

1 ***Plasmodium falciparum* CRK4 directs continuous rounds of DNA**
2 **replication during schizogony**

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30 *Plasmodium* parasites, the causative agents of malaria, have evolved a unique cell
31 division cycle in the clinically relevant asexual blood-stage of infection¹. DNA
32 replication commences approximately halfway through the intracellular development
33 following invasion and parasite growth. The schizont stage is associated with multiple
34 rounds of DNA replication and nuclear division without cytokinesis resulting in a
35 multinucleated cell. Nuclei divide asynchronously through schizogony, with only the
36 final round of DNA replication and segregation being synchronous and coordinated
37 with daughter cell assembly^{2,3}. However, the control mechanisms for this divergent
38 mode of replication are unknown. Here we show that the *Plasmodium*-specific kinase
39 *PfCRK4* is a key cell cycle regulator that orchestrates the multiple rounds of DNA
40 replication throughout schizogony in *P. falciparum*. *PfCRK4* depletion led to a complete
41 block in nuclear division and profoundly inhibited DNA replication. Quantitative
42 phosphoproteomic profiling identified a set of *PfCRK4*-regulated phosphoproteins with
43 greatest functional similarity to CDK2 substrates, particularly proteins involved in
44 origin of replication firing. *PfCRK4* was required for the initial and subsequent rounds
45 of DNA replication during schizogony, and in addition was essential for development in
46 the mosquito vector. Our results identified an essential S phase promoting factor of the
47 unconventional *P. falciparum* cell cycle. *PfCRK4* is required for both a prolonged
48 period of the intraerythrocytic blood-stage of malaria infection, as well as for
49 transmission, revealing a broad window for *PfCRK4*-targeted chemotherapeutics.

50 Malaria parasites proliferate through schizogony in the blood-stage of infection: a
51 series of rapid rounds of DNA replication and nuclear division produces a syncytial cell with
52 approximately 20 nuclei. In contrast to the synchronous nuclear division observed in other
53 multinucleated cells, such as the early *Drosophila* embryo⁴, *P. falciparum* nuclei divide
54 asynchronously during the blood-stage despite sharing the same cytoplasm^{2,3}, suggesting that

55 cell cycle progression is not governed by diffusible cytoplasmic factors. Knockout screens
56 have identified non-essential blood-stage *Plasmodium* kinases, and suggested those that could
57 be essential for the regulation of schizogony^{5,6}. The molecular mechanisms regulating this
58 diverged mode of replication are largely unknown yet the recent development of conditional
59 gene expression technologies now allows to unequivocally demonstrate gene essentiality in
60 *P. falciparum*⁷⁻⁹.

61 To directly identify critical regulators of schizogony and simultaneously determine
62 protein function, we adopted the destabilization domain (DD) conditional knockdown
63 approach^{7,10,11}. We generated endogenous DD fusions of 23 schizont-stage kinases in the *P.*
64 *falciparum* D10 strain and screened for vulnerability to destabilization by DD (Fig. 1a, b,
65 Supplementary Information Fig. 1a). Using this approach, we found that two kinases, the
66 cGMP-dependent protein kinase (*PfPKG*) and the *cdc2*-related protein kinase 4 (*PfCRK4*),
67 had profound proliferation defects in absence of Shield-1 and showed dose-dependency (Fig.
68 1c, Supplementary Information Fig. 1b, c). Chemical inhibition of *PfPKG* has previously
69 established its essential role in parasite egress from erythrocytes at the end of schizogony^{12,13},
70 now confirmed by conditional destabilization.

71 As previously observed^{14,15}, we found little correlation between the level of protein
72 knockdown and the ability to reveal essentiality of other kinases previously thought to be
73 essential⁶ using the DD approach (Fig. 1d, Supplementary Information Fig. 1d, 9c, d). This
74 could be due to insufficient destabilization of these proteins or because they are required at
75 inherently different levels for asexual proliferation.

76 The biological function of *PfCRK4* is unknown and we confirmed its essentiality in a
77 different parental line, *P. falciparum* P2G12¹⁶, which also produces gametocytes
78 (Supplementary Information Fig. 2). *PfCRK4* is a member of an Apicomplexa-specific kinase

79 subfamily related to cyclin-dependent kinases (CDK) (Supplementary Information Fig. 3,
80 Supplementary Data table 1)^{17,18}. In many organisms CDKs in complex with cyclins regulate
81 key steps of the cell cycle and other cellular functions^{19,20}. Compared to human CDK2, the
82 kinase domain of *PfCRK4* possesses multiple sequence inserts of unknown function (Fig. 2a,
83 Supplementary Information Fig. 4).

84 *PfCRK4* is localized to the nucleus of late trophozoites and schizonts, with the signal
85 greatly diminished in segmented schizonts that had undergone cytokinesis (Fig. 2b).
86 Following conditional depletion of *PfCRK4* (Fig. 1d) parasites arrested at the trophozoite-to-
87 schizont transition (Fig. 2c). At ≥ 40 hours post invasion (hpi), parasites [+] Shield-1 (*i.e.*,
88 wild type *PfCRK4* levels) segmented into daughter cells, containing nuclei and rhoptries
89 (apical organelles required for merozoite invasion), which are characteristic of mature
90 schizonts (Fig. 2d, top). In contrast, parasites cultured [-] Shield-1 (*i.e.*, *PfCRK4* is depleted)
91 showed no nuclear division or apical organelle biogenesis (Fig. 2d, bottom, Supplementary
92 Information Fig. 5a).

