC_{50} values down to the sub-micro-
molar level [57]. Progress towards achieving selectivity for
the parasite PDE by reducing activity against human PDE5
was also made by the same team by modifying the piperonyl
group [58]. Another sub-micromolar inhibitor of malaria
parasite PDEs, BIPPO, derived from a human PDE9 series,
has an EC_{50} of 0.4 μM against *P. falciparum* proliferation
and an IC_{50} of 150 nM against recombinant PDEδ, but still
retains an IC_{50} of 30 nM against human PDE9. Nonetheless,
this compound will also be a very useful tool for studying
cyclic nucleotide signalling in the parasite and could be an
important lead for drug discovery [50].

PDEδ has been deleted in *P. falciparum*, showing that it is
not essential for blood stage growth in *vitro*. Phenotype
analysis suggested that ring-stage parasites appeared earlier
in synchronized cultures of mutant parasites compared
with wild-type, but it was not clear whether this was related
to the absence of the PDEα gene. There was no evidence of
upregulation of mRNA levels of the other three PDEs to com-
 pense for the deleted gene. The study also highlighted the
existence of two splice variants of PIPDEα (A and B), one
of which encodes a protein with four predicted TM domains
rather than six, but the significance of this is not known.
cGMP-PDE activity in membrane fractions of the PDEα
mutant was 20% lower than those from wild-type parasites,
with no effect on cAMP-PDE activity. This is consistent
with the findings of the recombinant expression studies that
PDEα is cGMP specific. Interestingly, the authors also
found that addition of excess cGMP to blood-stage membrane
fractions from wild-type and PDEα knockout parasites greatly
reduced cAMP hydrolytic activity in PDE assays, suggesting
that one of the other PDEs expressed in blood stages is likely
to be dual specific [14]. PDEδ, which has proved refractory
to deletion in *P. falciparum* and *P. berghei* [10], is the most likely
candidate for this activity, but this is yet to be demonstrated.

PDEβ has been deleted in both *P. falciparum* [12] and *P. berghei* [6], showing that this gene is dispensable for blood
stage replication in *vitro*. Deletion of PDEβ in *P. falciparum*
led to a 50% reduction in native cGMP hydrolytic activity
in gametocytes compared with wild-type, with a correspond-
ing twofold increase in cellular cGMP levels in late-stage
gametocytes [12]. cAMP-PDE activity levels were consistently
low, and indistinguishable between wild-type and mutant
parasites. Interestingly, addition of Zaprinast to wild-type
gametocyte membrane fractions reduced cGMP hydrolytic
activity by around 75%, but it was totally ablated in PDEδ
mutants. This suggests that the residual PDE activity is more
sensitive to Zaprinast than PDEδ. Together these results
indicate that PDEδ has cGMP hydrolytic activity and that at
least one other cGMP-PDE is active in gametocytes. Although
PDEδ mRNA and protein are present in gametocytes, their
development up to maturity is not affected by the absence
of this gene. However, upon stimulation of gametogenesis
with xanthurenic acid, the ability of mutant parasites to
round up was significantly reduced, and exflagellation of
male gametocytes was reduced dramatically. Examination of
mutant and wild-type parasites by immunofluorescence
microscopy using an antibody against the human red cell
Table 2. Pharmacological agents used to study cyclic nucleotide signalling in malaria parasites. Details of compounds that have been used to target *Plasmodium* proteins involved in cyclic nucleotide signalling. The parasite and host targets are indicated, if known, along with the potency reported for parasite or host cell and biochemical/phenotypic effects (n.a., not applicable; n.d., not determined).

| compound | *P. falciparum* target | host target | published potency | published potency | biochemical effect | published phenotypic effect |
|----------|------------------------|-------------|------------------|------------------|-------------------|-----------------------------|
| Compound 1 | cGMP-dependent protein kinase | n/a | IC_{50} 0.49 μM asexual blood stage growth [45] | no data available | inhibition of parasite cGMP-dependent protein kinase | inhibition of blood stage schizont rupture [17], inhibition of XA stimulated rounding up of gametocytes and gamete emergence [18] |
| Compound 2 | cGMP-dependent protein kinase | n/a | EC_{50} 395.0 nM ± 21.9 asexual blood stage growth [46] | no data available | inhibition of parasite cGMP-dependent protein kinase | inhibition of blood stage schizont rupture [17], inhibition of XA stimulated rounding up of gametocytes and gamete emergence [18] |
| Zaprinast | phospho-diesterases | phospho-diesterase 5 | IC_{50} 33.7 ± 1.3 μM for gametocytes and 3.0 ± 1.2 μM for schizonts [18] | IC_{50} 1.3-3.2 μM PDE5 isoforms from cell lysate [47] | elevation of cellular cyclic nucleotide levels | induction of blood stage egress [48], induction of rounding up of gametocytes and gamete emergence [18] |
| Sildenafil | phospho-diesterases | phospho-diesterase 5 and 6 | IC_{50} 75.6 μM ± 1.6 for gametocytes and 22.5 μM ± 1.8 for schizonts [18] | IC_{50} 3.5 nM PDE5 33 nM PDE6 [49] | elevation of cellular cyclic nucleotide levels | induction of rounding up of gametocytes and gamete emergence [18] |
| BIPPO | phospho-diesterases | n/a | IC_{50} 0.4 μM ± 0.14 asexual blood stage growth [50] | no data available | elevation of cellular cyclic nucleotide levels | induction of blood stage egress [50] |
| IBMX | n/a | phospho-diesterases | IC_{50} >200 μM asexual blood stage growth [18] | IC_{50} 5.8-26.2 μM [51] | elevation of host cell cyclic nucleotide levels? | n/a |
| H89 | cAMP-dependent protein kinase catalytic domain | cAMP-dependent protein kinase catalytic domain | IC_{50} 2.9 μM ± 1.6 asexual blood stage growth [52] | Ki 0.05 μM [53] | inhibition of cAMP-dependent protein kinase | inhibition of blood stage schizont development [52], inhibition of blood stage merozite invasion [54], induces deformability in stage III gametocytes [16] |
| KT5720 | cAMP-dependent protein kinase catalytic domain | cAMP-dependent protein kinase catalytic domain | IC_{50} 10 μM asexual blood stage growth [54] | Ki 60 nM [53] | inhibition of cAMP-dependent protein kinase | inhibition of blood stage merozite invasion [54], induces deformability in stage III gametocytes [16] |
| KH7 | adenyl cyclase β | soluble adenylate cyclase | IC_{50} 8.5 μM asexual blood stage growth [33] | IC_{50} 10 μM [55] | inhibition of cAMP synthesis | inhibition of blood stage schizogony [33], inhibition of blood stage microneme release [5] |
| Forskolin | n/a | soluble adenylate cyclase | no effect on the blood stage enzyme [23] | IC_{50} 2-10 μM [56] | stimulation of host cell cAMP synthesis | n/a |
membrane protein Band 3 showed that although most mutant parasites could partially round up following stimulation of gametogenesis, very few parasites had emerged from the encapsulating red cell. This suggests that elevated levels of cGMP, and perhaps premature PKG activation in gametocytes lacking PDE8, have deleterious effects on gamete emergence. However, premature activation of gametocytes would be expected if this were the case, and so further work is needed to understand this phenotype. The results also indicate that PDE8 is important for regulation of cGMP levels during gametogenesis, and that both male and female gamete emergence is affected. This study also reported that PDEy, which is also expressed in gametocytes at the mRNA level (PlasmoDB.org), was dispensable for P. falciparum blood-stage growth and gametocyte development in vitro, but in contrast to PDE8 mutants, there was no effect on gametogenesis [12].

Perhaps surprisingly, the phenotype was different in P. berghei; deletion of PDE6 had no effect on gamete emergence [6]. This suggests that another PDE might be able to compensate for its absence in P. berghei by maintaining cGMP concentrations in mature gametocytes below the level required for PKG activation. However, deletion of P6PDE6 had a dramatic effect on ookinete development. During the first 12 h of development, PDE6- parasites progressed to the expected banana-shaped ookinetes, but by 24 h they had ‘dedifferentiated’ into rounded or stumpy forms. This resulted in a more than 94% decrease in the number of oocysts that developed subsequently, compared with wild-type parasites. Interestingly, if these mutant ookinetes were injected into the haemocoel of the mosquito, transmission was restored to wild-type levels. Therefore, the mutant ookinetes were viable, but could not cross the epithelium of the mosquito midgut due to a lack of forward gliding movement. This abnormality was explained by some marked ultrastructural abnormalities (gaps in the inner membrane complex and misorientation of the underlying subpellicular microtubules) rather than a lack of motor activity [6]. Interestingly, the dedifferentiation phenotype could be reversed by addition of a specific PKG inhibitor, confirming that inappropriate activation of this enzyme had taken place following PDE6 deletion resulting in an elevation of cGMP levels. This conclusion was further strengthened by the fact that Gcβ-PDE6- double mutant ookinetes also had normal morphology. Another interesting finding was that a CDPK3- PDE6- double mutant glided normally (prior to dedifferentiation), thereby reversing the CDPK3- mutant gliding phenotype. This indicated that elevated signalling through cGMP and PKG could compensate for the absence of this calcium-dependent kinase, which suggests there may be overlapping substrates for these two protein kinases.

Analysis of the localization of PDEy using an epitope tagged P. yoelii transgenic line showed low-level expression in blood stages that partially co-localized with a marker of the endoplasmic reticulum (ER), but strong expression in sporozoites [15]. Although deletion of PDEy caused a 50% reduction in peak blood-stage parasitaemia during the second half of a 20-day mouse infection, the gene is not essential for blood-stage replication. Furthermore, gametocyte and mosquito-stage development was unperturbed, including sporozoite formation in oocysts. However, there was a 55% reduction in the number of salivary gland sporozoites in mutant parasites compared with wild-type, and only very high sporozoite doses resulted in successful transmission to a proportion of mice, to establish a blood-stage infection. The authors demonstrated that this low level of transmission was due to the fact that the sporozoites lacking PDEy are defective in substrate-dependent gliding motility. Transcriptome analysis of PDEy- mutant sporozoites showed upregulation of PDEβ and PDEδ and downregulation of several mRNAs encoding proteins involved in sporozoite motility and hepatocyte invasion [15]. Mutant sporozoites were found to have an almost 20-fold higher cGMP concentration than wild-type sporozoites, strongly suggesting this to be the cause of the defective gliding phenotype. Consistent with this assertion, the PDE inhibitor Zaprinast reduced sporozoite gliding in a dose-dependent manner. The authors point out that according to these and previous observations, it is possible that cGMP and cAMP may have opposing effects on sporozoite function and that crossstalk between the two pathways could occur [15].

4. PKG is essential in all of the key stages of the malaria parasite life cycle

The apicomplexan PKGs have both structural and biochemical properties that distinguish them from mammalian isoforms. For example, apicomplexan PKGs have four recognizable cGMP-binding domains (A, B, C, D); only three of these are functional and domain C is degenerate. By contrast, mammalian PKGs have only two cGMP-binding domains. In the absence of cGMP, PKG adopts a globular configuration where a pseudo-substrate, autoinhibitory sequence occupies the active site and so the enzyme is kept in an inactive state (figure 3). On binding cGMP, the conformation becomes more open and extended, and the autoinhibition is released, thereby exposing the active site to allow phosphorylation of substrate proteins. In P. falciparum, cGMP-binding site D has by far the highest affinity for cGMP and selectivity over cAMP. Crystallography carried out on this binding site has identified the key determinants of cGMP binding and identified a distinct capping mechanism. A conditional reverse genetic approach was taken to show that substitution of either of these capping residues is lethal to blood-stage parasites [59].

Two PKG isoforms have been identified in the coccidian parasites Eimeria and Toxoplasma. One of these is dually acylated at the N-terminus; modifications that allow membrane association [60]. Mammalian PKGI is membrane associated by myristoylation [61]. By contrast, in Plasmodium PKGs, the acylation motifs are absent and although a subpopulation of the P. falciparum PKG (PFPKG; PF3D7_1436600) apparently becomes associated with the ER membrane in late schizonts, the means of attachment is unknown [62]. Mutational analysis to examine the properties of the individual apicomplexan cGMP-binding domains has revealed that, unlike mammalian PKGs, they are relatively insensitive to 8-substituted cGMP analogues [63,64]. Progress with studying the malaria parasite PKG has been greatly facilitated by the existence of specific inhibitors which were generated through pioneering work by a group at Merck who identified a tri-substituted pyrrole (Compound 1) in a whole cell screen of E. tenella, a related apicomplexan parasite of chickens [65]. Using a range of biochemical approaches, they found that the target of Compound 1 is the E. tenella PKG and a
subsequent medicinal chemistry effort led to the generation of a more potent imidazopyridine PKG inhibitor, Compound 2 [66]. At a similar time, another study carried out a biochemical analysis of the *P. falciparum* PKG and showed that it had a very similar structure and properties to the coccidian enzyme [64,67]. The two Merck PKG inhibitors were subsequently reported to be potent inhibitors of this malaria parasite PKG [18,45], and this proved crucial to functional...
studies when used in conjunction with an inhibitor-resistant transgenic line. This chemical genetic approach relies on the fact that the apicomplexan PKG has a structural feature, which is distinct from that of the human PKG and importantly the majority of serine threonine kinases, that allows inhibitor binding. The parasite PKG has a relatively small amino acid residue (threonine) in the so-called gatekeeper position (Thr618 in PfPKG). This is thought to allow access of the fluorophenyl group, shared by Compound 1 and Compound 2, to a small hydrophobic pocket which adjoins the kinase ATP-binding pocket. Substitution of the small gatekeeper residue with one with a bulkier side chain (such as glutamine or methionine) prevents access of the inhibitor to the gatekeeper pocket and renders the enzyme insensitive to it (figure 4). Introducing the gatekeeper mutant PKG into the parasite by complementation (Toxoplasma and Eimeria [68,69]) or allelic replacement (P. falciparum [17,18] and P. berghei [20]) confers inhibitor resistance to the transgenic line (the degree of resistance conferred varies with species and life-cycle stage). Comparison of the effects of the inhibitor on wild-type and gatekeeper mutant parasites first allowed demonstration that PKG is the primary target of the compounds, but importantly also allowed determination of the mechanism by which PKG inhibitors block blood stage schizont rupture and merozoite egress. It revealed that inhibition of PKG could not rupture, but whereas PKG inhibition prevented proteolytic processing of key merozoite proteins by the essential protease PfSUB1 (PF3D7_0507500) [73], the CDPK5 phenotype was downstream of this proteolytic activity and cGMP signalling [74]. A subsequent study examined the underlying mechanism by which PKG inhibitors block blood stage schizont rupture and merozoite egress. It revealed that inhibition of PKG blocked release of PfSUB1 from the apical organelles known as exosomes, indicating that cGMP signalling and PKG activity are required for release of PfSUB1 from these organelles. Furthermore, in PKG inhibitor-treated schizonts, AMA1 could not be released from the micronemes onto the merozoite surface [48]. A conditional knockdown approach using a fusion with the destabilization domain (DD) showed that depletion of PKG led to a block in P. falciparum blood stage egress [19], a phenotype that is consistent with that obtained with the chemical genetic approach.