93 Analysis of nuclei stained with fluorescent DNA-specific dyes confirmed that the
94 nuclei did not divide and revealed a substantially distorted nuclear morphology (Fig. 2e,
95 Supplementary Information Fig. 5b). Hemispindle structures were evident in *PfCRK4*-
96 depleted cells (Fig. 2f; Supplementary Information Fig. 5c, d); however, they were greatly
97 enlarged relative to spindles in wild type parasites and might account for the nuclear
98 distortion. Concordantly, division of the centriolar plaque^{3,21}, the parasite's microtubule
99 organizing centre, was diminished (Fig. 2g). To ascertain whether *PfCRK4* affects DNA
100 replication, we quantified the parasite's DNA content by flow cytometry^{22,23}. While in wild
101 type parasites DNA replication commences at 29-32 hpi, *i.e.*, parasites with C-values >1
102 appear, we found DNA replication to be profoundly inhibited in *PfCRK4*-depleted parasites

103 (Fig. 2h). In contrast, the development of the mitochondria and apicoplast organelles was
104 unaffected following depletion of *PfCRK4* (Supplementary Information Fig. 5e, f). The
105 *PfCRK4*-dependent block was completely reversible up to 38 hpi by addition of Shield-1.
106 However, reversion at 48 hpi led to very poor recovery (Fig. 2i), identifying a window within
107 which *PfCRK4* depletion is cytostatic before becoming cytotoxic.

108 To elucidate processes regulated by *PfCRK4*, we assessed changes in the
109 phosphoproteome of *PfCRK4*-depleted parasites at two time points (Fig. 3a): at 29 hpi, when
110 *PfCRK4* is already expressed at the onset of DNA replication (Fig. 2b, h), and also at 37 hpi,
111 where in wild type parasites DNA replication is evident and when the *PfCRK4*-dependent
112 arrest was still reversible (Fig. 2h, i) (Supplementary Information Fig. 6; Supplementary Data
113 table 2-5). Large changes occur in the phosphoproteome at 29 hpi indicating that *PfCRK4* is
114 active when the first round of DNA replication commences (Fig. 2h). We used k-means
115 clustering to identify the phosphosites most affected by the *PfCRK4* depletion, which showed
116 a reduction of about 2-fold (Supplementary Information Fig. 6b). Applying a threshold based
117 on a ≥ 2 -fold decrease in phosphorylation and p -values < 0.05 , we identified 220 proteins (80
118 from 29 hpi, 215 from 37 hpi with 75 proteins shared), suggesting that these belong to a
119 *PfCRK4*-regulated set of phosphoproteins (Fig. 3b, Supplementary Data table 4). Western
120 blot analysis confirmed differential phosphorylation of histone H3 (serine 28) in a different
121 genetic background (Supplementary Information Fig. 6c). The most enriched phosphorylation
122 motif within these proteins was a previously undescribed $S\Phi xK$ motif (Φ , hydrophobic; x ,
123 any amino acid), with S being phosphorylated. The canonical CDK motif $S/TP^{24,25}$, with S or
124 T phosphorylated, was also enriched (Supplementary Information Fig. 6d, e).

125 To examine the functional significance of the *PfCRK4*-regulated set of proteins, we
126 used gene ontology (GO) term enrichment analysis (Fig. 3c). The most enriched biological

127 process GO term was ‘DNA replication’, and ‘nucleus’ was the most enriched cellular
128 component GO term. Strikingly, six of the seven identifiable *P. falciparum* homologues (or
129 subunits thereof) of the 16 *Saccharomyces cerevisiae* factors required for origin licensing and
130 firing *in vitro*²⁶ showed reduced phosphorylation in *PfCRK4* depleted parasites (Fig. 3c, box;
131 Supplementary Information Fig. 6f, Supplementary Data table 3, 4). GO terms of proteins
132 with a ≥ 2 -fold increase in phosphorylation (p -value < 0.05) were enriched for biosynthetic
133 and intracellular transport processes at 37 hpi, perhaps indicating secondary effects; however,
134 no statistically significant enrichment was observed at 29 hpi (Supplementary Information
135 Fig. 6g).

136 We next compared the GO terms of the *PfCRK4*-regulated set with GO terms of other
137 available kinase substrate sets—one *S. cerevisiae* kinase²⁷ and 19 human kinases²⁸
138 (www.phosphosite.org)—each with ≥ 30 confirmed substrates (Fig. 3d). The substrate set of
139 human CDK2, a major S phase promoting factor, showed the highest percentage of shared
140 annotations, suggesting that *PfCRK4* acts in a related fashion to promote S phase in *P.*
141 *falciparum*.

142 *PfCRK4* expression increases through schizogony (Fig. 2b) and its transcripts peak
143 the latest of all putative *Plasmodium* CDK-like kinases²⁹. We therefore hypothesized that
144 *PfCRK4* is crucial for all rounds of DNA replication seen in schizogony. When we depleted
145 *PfCRK4* at points later in schizogony, we detected no further increase in DNA content (Fig.
146 4a), premature termination of nuclear division (Fig. 4b), and parasites failed to proliferate
147 (Fig. 4c). Thus demonstrating that *PfCRK4* function is critical throughout schizogony.