Another vital cellular process that was shown to be PKG-dependent using the gatekeeper mutant approach is calcium release from internal stores (figure 2). Elevation of cGMP...
levels stimulated by zaprinast, which triggers merozoite egress, was found to stimulate calcium release. This was measured by loading schizonts with the calcium-sensitive fluorescent dye, Fluo-4 AM. Addition of PKG inhibitor blocked calcium release in a dose-dependent manner. Analysis of phospholipids by mass spectrometry in *P. falciparum* wild-type and gatekeeper mutant schizonts treated simultaneously with zaprinast and Compound 2 showed PKG-dependent changes in phosphoinositide levels. For example, levels of PIP$_2$ (which in higher eukaryotes would be hydrolysed by phospholipase C (PLC) to generate IP$_3$, which would in turn lead to activation of IP$_3$-stimulated calcium channels leading to calcium release) were reduced by PKG inhibition. This study indicated that the process of PKG-dependent merozoite egress is mediated by calcium release which is linked to phosphoinositide metabolism [20]. To date it is not known what acts upstream of cGMP signalling to trigger egress, but there is evidence of events prior to PKG activation, with permeabilization of the parasitophorous vacuole membrane occurring 10–30 min before its PKG-dependent breakdown [75]. Interestingly, in *Toxoplasma* it has been shown that the plant hormone abscisic acid triggers egress in a calcium-dependent manner [76], but it remains to be seen whether this molecule has a role in malaria parasites.

The gatekeeper mutant and PKG inhibitor have also been used in conjunction with global phosphoproteomics to identify potential PKG substrates in *P. falciparum* schizonts. Peptides derived from schizont proteins from PKG inhibitor-treated and untreated cultures of wild-type and gatekeeper mutant parasites were labelled with isobaric tags and analysed using quantitative mass spectrometry. A total of 107 phosphosites from 69 individual proteins were regulated in a PKG-dependent manner, revealing involvement of cGMP signalling in a range of cellular processes including egress/invasion, motor function, signal transduction, ion and protein transport, and chromatin regulation. Interestingly, PKG-dependent phosphosites were identified on GCa, PDEβ and ACβ suggesting feedback loops involving cGMP and possible crosstalk with cAMP signalling. Work is under way to determine the physiological significance of a number of these phosphosites in key proteins. This approach identified PKG-dependent phosphorylation events in schizonts, but in many cases it is likely that the proteins are actually phosphorylated by kinases that act downstream of PKG. One protein that was found to be a direct substrate of PKG in schizonts is CDPK1 (PF3D7_0217500), suggesting that PKG may play a dual regulatory role in calcium signalling, stimulating calcium release and direct phosphorylation of calcium-dependent kinases [35]. A recent study reported the successful generation of a *P. falciparum* CDPK1 gatekeeper mutant (harbouring a Thr$_{35}$Ala substitution). This line had a significantly reduced kinase activity, but interestingly had a greater sensitivity to the PKG inhibitor Compound 2 than wild-type parasites. This was interpreted as indicating that PKG was probably compensating for the reduced activity of the mutant CDPK1 enzyme. We believe it is more likely that the increased sensitivity of this line to Compound 2 is due to the fact that CDPK1 is also a target of this inhibitor and that because the gatekeeper mutant CDPK1 cannot bind Compound 2, the amount of free compound available to block PKG function is elevated, resulting in a lower EC$_{50}$ value than in wild-type. The increased transcript levels of CDPK5 and CDPK6 (PF3D7_1122800) measured by the authors in the mutant line probably indicate that these closely related kinases compensate for the reduced CDPK1 activity [77]. CDPK7 has been shown to be important for early blood-stage development and is therefore likely to operate independently of PKG. This kinase is unique among the CDPKs in having a pleckstrin homology domain that has been shown to bind to PIP$_2$ and is proposed to guide its subcellular localization [78].

PKG has also been shown to play an essential role in *P. berghei* ookinetes. The PKG inhibitor Compound 1 prevented ookinete gliding with an EC$_{50}$ of 100 nM. PKG fused to the fluorescent reporter GFP identified a largely cytosolic location in ookinetes [79]. A subsequent study examined in more detail how cGMP signalling and PKG activity might regulate ookinete gliding. The study used quantitative mass spectrometry to analyse cGMP-dependent phosphorylation events using two parallel genetic approaches. The first used a GCβ knockout line and the second used the PKG inhibitor Compound 2 in conjunction with a *P. berghei* gatekeeper mutant parasite line (expressing a Thr$_{35}$Gln substitution). Changes in phosphorylation levels in either of the approaches detected key sites in PKG itself as well as in GCβ and PDEβ, showing potential mechanisms for self-regulation of the pathway. Interestingly, regulation of proteins involved in phosphoinositide metabolism and vesicle trafficking was measured in both of these two approaches in *P. berghei*. For example, phosphopeptides identified in three of the four P4 K (PBANKA_110940; PBANKA_072200) and PIP5 K (PBANKA_020310) isoforms were reduced significantly, suggesting that phosphoinositide metabolism is required for ookinete motility and that it is regulated by PKG. The lack of any change in phosphosites on PLC suggests that regulation is at the level of sequential phosphorylation of phosphatidylinositol (PI) rather than at the level of PIP$_2$ hydrolysis [20]. Interestingly, *P. berghei* ookinetes containing a P4 K (PBANKA_110940) phosphomutant (Ser$_{343}$Ala) showed a reduced gliding speed compared with wild-type parasites. Furthermore, a *P. berghei* line expressing HA-tagged PIP5 K (PBANKA_020310) showed a change in distribution from the cell periphery to the cytosol when treated with PKG inhibitor. Intriguingly, this PIP5 K has a predicted bifunctional structure with an N-terminal domain with homology to the EF-hand-containing neuronal calcium sensors, a subfamily of which activate guanylyl cyclase in mammals. The *P. falciparum* orthologue (PPIP5 K/NCS; PF3D7_0110600) has been shown to be activated by ADP-ribosylation factor (ARF1) to generate PIP$_2$ [80], providing another potential link between phosphoinositide metabolism and vesicular trafficking. Both of these observations strongly suggest a link between PKG and phosphoinositide metabolism. Analysis of ookinete phospholipids by quantitative mass spectrometry in the presence and absence of PKG inhibitor revealed reduced levels of PI4P and PI$_2$ synthesis following PKG inhibition. This in turn suggested a potential link to calcium release via hydrolysis of PIP$_2$ (by PLC) to give IP$_3$ (and diacylglycerol) that in other systems activates calcium channels. An IP$_3$-gated calcium channel sequence has not been identified in *Plasmodium* genomes, but it is likely that a protein exists in the parasite which performs this role. A transgenic *P. berghei* line expressing a fluorescent calcium reporter showed clearly that inhibition of PKG led to rapidly reduced levels of cytosolic calcium in ookinetes. Furthermore, the PKG-
mediated stimulation of gametogenesis by reduced temperature and xanthuric acid was also shown to stimulate calcium release, and this was detectable just 10 s after stimulation [20]. The *P. berghei* line expressing gatekeeper mutant PKG was used to demonstrate that the effects of PKG inhibition were specific.

In addition to its role in blood-stage egress and invasion, gametogenesis and ookinet motility, PKG has also been shown to be essential for late-liver-stage development in *P. berghei*. As it was not possible to delete the gene, an elegant conditional knockout approach was used, whereby gene excision was triggered during mosquito stage development by a drop in temperature [21]. Gene excision was detectable in the sporozoite stage. Injection of mice with these sporozoites did not lead to a subsequent blood-stage infection. Examination of liver stages in HepG2 cell cultures showed that development could apparently proceed normally up to 40 h post-infection; however, after 72 h, cultures appeared blocked at the late liver schizont stage with few merozoites (vesicles containing merozoites) released. A follow-up study using the PKG inhibitor Compound 1 showed that it blocked sporozoite gliding, invasion and infection, but in the light of the previous findings this was interpreted as off-target activity. However, a more recent study used the gatekeeper mutant approach in *P. berghei* for both PKG and CDPK4 (PF3D7_0717500) in conjunction with either a PKG inhibitor or a bumped kinase inhibitor (BK1 1294), used as a CDPK4 inhibitor, to demonstrate that PKG is in fact required for sporozoite gliding motility (and that Compound 1 was on target), microneme exocytosis and hepatocyte invasion, with CDPK4 also having a role in hepatocyte invasion [30]. The findings of this paper are seemingly at odds with the earlier work described above, which suggested that PKG is not required for sporozoite function. The authors pointed out that this discrepancy is likely to be due to residual PKG protein synthesized prior to conditional excision carried over into the sporozoites [30].

It is clear from the studies described above that PKG plays an essential role at almost every stage of the malaria parasite life cycle (figure 1). This emphasizes that this molecule might be worth pursuing as an antimalarial drug target. *In vivo* proof of concept has been established following a medicinal chemistry effort to optimize the imidazopyridines against the *P. falciparum* PKG. Oral dosing of a *P. falciparum* mouse model was able to clear blood-stage infection as well as block transmission of gametocytes to mosquitoes. This study also reported co-crystal structures that confirmed the importance of compound interactions within the gatekeeper pocket of the kinase [46]. An important related finding that has emerged over time is that PKG usually acts in concert with a stage-specific CDPK (figure 1). In some cases (e.g. gametogenesis and blood stage merozoite egress), it appears that PKG triggers release of calcium into the cytosol, which leads to the activation of the corresponding CDPK, but evidence suggests that PKG can also phosphorylate the partner CDPKs, which might fine-tune the stage-specific calcium response.

5. A role for PKA in multiple life-cycle stages is emerging

Unlike PKG, the cAMP-dependent protein kinase, PKA, is made up of distinct polypeptides encoded by separate genes (figure 3). The inactive enzyme is made up of a catalytic subunit (PKAc) and a regulatory subunit (PKAr) which contains a pair of cAMP-binding sites. When cAMP binds to PKAr, the complex dissociates and PKAc is thereby activated and can bind to substrate proteins which are then phosphorylated. When bound to PKAr, the PKAc enzyme is inactive, with part of PKAr binding to the active site via a substrate-like autoinhibitory sequence. In mammals, there are four regulatory subunits (two type R1 and two type R2), and a complex of four polypeptides made up of two catalytic subunits (three C types) and two regulatory subunits constitutes the inactive heterotetrameric state [81]. By contrast, in *Plasmodium*, there are single genes encoding both PKAc (PF3D7_0934800) [82] and PKAr (PF3D7_1223100) subunits. A unique feature of the malaria parasite PKA regulatory subunit sequence is that the dimerization domain characteristic of PKAr in other species is absent [83]. The sequence of *Plasmodium* PKAr suggests it may be a hybrid of the human R1 and R2 types [84], and crystallographic analysis confirmed that it has features that are characteristic of both human R1 and R2 types [85].

In mammals, the localization and function of PKA is determined by interaction with a family of A kinase activating proteins (AKAPs) [86]. AKAPs ensure that PKA activity is targeted to the correct cellular compartment and ensures its specificity of response to discrete cellular stimuli. However, the dimerization/docking domain of PKAr in other species allows tethering to AKAP in the *Plasmodium* PKAr sequence [83]. To date only one gene encoding an AKAP-like protein (PIKAL; PF3D7_0512900) has been identified in *P. falciparum*, and while no direct interaction with the parasite PKAr could be detected, there was evidence that it binds to Pf14–3-3 (PF3D7_0181200; which in turn binds to PKAr) and also to AMP and cAMP [87], suggesting a divergent AKAP mechanism in *Plasmodium* species. Intriguingly, *P. falciparum* Rab5A (PF3D7_0211200) and Rab7 (PF3D7_0903200) proteins have also been identified as potential PKAc interactors, and the authors suggested that Rabs may act as surrogate AKAPs [88,89].

The reported role of PKA in *P. falciparum* is varied, acting at multiple stages of the parasite life cycle. An early study detected *P. falciparum* blood stage parasite proteins that were phosphorylated in a cAMP-dependent manner, and further isolated a kinase activity that was cAMP-dependent thereby revealing the presence of PKA in malaria parasites [90]. The *P. falciparum* PKA activity detected in asexual blood-stage parasites was shown, as with the human red-cell-derived activity, to be sensitive to the PKA inhibitors H89 and PKI [52]. However, a later study using expression of the *P. falciparum* PKAc in a cell-free system demonstrated that it was much less susceptible to these inhibitors than mammalian PKA [91,92]. Two *P. falciparum* cloned lines (LE5 and T9/96) that differ in their ability to produce gametocytes were examined to determine whether they have differences in their adenyl cyclase or PKA activity. This was prompted by a report that addition of 1 mM cAMP to static cultures of the *P. falciparum* Z isolate could trigger conversion of almost 100% rings to gametocytes [93]. While AC activity was the same in both parasite clones, PKA activity was found to be significantly higher in the gametocyte-producing line, suggesting that this could reflect a causal relationship [90]. By contrast, a subsequent study attempting to reproduce the experiment performed by Kaushal *et al.* [93]
reported a negative effect of cAMP on asexual growth and gametocytogenesis [94], potentially indicating a cryptic pleiotropic effect of this molecule on the parasite cell cycle. A number of groups have investigated this enigmatic link between cAMP signalling and gametocyte formation. For example, Dyer & Day showed that treatment of malaria parasites with cholera toxin (which in mammals stimulates AC by activation of the Gs class of heterotrimeric G proteins) enhanced gametocyte formation [95]. The absence of genes encoding heterotrimeric G proteins from the Plasmodium genome, however, makes these results difficult to interpret. Others have found that cholera toxin has an effect on malaria parasites, but concluded that its target is unclear [31].

It has been reported that the human hormone melatonin causes in vivo synchronization of the rodent malaria parasite P. chabaudi and that this is achieved through calcium signalling [96]. Intriguingly, the same group has shown that 100 nM melatonin can promote early maturation of schizonts and stimulate elevated cAMP levels and PKA activity in P. falciparum blood stages in a phospholipase C-dependent manner, suggesting the involvement of calcium signalling [97]. The same study also showed that addition of 20 μM 6-Bz-cAMP, a cell-permeable cAMP analogue, can induce Ca<sup>2+</sup> release in the parasite cytosol, implicating cAMP in modulating downstream calcium-dependent processes. No change in cAMP was stimulated by forskolin, suggesting that the human red cell AC was absent from the preparation. The same two pathways have been implicated in mediating the effects of melatonin in P. chabaudi [98]. In a more recent study, this group have examined gene expression changes caused by melatonin and cAMP using RNA-Seq. Addition of melatonin led to a change in expression of 38 genes in trophozoites (101 with cAMP) which was dependent on the presence of an intact P. falciparum protein kinase 7 (PK7; PF3D7_0213400) gene; P. falciparum PK7 knockout line is unable to respond to melatonin. Gene ontology and KEGG pathway analysis indicated that components of the ubiquitin–proteasome system (UPS) pathway were regulated by both treatments in trophozoites [99].