148 To determine a role for *PfCRK4* in other life cycle stages, we used the P2G12-
149 *PfCRK4*-HA-DD line that produces high levels of gametocytes to analyse *PfCRK4* function
150 during parasite transmission to the *Anopheles gambiae* mosquito vector. While we detected

151 nuclear expression of *PfCRK4* in gametocytes (Fig. 4d, Supplementary Information Fig. 7a),
152 depletion during the latter portion of gametocyte development (*i.e.* from day 6 to 16 post-
153 induction) had no effect on gametocytemia nor on the male-to-female gametocyte ratio
154 (Supplementary Information Table 1). *PfCRK4* depletion also did not impair exflagellation
155 (Supplementary Information Fig. 7b, c, d). We observed a reduced *PfCRK4* signal in
156 ookinetes (Supplementary Information Fig. 7e), and the numbers of oocysts on *An. gambiae*
157 female midguts was greatly diminished upon *PfCRK4* depletion (Fig. 4e, Supplementary
158 Information Fig. 8). Oocysts derived from gametocyte cultures in the presence of *PfCRK4*
159 showed multiple nuclei, well-expanded organelles, and a well-developed capsule (Fig. 4f,
160 top). In contrast, infections with *PfCRK4*-depleted gametocytes resulted in necrotic parasites
161 reminiscent of dead ookinetes, the motile zygote of *Plasmodium* parasites (Fig. 4f, bottom;
162 Supplementary Information Fig. 8). This indicates that *PfCRK4* is required at some point
163 during ookinete formation and early oocyst development, and likely plays a role during DNA
164 replication in zygotes.

165 Our data provide evidence that *PfCRK4* is an essential protein kinase promoting S
166 phase through the initiation of multiple rounds of DNA replication during schizogony. While
167 phosphorylation is likely important for the regulation of *PfCRK4* (Supplementary
168 Information Fig. 3b, 4a), a potential cyclin-dependence is in question as the cyclin binding
169 domain is poorly conserved (Supplementary Information Fig. 4a)¹⁷. Interestingly,
170 homologues of Group I cyclins are not found in the *Plasmodium* genome, including cyclin E,
171 which is critical for G1 to S transition in other systems³⁰. Interventions targeting *PfCRK4*
172 function would be efficacious throughout the extended period of schizogony in the clinically
173 relevant blood-stage, as well as at the transmission stages, which taken together are attractive
174 features for future drug development.

175 **Supplementary Information** is linked to the online version of the paper at
176 www.nature.com/nmicrobiol/

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195 **Author contributions**

196 M.G. designed, performed, and interpreted much of the experimental work; J.M.G. analysed
197 the phosphoproteomic data and provided bioinformatics support; J.D.D., J.A.P., J.G.K.,
198 A.K.T., A.S.P., and I.C. designed and performed specific experimental work; J.Y.,

199 constructed plasmids; R.H.Y.J. provided bioinformatics support; B.E. performed Western
200 blots; D.A.B., R.R.D., and S.P.G. provided reagents and intellectual input into study design;
201 M.G., J.D.D., and M.T.D. conceived the study; M.G., J.M.G, and M.T.D wrote the
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203 **Author information**

204 The authors declare that they have no competing financial interests. Reprints and permissions
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207 **Methods**

208 **Reagents and oligonucleotide primers.** We purchased chemicals from Sigma-Aldrich
209 (unless otherwise noted), primers Integrated DNA Technologies (sequences are available
210 upon request), and restriction enzymes from New England Biolabs.

211 **Construction of plasmids.** The terminal 1-2 kb of PKB (PF3D7_1246900), TLK4
212 (PF3D7_0623800), and PKG (PF3D7_1436600) were sub-cloned into the 3' replacement
213 vector pJDD41⁷, generating HA-DD single crossover tagging plasmids. The terminal ~1 kb
214 of PF3D7_0420100 was amplified using an extended reverse primer, introducing two
215 additional HA-tags. Cloning into pJDD41 resulted in a 3xHA-DD single crossover tagging
216 plasmid. All other targeting fragments were sub-cloned into this 3xHA-DD plasmid
217 (sequences available upon request).

218 **Parasite culture, transfection and synchronization.** *P. falciparum* D10 was obtained from
219 the Walter and Eliza Hall Institute (Melbourne, Australia). *P. falciparum* P2G12 (clone of
220 3D7) was obtained from Harvard T.H. Chan School of Public Health (Boston, USA).
221 Parasites were cultured as previously described in RPMI-1640 media supplemented with

222 0.5% Albumax-II (Invitrogen), 50 mg / l hypoxanthine, 0.21% Sodium Bicarbonate, and 25
223 mM HEPES (EMD Biosciences)³¹. Human O+ erythrocytes (Research Blood Components)
224 were diluted to 2-4% haematocrit. Transgenic parasites were generated by electroporation of
225 synchronized ring-stage parasites as previously described³². Single homologous
226 recombination events were selected in presence of 500 nM Shield-1 by cycling on and off
227 WR99210 (Jacobus Pharmaceutical Company) and cloned by limiting dilution. Parasites
228 were synchronized by a combination of heparin and sorbitol treatments as previously
229 described³³. Unless otherwise noted, DD-tagged parasites were cultured in 250 nM Shield-1.
230 When desired, Shield-1 was removed by triple washes with excess of RPMI.