It is known that the parasite transports many proteins into the host red cell; some of these are involved in the binding of infected red cells to host endothelial cells and some are involved in nutrient uptake and disposal. It has been shown in uninfected red cells that anion conductance can be activated by cAMP [100] and that PKA plays a role in anion conductance in the membrane of P. falciparum-infected red blood cells [83]. Injection of recombinant P. falciparum PKAr into the infected red cell using a patch-clamping pipette led to a decrease in anion conductance. Whether this is due to inhibition of erythrocyte-derived PKA is unclear, as it may be that PKA injected into the erythrocyte cytosol cannot access parasite PKA contained within internal limiting membranes. Furthermore, transgenic overexpression of PfPKAr had an inhibitory growth effect on the parasite, in conjunction with decreased anion conductance, indicating a role for this pathway in parasite cell cycle progression. This growth defect could be reversed by addition of IBMX (a PDE inhibitor expected to elevate cAMP levels) to the culture medium, which may indicate a role of erythrocyte host derived PDEs in this process as IBMX does not inhibit parasite PDEs (table 2). What is not clear is whether the cAMP production and transduction machinery involved in anion conductance in parasite-infected red cells is derived from the red cell or the parasite [100,101]. A recent conditional knockout study has clearly implicated the parasite derived RhopH complex in facilitating the increase in anion conductance [102].

A role for PKA in P. falciparum red cell invasion has also been reported [54]. AMA1, a protein released from the micro- nemes onto the merozoite surface and involved in tight junction formation, was shown to be phosphorylated at Ser540 in the cytoplasmic tail. Pharmacological and genetic evidence indicated that this event was carried out by the parasite PKA. The event was enhanced by addition of a cAMP analogue and reduced by PKA inhibition in parasite lysates. Substitution of AMA1 Ser540 with an alanine using a conditional transgenic approach led to a dramatic reduction in merozoite invasion [54]. A later report confirmed the PKA-dependent phosphorylation of AMA Ser540 and showed that phosphorylation of Ser540 is an initial step that is needed before subsequent phosphorylation of Thr<sub>513</sub> by the parasite glycoprotein synthase kinase 3 (GSK3; PF3D7_0312400) required for efficient invasion [103]. As mentioned above, a regulatory role for cAMP and PfAC<sub>b</sub> in calcium release and microneme secretion has been demonstrated using a low K<sup>+</sup> buffer system [5]. Free K<sup>+</sup> ions cannot readily cross membrane barriers and require specific ion channels to facilitate this process. In the light of this, the details of how this drop in K<sup>+</sup> concentration is sensed by the parasite and how it relates to a change in cAMP within the cell remain enigmatic. Evidence was presented suggesting that, once elevated, cAMP acts through two different pathways. First, experiments using in vitro kinase assays and a transgenic line overexpressing PfPKAr pointed to a role for PKA activity in regulating microneme release. Second, evidence derived from a pharmacological approach implicated an exchange protein directly activated by cAMP (Epac)-like molecule (PiEpac; PF3D7_1417400) in the calcium mobilization required for microneme release and merozoite invasion [5]. PiEpac contains consensic cyclic nucleotide-binding domains [6], but lacks other characteristic Epac signatures. In other organisms, Epac stimulates a CAMP-dependent calcium release by activating PLC via the small G protein rap1, but no orthologue has been identified to date in P. falciparum. Interestingly, it has been shown recently that culture-adapted P. falciparum isolates have a high propensity to undergo truncation of the PiEpac gene [22], suggesting that expression of the full-length gene represents a fitness cost in vitro and a non-essential role in blood-stage parasites.

A global phosphoproteome analysis carried out on P. falciparum schizonts revealed that 425 of the 2541 unique phosphoproteins conformed to one of the mammalian PKA substrate consensus sequences [R/K] R/K <i>+</i> S/T, suggesting that this kinase is likely to have a key role at this stage of the life cycle [104]. Bioinformatic approaches determined that a number of these potential PKA substrates are found in the ‘locomotion and entry into host’ Gene Ontology category which is required for merozoite invasion. In vitro phosphorylation of a synthetic peptide derived from Myosin A and recombinant CDPK1 and GAP43 (glideosome-associated protein 45; PF3D7_1222700) by bovine PKA suggested that these motor proteins could be PKA substrates in P. falciparum schizonts. The analysis also revealed phosphorylation in schizonts of several proteins involved in phosphoinositide metabolism, suggesting a key role for phosphatidylinositol signalling at this stage of the life cycle [104]. The authors point out that four phosphatidylinositol kinases
are phosphorylated at sites resembling consensus PKA sites; however, consensus PKA and PKG sites are very similar [105], and previous work on P. berghei ookinetes has suggested that phosphatidylinositol kinases are phosphorylated in a PKG-dependent manner [20]. The question of whether this phosphorylation is carried out directly by PKG or indirectly acting via downstream PKA phosphorylation will need further work.

A role for cAMP and PKA has also been found in gametocyte development. It is thought that the non-deformable nature of immature P. falciparum gametocytes allows them to sequester in bone marrow tissue to prevent splenic clearance [106–109]. However, to transmit to mosquitoes, mature gametocytes must re-enter the circulation and evidence suggests that to do this they need to become deformable. Using a microbead separation system designed to emulate splenic filtration, agents which promote cAMP signalling (such as cAMP analogues, AC activators and PDE inhibitors) through PKA were found to cause gametocytes to become more rigid (and retard their passage through the micro bead matrix), whereas reduced signalling stimulated by, for example, PKA inhibitors (but not a PKG inhibitor) caused the gametocytes to become more deformable [16]. In a parallel genetic approach in the same study, a line that overexpresses the regulatory subunit of PfPDE showed higher deformability in immature gametocytes. In addition, gametocytes from a PfPDE knockout line displayed a twofold increase in cAMP levels and a concomitant decrease in deformability of mature stages. Whether this effect on gametocyte cAMP levels is directly or indirectly attributed to PfPDE remains to be established, however consensus PKA and PKG sites are very similar [105], whether this is via host PKA or by PfPKA remains to be answered. A comprehensive research effort will be required to determine the biological significance of the individual phosphorylation events driven by PKA and PKG. It will also be fascinating to discover how these two pathways interact to coordinate merozoite egress and invasion. Another important question remains regarding how specificity is maintained when PKA and PKG have such similar consensus substrate sequences. How is it that all proteins having this basic amino acid motif are not haphazardly phosphorylated by either kinase because they are both expressed in late schizonts and merozoites? However, we do not know whether peaks of cAMP and cGMP coincide temporally within schizonts and free merozoites. It is also likely that, when active, the two kinases are located in distinct cellular compartments and/or are tethered within distinct protein complexes that bring them into the proximity of their respective substrates. The revelation that enzymatic components of the pathways play such pivotal roles in life-cycle progression has provided important target validation data for drug discovery projects. In partnership with medicinal chemists, it is possible that effective future antimalarial drugs will be developed that selectively target cAMP and cGMP signalling in malaria parasites.