231 **Southern blot analysis.** Genomic DNA was harvested with QIAamp Blood Mini Kit
232 (Qiagen) and digested with the enzymes indicated in Supplementary Fig. 1a, resolved on
233 0.8% agarose gels, transferred to GeneScreen Plus (Perkin Elmer), and hybridized with
234 specific radiolabeled probes.

235 **May-Grünwald-Giemsa staining and imaging of blood-stage parasites.** Air-dried thin-
236 smear blood films were fixed and stained according to manufacturers instructions and imaged
237 on a Zeiss AxioCam microscope equipped with a 100X oil-immersion objective. Raw images
238 were analysed using ImageJ³⁴.

239 **Dose response experiments.** Dose response curves were generated as previously described
240 using SYBR® Green I³⁵. In brief, triplicate 2-fold Shield-1 dilution series were set up in 100
241 µl parasite cultures (0.075% parasitemia in 2% haematocrit). After two cycles, parasites were
242 lysed with 20 µl of 6x SYBR® green I lysis buffer (0.16% saponin; 20 mM Tris-HCl, 5 mM
243 EDTA, 1.6% Triton X 100, pH 7.4), supplemented with 1:1.000 SYBR® green I (from
244 10.000x stock, Thermo Fisher). Fluorescence intensity was measured on a SpectraMax M5
245 plate reader (Molecular Devices).

246 **Conditional destabilization screen of schizont-stage kinases.** We used the relative log₁₀
247 parasite multiplication rate (PMR) [-] over [+] Shield-1 as a measure of the effect of
248 conditional destabilization of schizont-stage kinases. Synchronized ring-stage parasites of all
249 *P. falciparum* kinase-DD fusion lines were seeded in triplicate at 0.05% parasitemia, 0.5%
250 haematocrit [+] or [-] Shield-1. Parasitemia was monitored by flow cytometry (see below)
251 and expressed as log₁₀. Linear regression analysis (Prism, GraphPad) allowed calculating the
252 PMR. The effect of protein knockdown ([-] Shield-1) relative to [+] Shield-1 was plotted as
253 log₁₀ PMR [-] Shield-1 over log₁₀ PMR [+] Shield-1. The error (standard deviation) for

254
$$f = \frac{A}{B}$$

255 was propagated as follows:

256
$$s_f = |f| \sqrt{\left(\frac{s_A}{A}\right)^2 + \left(\frac{s_B}{B}\right)^2}$$

257 Where A is the log₁₀ PMR [-] Shield-1, B is the log₁₀ PMR [+] Shield-1 and s_A and s_B are the
258 sample standard deviations in A and B, respectively³⁶.

259 **Parasitemia quantification by flow cytometry.** Parasitized erythrocytes were fixed as
260 previously described³⁷, stained with 1:2.000 SYBR® green I (Life Technologies) in PBS, and
261 analysed on a Miltenyi MACSQuant® flow cytometer. Data were analysed using FlowJo
262 (Tree Star) and Prism (GraphPad) software.

263 **Immunoblot analysis.** Synchronized parasites were harvested and uninfected erythrocytes
264 were removed using 0.05% saponin in PBS plus 1x cComplete™ protease inhibitor cocktail
265 (Roche). Upon washes, parasites were lysed in Laemmli sample buffer and proteins were
266 resolved by SDS-PAGE, transferred to nitrocellulose, and probed with the following primary
267 antibodies: anti-HA (1:1.000; Roche; clone 3F10), anti-*Pf*lactate dehydrogenase (LDH)

268 (1:2.000; gift of Michael T. Makler), anti-histone H3 (1:1.000; Abcam; ab1791), anti-pS28
269 histone H3 (1:1.000; Abcam; ab5169), anti-Pfaldolase (1:40.000, Abcam; ab38905). Signal
270 intensities were quantitated using ImagJ.

271 **Gametocyte assays and *An. gambiae* infections.** Gametocyte cultures were seeded at 0.5%
272 parasitemia in 5 ml RPMI with 10% serum, 4% haematocrit, and 1 μ M Shield-1 and kept in a
273 candle jar at 37 °C. From day six onwards parasites were cultured [+] and [-] Shield-1.
274 Estimates for the various stages and sexes were determined from a Giemsa-stained thin-
275 smear blood film. On days 15 to 18 of culture, parasites were harvested and male gametocyte
276 exflagellation was assessed by microscopy. For the infection of *An. gambiae* KEELE strain
277 females, cultures were washed and gametocytemia was adjusted to 0.3% in normal human
278 serum with 45% haematocrit. Subsequently, females were infected in a Standard Membrane
279 Feeding Assay as previously described³⁸.

280 **Oocyst quantification.** Midguts from infected *An. gambiae* females were dissected on days 7
281 or 8 post blood-feeding and subsequently the number of oocysts were enumerated following
282 0.1% mercurochrome staining and analysis by light microscopy (20X objective).