6. Conclusion

In recent years, important steps have been made in determining the function of cyclic nucleotide signalling in malaria parasites, but recent advances in our genetic tools will ensure that we will have a more complete picture in the next 5–10 years. One of the big remaining questions is what operates upstream of cyclic nucleotide signalling in malaria parasites; only in gametogenesis do we have an answer to this. A comprehensive research effort will be required to determine the biological significance of the individual phosphorylation events driven by PKA and PKG. We declare we have no competing interests.

Data accessibility. This article has no additional data.

Funding. We are grateful to the Wellcome Trust (grant ref: 106240/Z/14/Z) for supporting this work.

References

1. World Health Organization. 2016 World malaria report 2016. Geneva, Switzerland: World Health Organization.
2. Burrows JN, Duparc S, Gutteridge WE, Hoof van Huijpsduijnen R, Kaszuboka W, Macintyre F, Mazzuri S, Möhrle JJ, Wells TNC. 2017 New developments in anti-malarial target candidate and product profiles. Malar. J. 16, 26. (doi:10.1186/s12936-016-1675-x)
3. Schaap P. 2016 Evolution of developmental signalling in dictyostelid social amoebas. Curr. Opin. Genet. Dev. 39, 29–34. (doi:10.1016/j.gde.2016.05.014)
4. Linder JU, Engel P, Reimer A, Krüger T, Plattner H, Schultz A, Schultz JE. 1999 Guanylyl cyclases with the topology of mammalian adenyly cyclases and
14. Wentzinger L, Bopp S, Tenor H, Klar J, Brun R, Beck HP, Scheck T. 2008 Cyclic nucleotide-specific phosphodiesterases of Plasmodium falciparum: PPDDeA1pha, a non-essential cGMP-specific PDE that is an integral membrane protein. Int. J. Parasitol. 38, 1625 – 1637. (doi:10.1016/j.ijpara.2008.05.016)

15. Lakshmanan V, Fishbaugh ME, Morrison B, Baldwin M, Macaruly M, Vaughan AM, Mikolajczak SA, Kappe SH. L. 2015 Cyclic GMP balance is critical for malaria parasite transmission from the mosquito to the mammalian host. MBio 6, e02330. (doi:10.1128/mBio.02330-14)

16. Ramdani G et al. 2015 cAMP-signalling regulates gametocyte-infected erythrocyte deformability required for malaria parasite transmission. PLoS Pathog. 11, e1004815. (doi:10.1371/journal.ppat.1004815)

17. Taylor HM, McRobert L, Grainger M, Sicard A, Dluozewski AR, Hopp CS, Holder AA, Baker DA. 2010 The malaria parasite cyclic GMP-dependent protein kinase plays a central role in blood-stage schizogony. Eukaryot. Cell 9, 37 – 45. (doi:10.1128/EC.00186-09)

18. McRobert L, Taylor CJ, Deng W, Fivelman QL, Cummings RM, Polley SD, Billker O, Baker DA. 2008 Gametogenesis in malaria parasites is mediated by the cGMP-dependent protein kinase. PLoS Biol. 6, e139. (doi:10.1371/journal.pbio.0060139)

19. Ganter M et al. 2017 Plasmodium falciparum CRK4 directs continuous rounds of DNA replication during schizogony. Nat. Microbiol. 2, 17017. (doi:10.1038/nm-radiobiology.2017.17)

20. Brochet M et al. 2014 Phosphonositolide metabolism links cGMP-dependent protein kinase G to essential Ca2+ signals at key decision points in the life cycle of malaria parasites. PLoS Biol. 12, e1001806. (doi:10.1371/journal.pbiol.001806)

21. Faale A, Gombe A, Amaladoss A, Carvalho T, Mendr R, Bhattacharjee P. 2010 Role of Plasmodium berghei cGMP-dependent protein kinase in late liver stage development. J. Biol. Chem. 285, 3282 – 3288. (doi:10.1074/jbc.M107.036376)

22. Claessens A, Affara M, Assifu SA, Kwiatkowski DP, Conway DJ. 2017 Culture adaptation of malaria parasites selects for convergent loss-of-function mutants. Sci. Rep. 7, 41303. (doi:10.1038/srep41303)

23. Read LK, Mikkelson RB. 1991 Plasmodium falciparum-infected erythrocytes contain an adenylate cyclase with properties which differ fromthose of the host enzyme. Mol. Biochem. Parasitol. 43, 109 – 119. (doi:10.1016/0166-6851(91)9032-2)

24. Baker DA, Kelly JM. 2004 Purine nucleotide cyclases in the malaria parasite. Trends Parasitol. 20, 227 – 232. (doi:10.1016/j.pt.2004.02.007)

25. Muhia DK, Sawaes CA, Eckstein-Ludwig U, Sasan, P, Polley SD, Kelly JM, Schap P, Krishna S, Baker DA. 2003 Multiple splice variants encode a novel adenyl cyclase of Plasmodium falciparum. Mol. Biochem. Parasitol. 143, 109 – 119. (doi:10.1016/S0166-6851(03)00257-2)

26. Weber JH, Vishnyakov A, Hambach K, Schultz A, Billker O. 2004 Initiation of Plasmodium falciparum development. PLoS Pathog. 1, 1000746. (doi:10.1371/journal.ppat.1000746)

27. Carucci DJ et al. 2000 Guanylyl cyclase activity associated with putative bifunctional integral membrane proteins in Plasmodium falciparum. J. Biol. Chem. 275, 22147 – 22156. (doi:10.1074/jbc.M00121200)

28. Kawamoto F, Abe-Blocco R, Fleck SL, Kawamoto Y, Sinden RE. 1999 Possible roles of Ca2+/ and cGMP as mediators of the exitflagellation of Plasmodium berghei and Plasmodium falciparum. Mol. Biochem. Parasitol. 42, 101 – 108. (doi:10.1016/0166-6851(99)01117-5)

29. Kawamoto F, Fujioka H, Murakami K, Safufridin HM, Ishikawa T, Hidaka H. 1993 The roles of Ca2+ /cGMP-signalling pathways in gametogenesis of a rodent malaria parasite, Plasmodium berghei. Eukaryot. Cell 2, 101 – 107. (doi:10.1242/jcs.177386)

30. Martin SK, Miller LH, Nijhout MM, Carter R. 1978 Plasmodium gallinaceum: induction of male gametocyte exflagellation by phosphodiesterase inhibitors. Exp. Parasitol. 44, 239 – 242. (doi:10.1016/0014-4894(78)90104-2)
40. Garcia GE, Wirtz RA, Barr JR, Woolfitt A, Rosenberg R. 1998 Xanthurenic acid induces gametogenesis in Plasmodium, the malaria parasite. J. Biol. Chem. 273, 12 003 – 12 005. (doi:10.1074/jbc.273.20.12003)

41. Billker O, Lindo V, Panico M, Etienne AE, Paxton T, Dell A, Rogers M, Sinden RE, Morris HR. 1998 Identification of xanthurenic acid as the putative inducer of malaria development in the mosquito. Nature 392, 289 – 292. (doi:10.1038/32667)

42. Muhia DK, Swales CA, Deng W, Kelly JM, Baker DA. 2001 The gametocyte-activating factor xanthurenic acid stimulates an increase in membrane-associated guanylyl cyclase activity in the human malaria parasite Plasmodium falciparum. Mol. Microbiol. 42, 553 – 560. (doi:10.1046/j.1365-2958.2001.02665.x)

43. Maurice DH, Ke H, Ahmad F, Wang Y, Chung J, Manganiello VC. 2014 Advances in targeting cyclic nucleotide phosphodiesterases: considerations for antimalarial drug design. J. Comput. Aided Mol. Des 28, 753 – 762. (doi:10.1007/s10822-011-9458-5)

44. Diaz CA, Allocco J, Powles MA, Yeung L, Donald RGK, Anderson JW, Liberator PA. 2006 Characterization of Plasmodium falciparum cGMP-dependent protein kinase (PKPk): antiparasitic activity of a PKG inhibitor. Mol. Biochem. Parasitol. 146, 78 – 88. (doi:10.1016/j.molbiopara.2005.10.020)

45. Baker DA et al. 2017 A potent series targeting the malarial cGMP-dependent protein kinase clears infection and blocks transmission. Nat. Commun. 8, 430. (doi:10.1038/s41467-017-00572-x)

46. Lin CS, Lau A, Tu R, Lue TF. 2000 Expression of the human parasite Plasmodium falciparum cGMP-specific phosphodiesterase (PDE5) in human penile cavernosum. Biochem. Biophys. Res. Commun. 268, 628 – 635. (doi:10.1006/bbrc.2000.2187)

47. Collins CR, Hackett F, Strath M, Penzo M, Withers-Martinez C, Baker DA, Blackman MJ. 2013 Malaria parasite cGMP-dependent protein kinase regulates blood stage merozoite secretion organelle discharge and egress. PLoS Pathog. 9, e1003344. (doi:10.1371/journal.ppat.1003344)

48. Ballard SA, Gimpell CJ, Tang K, Turner LA, Price ME, Naylor AM. 1998 Effects of sildenafil on the relaxation of human corpus cavernosum tissue in vitro and on the activities of cyclic nucleotide phosphodiesterase isozymes. J. Urol. 159, 2164 – 2171. (doi:10.1016/S0022-5347(01)63299-3)

49. Howard BL et al. 2015 Identification of potent phosphodiesterase inhibitors that demonstrate cyclic nucleotide-dependent functions in apicomplexan parasites. ACS Chem. Biol. 10, 1145 – 1154. (doi:10.1021/cb501004q)

50. Befekat T et al. 1992 Phosphodiesterase inhibition by enoximone in preparations from nonfailing and failing human hearts. Arzneimittel-Forschung 42, 437 – 445.

51. Syin C et al. 2001 The H89 cAMP-dependent protein kinase inhibitor blocks Plasmodium falciparum development in infected erythrocytes. Eur. J. Biochem. 268, 4842 – 4849. (doi:10.1046/j.1432-1327.2001.02403.x)

52. Murray AJ. 2008 Pharmacological PKA inhibition: all may not be what it seems. Sci. Signal. 1, re4. (doi:10.1126/scisignal.212re4)

53. Leykau K, Treek M, Gilson PB, Nebi T, Braille T, Cowman AF, Gilberger TW, Crabb BS. 2010 Protein kinase a dependent phosphorylation of apical membrane antigen 1 plays an important role in erythrocyte invasion by the malaria parasite. PLoS Pathog. 6, e1000941. (doi:10.1371/journal.ppat.1000941)

54. Hess KC et al. 2005 The ’soluble’ adenylyl cyclase in sperm mediates multiple signalling events required for fertilization. Dev. Cell 9, 249 – 259. (doi:10.1016/j.devcel.2005.06.007)

55. Seaman KB. 1984 Chapter 29. forosokin and adenylyl cyclase: new opportunities in drug design. Annu. Rev. Med. Chem. 19, 293 – 302. (doi:10.1006/jrme.1990.0070)

56. Beghyn TB, Charton J, Leroux F, Laconte G, Bourin A, Cos P, Maes L, Deprez B. 2011 Drug to genome to drug: discovery of new antimalarial compounds. J. Med. Chem. 54, 3222 – 3240. (doi:10.1021/jm2014617)

57. Kim JJ et al. 2015 Crystal structures of the carboxyl cGMP binding domain of the Plasmodium falciparum cGMP-dependent protein kinase reveal a novel capping triad crucial for merozoite egress. Cell 165, 165 – 175. (doi:10.1016/j.cell.2016.05.007)

58. Hopp CS, Flueck C, Solyakov L, Tobin A, Baker DA. 2002 The role of a parasite-specific allosteric site in phosphodiesterase isozymes. J. Biol. Chem. 277, 22003 – 22010. (doi:10.1074/jbc.M201389320)

59. Donald RK, Zhong T, Wiersma H, Nare B, Yao D, Lee A, Allocco J, Liberator PA. 2006 Anticoccidial protein kinase inhibitors: identification of protein kinase targets secondary to cGMP-dependent protein kinase. Mol. Biochem. Parasitol. 149, 86 – 98. (doi:10.1016/j.molbiopara.2006.05.003)

60. Deng W, Baker DA. 2002 A novel cyclic GMP-dependent protein kinase is expressed in the ring stage of the Plasmodium falciparum life cycle. Mol. Microbiol. 44, 1141 – 1151. (doi:10.1046/j.1365-2958.2002.02948.x)

61. Wiersma HI, Galuska SE, Tomley FM, Sibley LD, Liberator PA, Donald RGK. 2004 A role for cyclodin cGMP-dependent protein kinase in motility and invasion. Int. J. Parasitol. 34, 369 – 380. (doi:10.1016/j.ijpara.2003.11.019)

62. Billker O, Dechamps S, Tewari R, Weng G, Franke-Fayard B, Brinkmann V. 2004 Calcium and a calcium-dependent protein kinase regulate gamete formation and mosquito transmission in a malaria parasite. Cell 117, 503 – 514. (doi:10.1016/S0002-8474(04)00490-0)

63. Fang H, Klages N, Baechler B, Hillner E, Yu L, Pardo M, Choudhary J, Brochet M. 2017 Multiple short windows of calcium-dependent protein kinase 4 activity coordinate distinct cell cycle events during Plasmodium gametogenesis. eLife 6, e26524. (doi:10.7554/eLife.26524)

64. Bansal A, Molina-Cruz A, Bzostowski J, Mu J, Miller LH. 2017 Plasmodium falciparum calcium-dependent protein kinase 2 is critical for male gametocyte egress but not essential for asexual proliferation. MolBio 18, 7. (doi:10.1128/mbio.01656-17)

65. Yeoh S et al. 2007 Subcellular discharge of a serine protease mediates release of invasive malaria parasites from host erythrocytes. Cell 131, 1072 – 1083. (doi:10.1016/j.cell.2007.10.049)

66. Dvorin JD et al. 2010 A plant-like kinase in Plasmodium falciparum regulates parasite egress from erythrocytes. Science 328, 910 – 912. (doi:10.1126/science.1188191)

67. Hale VL et al. 2017 Parasitophorous vacuole poration precedes its rupture and rapid host erythrocyte cytoskeleton collapse in Plasmodium falciparum egress. Proc. Natl Acad. Sci. USA 114, 3439 – 3444. (doi:10.1073/pnas.1619441114)

68. Nagamura K, Hicks LM, Fux B, Bressler F, Chini EN, Sibley LD. 2008 Abscisic acid controls calcium-
dependent egress and development in Toxoplasma gondii. *Nature* **451**, 207–210. (doi:10.1038/nature06478)

77. Bansal A, Ojo KK, Mu J, Malay DJ, Van Voorhis WC, Miller LH. 2016 Reduced activity of mutant calcium-dependent protein kinase 1 is compensated in *Plasmodium falciparum* through the action of protein kinase G. *Mbio* **7**, 16. (doi:10.1128/mBio.02011-16)