283 **DNA content analysis.** The DNA-content of parasites was determined by flow cytometry.
284 Parasitized erythrocytes were fixed in 4% paraformaldehyde and 0.0075% glutaraldehyde for
285 45 min at room temperature³⁷. To reduce RNA-derived background signal cells were
286 permeabilized with 0.1% Triton X-100, treated with 0.3 mg / ml RNase A for 20' at 37 °C,
287 and subsequently washed. Cells were stained with 1:2.000 SYBR® green I (Life
288 Technologies) in PBS, washed, and analysed on a Miltenyi MACSQuant® flow cytometer.
289 Data were analysed using FlowJo (Tree Star), Excel (Microsoft), and Prism (GraphPad)
290 software. Second cycle ring-stage parasites were omitted for DNA content analysis of 48 hpi
291 schizont-stage parasites.

292 **Immunofluorescence assays.** Immunofluorescence assays of blood-stage parasites was done
293 as previously described^{37,39}. In brief, parasitized erythrocytes were air-dried, methanol-
294 fixed³⁹, and analysed using rat anti-HA (1:100, Roche, clone 3F10), mouse anti-centrin
295 (1:500, Millipore, clone 20H5), mouse anti- α -tubulin (1:2.000, Sigma-Aldrich, clone B-5-1-
296 2), rabbit anti-GAP45 (1:2.000, gift of Julian C. Rayner), or rabbit anti-*Pf*EBA175 (1:500,
297 Malaria Research and Reference Reagent Resource Center, R3347) antibodies. Alternatively,
298 parasites were fixed in solution³⁷ for 20 min at room temperature, immobilized on poly-L-
299 lysine coated cover slips (Corning®), and analysed using mouse anti- α -tubulin (1:2.000,
300 Sigma-Aldrich, clone B-5-1-2) or rabbit anti-*P. falciparum* acyl carrier protein⁴⁰ (1:500, gift
301 of Sean T. Prigge) antibodies. Primary antibodies were detected using Alexa Fluor®
302 conjugated secondary antibodies (1:2.000, Thermo Fisher Scientific). Cells were mounted
303 with DAPI Fluoromount-G® (SouthernBiotech) or cells were stained with Hoechst 33342
304 (Thermo Fisher Scientific) and mounted Fluoromount-G® (SouthernBiotech), imaged on a
305 Nikon Eclipse TE300 microscope equipped with a Hamamatsu C10600 Orca R2 digital
306 camera, and analysed employing ImageJ software.

307 For immunofluorescence analysis of exflagellation, mature gametocytes were
308 resuspended in RPMI-1640 media supplemented with 20% human serum (AB+), 50 mg / l
309 hypoxanthine, 0.21% Sodium Bicarbonate, 25 mM HEPES (EMD Biosciences), and 50 μ M
310 xanthurenic acid at room temperature and pH 8 to induce exflagellation⁴¹. Following 20 min
311 incubation at room temperature, cells were washed, spotted on a glass slide, air-dried,
312 methanol-fixed, and further processed as described above.

313 For immunofluorescence analysis of ookinetes, female *An. gambiae* mosquitoes were
314 fed with late stage *Pf*CRK4-HA-DD gametocyte cultures, which were maintained [+] and [-]
315 Shield-1 from day 6 post-induction. Approximately 24 hours post-feeding, the blood bolus

316 from the mosquito midgut lumen was isolated. Bolus material was spotted on glass slides and
317 air-dried preceding immunofluorescence analysis.

318 Immunofluorescence analysis of *P. falciparum* late midgut oocysts was done as
319 previously described⁴². In brief, dissected midguts were fixed in 4% paraformaldehyde for 1h
320 at room temperature, washed, and permeabilized, and blocked using 0.2% Triton X-100 in
321 1% bovine serum albumin in PBS. Midgut oocysts were analysed using a mouse anti-
322 circumsporozoite (CS) protein antibody (15 µg/ml, gift of Photini Sinnis), an Alexa Flour®
323 594 secondary antibody (1:1.000), and DAPI nuclear stain, mounted on a glass slide in 10%
324 glycerol in PBS, and imaged on a Nikon Eclipse 90i microscope. Raw images were analysed
325 using Volocity software (Perkin-Elmer).

326 **JC-1 staining of *P. falciparum* mitochondria.** Parasite mitochondria were stained with the
327 cationic mitochondrial membrane potential sensor JC-1 (Life Technologies) as previously
328 described⁴³. In brief, 2 µM JC-1 in 37 °C warm RPMI was filtered using an Acrodisc®
329 Syringe Filter (0.2 µm HT Tuffryn® Membrane, Pall) to removed preformed JC-1
330 aggregates. Parasites were incubated with JC-1 containing media for 20 min at 37 °C,
331 washed, and imaged immediately on a Nikon Eclipse TE300 microscope with equipped a
332 Hamamatsu C10600 Orca R2 digital camera and analysed employing ImageJ software.

333 **Electron microscopy.** Synchronized blood-stage parasites were isolated by magnetic affinity
334 purification using a MACS LS column (Miltenyi Biotec), adjusted to ~50% parasitemia with
335 uninfected erythrocytes, and resuspended in 200 µl 3% BSA in PBS. Cells were fixed for 1h
336 at room temperature with an equal volume of 2.5% paraformaldehyde, 5.0% glutaraldehyde,
337 and 0.06% picric acid in 0.2 M cacodylate buffer. Cells were washed in cacodylate buffer,
338 post-fixed for 1 h with 1% osmium tetroxide (OsO₄), 1.5% potassium ferrocyanide
339 (K₄Fe(CN)₆), and washed in H₂O. Following 1h incubation in 1% aqueous uranyl acetate
340 solution, samples were washed in H₂O and subsequently dehydrated in grades of alcohol. The

341 samples were placed in propyleneoxide for 1 h and infiltrated in a 1:1 mixture of
342 propyleneoxide and TAAB Epon (Marivac) over night. Samples were embedded in TAAB
343 Epon and polymerized at 60 °C for 48 h. Ultrathin sections from a Reichert Ultracut-S
344 microtome were mounted on copper grids and stained with lead citrate. Samples were imaged
345 on a TecnaiG² Spirit BioTWIN with an AMT 2k CCD camera under 80 kV.

346 For *P. falciparum* oocyst thin-section transmission electron microscopy, infected
347 midguts were fixed at day 7 post infection in 2.5% glutaraldehyde (Electron Microscopy
348 Sciences) and processed as described⁴⁴. Stained sections were examined with a Philips
349 CM120 EM under 80 kV.

350 **Phosphoproteomic profiling.** *Cell culture and harvest.* Synchronized D10-*Pf*CRK4-HA-DD
351 parasites were cultured for 29 or 37 hours post invasion (hpi) [+] or [-] Shield-1. At 29 hpi
352 duplicate samples were harvested, at 37 hpi triplicate samples were harvested for each [+]
353 and [-] Shield-1. For each of the ten samples $\sim 1.25 \times 10^9$ parasites were isolated and
354 uninfected erythrocytes were removed using saponin buffer (0.05% saponin in PBS plus 1x
355 PhosSTOP (Roche), 1x cOmpleteTM protease inhibitor cocktail (Roche), and 1 mM
356 phenylmethanesulfonyl fluoride (PMSF)). Subsequently, cells were washed in saponin buffer
357 to further remove erythrocytes and erythrocyte debris, followed by washes with PBS plus 1x
358 PhosSTOP, 1x cOmpleteTM protease inhibitor cocktail, and 1 mM PMSF.

359 *Cell lysis and protein digestion.* Lysates were prepared as previously described^{45,46}. In
360 brief, cells were lysed in 8 M urea, 100 mM sodium chloride, 25 mM TRIS pH 8 plus 1x
361 PhosSTOP, 1x cOmpleteTM protease inhibitor cocktail, and 1 mM PMSF in PBS. The protein
362 concentration was estimated in the Bio-Rad Protein Assay (Bio-Rad). Disulfide bonds were
363 reduced with 5 mM tris(2-chloroethyl) phosphate (TCEP) for 30 min at 37 °C. Cysteines
364 were alkylated with 15 mM iodoacetamide for 30 min at room temperature in the dark,
365 followed by incubation with 5 mM DTT for 15 min at room temperature in the dark to

366 quench excess of iodoacetamide.

367 Chloroform-methanol precipitated proteins⁴⁷ were resuspended in 8 M urea, 50 mM
368 HEPES pH 8.5 and subsequently diluted to 1 M urea 50 mM HEPES pH 8.5 for digestion
369 with LysC protease (1:100 protease-to-protein ratio, 3 h at 37 °C) before addition of trypsin
370 (1:100 protease-to-protein ratio) and continued digest overnight at 37 °C. The reaction was
371 quenched with 1% formic acid, subjected to C18 solid phase extraction (Sep-Pak, Waters),
372 followed by vacuum-centrifugation.

373 *Isobaric labelling with tandem mass tags (TMT).* Peptides (200 µg) from each sample
374 were dissolved in 100 mM HEPES pH 8.5. Labelling with TMT reagents (Thermo Fisher
375 Scientific) was done according the manufactures instructions and as previously described⁴⁸.
376 The TMT-labelled samples were combined at a 1:1 ratio across all samples, vacuum
377 centrifuged to near dryness, and subjected to C18 solid phase extraction (Sep-Pak, Waters).

378 *Phosphopeptide enrichment.* Peptides were resuspended in 100 mM HEPES pH 8.5
379 and phosphopeptides were enriched using Titanosphere TiO₂ 5 µm particles (GL
380 Biosciences) as previously described^{46,48,49}. The flow-through was kept for proteome
381 analysis. Enriched TMT-labelled phosphopeptides and peptides were dried via vacuum
382 centrifugation.

383 *Off-line basic pH reversed-phase (BPRP) fractionation.* For protein level analysis, we
384 fractionated the pooled TMT-labelled peptide sample via BPRP HPLC as previously
385 described^{48,50}. Fractions were desalted, dried, and reconstituted in 5% acetonitrile, 5% formic
386 acid for LC-MS/MS processing.

387 For phosphopeptide analysis, we used the Pierce Off-line basic pH reversed-phase
388 (BPRP) fractionation kit (Thermo Scientific). Fractions collected included organic buffer
389 bumps with 5, 7.5, 10, 12.5, 15, 17.5, 20, 50, and 70% acetonitrile, in addition to the wash

390 and flow through fractions. These fractions were vacuum-centrifuged to near dryness,
391 desalted via StageTip, dried, and reconstituted in 5% acetonitrile, 5% formic acid for LC-
392 MS/MS processing.