78. Kumar P, Tripathi A, Ranjan R, Halbert J, Gilberger T, Doering C, Sharma P. 2014 Regulation of *Plasmodium falciparum* development by calcium-dependent protein kinase 7 (PfPKC7). *J. Biol. Chem.* **289**, 20 386 – 20 395. (doi:10.1074/jbc.M114.561760)

79. Moon RW, Hall J, Rangkuti F, Ho YS, Almond N, et al. 2013 Adaptation of the genetically tractable malaria pathogen *Plasmodium knowlesi* to continuous culture in human erythrocytes. *Proc. Natl Acad. Sci. USA* **110**, 531 – 536. (doi:10.1073/pnas.1216457110)

80. Leber W, Skippen A, Rangkuti F, Ho YS, Almond N, Mitchell GH, Pain A, Holder AA, Blackman MJ. 2013 Assembly of allosteric macromolecular switches: lessons from *Plasmodium falciparum*. *Mol. Biochem. Parasitol.* **39**, 645 – 653. (doi:10.1016/j.jpara.2008.11.015)

81. Taylor SS, Blouz R, Zhang P, Kornev AP. 2012 Assembly of allosteric macromolecular switches: lessons from *Plasmodium falciparum*. *Mol. Biochem. Parasitol.* **192** – 198. (doi:10.1016/j.molbiopara.2008.03.011)

82. Li J, Cox LS. 2000 Isolation and characterisation of a cAMP-dependent PKA and anion channel conductance. *Parasitol. Res.* **86**, 192 – 198. (doi:10.1007/s004360001015)

83. Merckx A, Nivez M-P, Bouyer G, Alano P, Langsley G, Holder AA, Langsley G, Alano P. 2012 A switch in infected erythrocyte deformability enabling malaria parasite egress. *Blood Cells Mol. Dis.* **46**, 192 – 198. (doi:10.1016/j.bcmd.2004.01.008)

84. Choudhury A, Decherf G, Egeé S, Staines HM, Ellory JC, Thomas S, Doerig C, Sharma P. 2014 Regulation of parasite deformability of *Plasmodium falciparum* infected human red blood cells. *Blood Cells Mol. Dis.* **46**, 192 – 198. (doi:10.1016/j.bcmd.2004.01.008)

85. Inselburg J. 1983 Stage-specific inhibitory effect of melatonin and other indoleamines on the in vitro development of *Plasmodium falciparum*. *J. Pineal Res.* **3**, 91 – 99. (doi:10.1002/jpe.480030208)

86. Bandjé K et al. 2016 Characterization of an A-kinase anchoring protein-like suggests an alternative way of PKA anchoring in *Plasmodium falciparum*. *Malar. J.* **15**, 248. (doi:10.1186/s12936-016-1275-9)

87. Rached FB et al. 2012 Construction of a *Plasmodium falciparum* Rab-interactome identifies CTK and PKA as Rab-effector kinases in malaria parasites. *Biol. Cell* **104**, 34 – 47. (doi:10.1111/boc.20100081)

88. Holder AA, Holder AA, Langsley G, Alano P. 2013 Characterization of an A-kinase anchoring protein-like suggests an alternative way of PKA anchoring in *Plasmodium falciparum*. *Malar. J.* **15**, 248. (doi:10.1186/s12936-016-1275-9)

89. Haidar M, Ramdani G, Kennedy EJ, Langsley G. 2016 PKA and apicomplexan parasite diseases. *Horm. Metab. Res.* **49**, 296 – 300. (doi:10.1055/s-0042-17847)

90. Toto D, Schreck MA, Desai SA. 2017 An essential dual-function complex mediates erythrocyte invasion and channel-mediated nutrient uptake in malaria parasites. *elife* **6**, e23485. (doi:10.7554/ elife.23485)

91. Prinz B et al. 2016 Hierarchical phosphorylation of apical membrane antigen 1 is required for efficient red blood cell invasion by malaria parasites. *Sci. Rep.* **6**, 24479. (doi:10.1038/srep34479)

92. Francis SH, Busch JL, Corbin JD, Sibley DL. 2010 cGMP-dependent protein kinases and cGMP phosphodiesterases in nitric oxide and cGMP action. *Pharmacol. Rev.* **62**, 532 – 563. (doi:10.1124/pr.110.002907)

93. Smalley ME, Abdalla S, Brown J. 1981 The distribution of *Plasmodium falciparum* in the peripheral blood and bone marrow of Gambian children. *Trans. R. Soc. Trop. Med. Hyg.* **75**, 103 – 105. (doi:10.1016/S0035-9203(81)80019-5)

94. Radoson E, Green JL, Camarda G, Talabani H, Holder AA, Langsley G, Alano P. 2012 The *Plasmodium falciparum* schizont phosphoproteome reveals extensive phosphoalstatidinostol and cAMP-protein kinase A signalling. *J. Proteome Res.* **11**, 3523 – 3537. (doi:10.1021/pr300557m)

95. Decherf G, Egeé S, Staines HM, Ellory JC, Thomas S, Doerig C, Sharma P. 2014 Regulation of parasite deformability of *Plasmodium falciparum* infected human red blood cells. *Blood Cells Mol. Dis.* **46**, 192 – 198. (doi:10.1016/j.bcmd.2004.01.008)

96. Rached FB et al. 2012 Construction of a *Plasmodium falciparum* Rab-interactome identifies CTK and PKA as Rab-effector kinases in malaria parasites. *Biol. Cell* **104**, 34 – 47. (doi:10.1111/boc.20100081)

97. Holder AA, Holder AA, Langsley G, Alano P. 2013 Characterization of an A-kinase anchoring protein-like suggests an alternative way of PKA anchoring in *Plasmodium falciparum*. *Malar. J.* **15**, 248. (doi:10.1186/s12936-016-1275-9)

98. Haidar M, Ramdani G, Kennedy EJ, Langsley G. 2016 PKA and apicomplexan parasite diseases. *Horm. Metab. Res.* **49**, 296 – 300. (doi:10.1055/s-0042-17847)

99. Toto D, Schreck MA, Desai SA. 2017 An essential dual-function complex mediates erythrocyte invasion and channel-mediated nutrient uptake in malaria parasites. *elife* **6**, e23485. (doi:10.7554/ elife.23485)

100. Francis SH, Busch JL, Corbin JD, Sibley DL. 2010 cGMP-dependent protein kinases and cGMP phosphodiesterases in nitric oxide and cGMP action. *Pharmacol. Rev.* **62**, 532 – 563. (doi:10.1124/pr.110.002907)

101. Smalley ME, Abdalla S, Brown J. 1981 The distribution of *Plasmodium falciparum* in the peripheral blood and bone marrow of Gambian children. *Trans. R. Soc. Trop. Med. Hyg.* **75**, 103 – 105. (doi:10.1016/S0035-9203(81)80019-5)

102. Aingaran M et al. 2012 Host cell deformability is linked to transmission in the human malaria parasite *Plasmodium falciparum*. *Cell. Microbiol.* **14**, 983 – 993. (doi:10.1111/j.1462-5822.2012.01786.x)

103. Joice R et al. 2014 *Plasmodium falciparum* transmission stages accumulate in the human bone marrow. *Sci Transl. Med.* **6**, 244re5. (doi:10.1126/scitranslmed.3008882)

104. Naissett B et al. 2016 *Plasmodium falciparum* STEVOR phosphorylation regulates host erythrocyte deformability enabling malaria parasite transmission. *Blood* **127**, e42 – e53. (doi:10.1182/ blood-2016-01-690776)