393 *Liquid chromatography and tandem mass spectrometry.* Data were collected using an
394 Orbitrap Fusion mass spectrometer (Thermo Fisher Scientific) coupled to a Proxeon EASY-
395 nLC II liquid chromatography (LC) pump (Thermo Fisher Scientific). Peptides were
396 fractionated on a 75 μm inner diameter micro capillary column packed with ~0.5 cm of
397 Magic C4 resin (5 μm , 100 \AA , Michrom Bioresources) followed by ~35 cm of GP-18 resin
398 (1.8 μm , 200 \AA , Sepax). For each analysis, we loaded ~1 μg onto the column.

399 For total proteome analysis, peptides were separated using a 180 min gradient of 6-
400 23% acetonitrile in 0.125% formic acid (flow rate: ~600 nl / min). Each analysis used the
401 multi-notch MS3-based TMT method⁵¹. The scan sequence began with an MS1 spectrum
402 (Orbitrap analysis; resolution 120.000; mass range 400-1,400 m/z; automatic gain control
403 (AGC) target 2.0×10^5 ; maximum injection time 100 ms). Precursors for MS2/MS3 analysis
404 were selected using a TopSpeed 2 seconds method. MS2 analysis consisted of collision-
405 induced dissociation (quadrupole ion trap analysis; AGC 4×10^3 ; normalized collision
406 energy (NCE) 35; maximum injection time 150 ms). Following acquisition of each MS2
407 spectrum, we collected an MS3 spectrum using a recently described method in which
408 multiple MS2 fragment ions were captured in the MS3 precursor population using isolation
409 waveforms with multiple frequency notches⁵¹. MS3 precursors were fragmented by high-
410 energy collision-induced dissociation (HCD) and analysed (NCE 55; AGC 5×10^4 ;
411 maximum injection time 150 ms, resolution was 60.000 at 200 Th).

412 For phosphoproteome analysis, peptides were separated using a 120 min gradient of
413 3-23% acetonitrile in 0.125% formic acid (flow rate: ~575 nl / min). Each analysis used the
414 multi-notch MS3-based TMT method⁵¹. The scan sequence was identical to the proteome

415 scan, except the maximum injection time was 200 ms for MS2 analysis and 250 ms for MS3
416 analysis.

417 *Initial mass spectrometry data analysis.* Mass spectra were processed using a
418 SEQUEST-based in-house software pipeline⁵². Spectra were converted to mzXML using a
419 modified version of ReAdW.exe. Database searching included all entries from a combined
420 Uniprot human and *Plasmodium* database (February, 2015). This database was concatenated
421 with one composed of all protein sequences in the reversed order. Total protein level analysis
422 searches were performed using a 50 ppm precursor ion tolerance, product ion tolerance was
423 set to 0.9 Da. These wide mass tolerance windows were chosen to maximize sensitivity in
424 conjunction with SEQUEST searches and linear discriminant analysis^{52,53}. TMT tags on
425 lysine residues and peptide N termini (+229.163 Da) and carbamidomethylation of cysteine
426 residues (+57.021 Da) were set as static modifications, while oxidation of methionine
427 residues (+15.995 Da) was set as a variable modification for both total proteome and
428 phosphoproteome datasets. For phosphoprotein analysis, +79.966 Da was set as a variable
429 modification on serine, threonine, and tyrosine residues.

430 Peptide-spectral matches (PSMs) were adjusted to a 1% false discovery rate
431 (FDR)^{54,55}. PSM filtering used a linear discriminant analysis as previously described⁵², while
432 considering the following parameters: XCorr, ΔCn , missed cleavages, peptide length, charge
433 state, and precursor mass accuracy. For the phosphorylation dataset, site localization was
434 evaluated via AScore⁵³. For TMT-based reporter ion quantitation, we extracted the summed
435 signal-to-noise (S/N) ratio for each TMT channel and found the closest matching centroid to
436 the expected mass of the TMT reporter ion.

437 The search space for each reporter ion was limited to a range of 0.003 Th to prevent
438 overlap between the isobaric reporter ions. For protein-level comparisons, PSMs were
439 identified, quantified, collapsed to a 1% peptide false discovery rate (FDR), and further

440 collapsed to a final protein-level FDR of 1%. Protein assembly was guided by principles of
441 parsimony to produce the smallest set of proteins necessary to account for observed peptides.

442 Proteins were quantified by summing reporter ion counts across all matching PSMs
443 using in-house software, as previously described⁵². Poor quality PSMs, MS3 spectra with >8
444 TMT reporter ion channels missing, MS3 spectra with TMT reporter summed signal-to-noise
445 ratio <100, or no MS3 spectra were excluded from quantitation⁵¹. Protein quantitation values
446 were exported for further analysis in Microsoft Excel or SAS JMP. Each reporter ion channel
447 was summed across all quantified proteins and normalized assuming equal protein loading of
448 all 10 samples.

449 **Evolutionary relationship of *PfCRK4*.** Protein kinase amino acid sequences from the
450 cyclin-dependent kinase (CDK) families of apicomplexans and model organism and non-
451 CDK out-group kinases were chosen for comparison with *PfCRK4* (Supplementary Data
452 table 1). Kinase domain sequences were trimmed, aligned using MUSCLE⁵⁶ and analysed
453 using RAxML⁵⁷.

454 **Domain drawings of *PfCRK4* and its orthologues in *Plasmodium* spp.** Domain boundaries
455 were identified by searching sequences against Pfam⁵⁸ using the HMMER package⁵⁹.
456 Domains were illustrated using Protdraw (<http://sourceforge.net/projects/protdraw/>).

457 **Homology modelling of *PfCRK4*.** The protein kinase domain of *PfCRK4* was modelled
458 using the co-crystal structure of the human CDK2-Cyclin A-substrate peptide complex
459 (1QMZ.pdb)²⁵. We used this structure as a template in order to model *PfCRK4* in a
460 potentially active form. The kinase domains of the target and query sequences were aligned
461 using MUSCLE⁵⁶ followed by manual review and adjustment. Homology models were made
462 using MODELLER 9.15⁶⁰. Structural alignments were reviewed and adjusted using
463 Chimera⁶¹.

464 **Identification of conditionally related clusters in proteomic and phosphoproteomic data.**

465 The total sum signal-to-noise values of peptides collapsed into proteins are given in
466 Supplementary Data table 3. For phosphopeptides values are given in Supplementary Data
467 table 4. Contaminants and false positives were removed, and values for each channel were
468 normalized by the column sum. Values were placed on a scale of 0 - 1 by dividing by the
469 maximum value for the protein or phosphopeptide. Scaling was done for all channels at once
470 and separately for experiments at 29 hpi and 37 hpi. Phosphopeptides for which the total sum
471 signal-to-noise sum ≤ 100 were not included in clustering analysis, and the total sum signal-
472 to-noise values were adjusted to reflect protein levels from the proteome analysis. K-means
473 clustering⁶² was done using R (<http://cran.r-project.org/>) with the Hartigan-Wong method
474 using 1,000 starting configurations and 10,000 iterations. Clustering results, log₂-fold
475 changes, and $-\log_{10} p$ -values (Student's t-test) are included in Supplementary Data tables 3
476 and 4.

477 **Gene orthology relationships.** Relationships between *P. falciparum* proteins and
478 *Saccharomyces cerevisiae* orthologues for Figure 3F Box (Supplementary Data table 3) were
479 investigated using a downloaded version of the OrthoMCL database, version 5⁶³. A
480 comprehensive orthology analysis is provided in Supplementary Data table 6.

481 **Functional annotation and enrichment analysis.** The functional enrichment of sets of
482 proteins with phosphopeptides most reduced in knockdown experiments at 29 hpi and 37 hpi
483 was assessed using Ontologizer⁶⁴. Gene association files containing gene ontology (GO)
484 terms for *Plasmodium falciparum*, *Saccharomyces cerevisiae*, and *Homo sapiens* were from
485 <http://www.geneontology.org/gene-associations>⁶⁵ downloaded on May 21, 2015. To generate
486 comparative heat maps of GO terms for both time points we selected the most specific terms,
487 *i.e.* those without children, which we refer to as “tips” for convenience. Enrichment p -values
488 were included in the heat map if they were present at either time point; p -values for non-tip

489 terms were included if the term was a tip at the other time point. GO term annotations of the
490 proteins detected in this study are given in Supplementary Data table 3 and defined in
491 Supplementary Data table 7.

492 **Functional comparison of the *PfCRK4*-regulated set of phosphoproteins.** The functional
493 similarity between *PfCRK4* and other protein kinases was assessed by comparing GO terms
494 of the *PfCRK4*-regulated set with GO terms of reference kinase substrate sets. Reference
495 human kinase substrate sets were obtained from PhosphoSitePlus
496 (<http://www.phosphosite.org/staticDownloads.do>)²⁸ on May 4, 2015, the substrate set from
497 *Saccharomyces cerevisiae* (*Sc*) *cdc28* came from²⁷. Reference kinases with less than 30
498 identified substrates were omitted in the analysis. Functional enrichment was assessed as
499 described above. To calculate the relatedness of the substrate sets of kinases with *PfCRK4*,
500 the number of terms in the intersect of the set of significantly enriched functions from the
501 *PfCRK4*-regulated set and the substrate set from the reference kinases was divided by the
502 number of significant functional terms in the substrate set from the reference kinase.

503 **Identification and enrichment analysis of phosphorylation motifs.** Alignments of
504 phospho-peptides with reduced abundance in the absence of Shield-1 were manually
505 inspected. The two motifs discernable in this set may be retrieved from tab-delimited tables
506 of triskaidekapeptides by the regular expressions⁶⁶: $\backslash\{t[A-Z]\{6\}[ST][IVLFMWA].[RKH]$ (the
507 $S\phi xK$ motif; ϕ , hydrophobic amino acid; x, any amino acid) and $\backslash\{t[A-Z]\{6\}[ST]P$ (the S/TP
508 motif). The set of phosphopeptides detected in this study (Supplementary Data table 4) was
509 searched with these regular expressions and the percentage peptides comprised of these
510 motifs was calculated. Phosphopeptides containing $S\phi xK$ and S/TP motifs are identified in
511 Supplementary Data table 4. Sequence logos were made from alignments of phosphopeptides
512 containing $S\phi xK$ and S/TP motifs from sets of phosphopeptides reduced in the absence of
513 Shield-1 using Weblogo (<http://weblogo.berkeley.edu>)⁶⁷.

514 **Data availability**

515 The data supporting the results of this study are available within the paper and its
516 Supplementary Information and Supplementary Data. Sequences of oligonucleotides and
517 plasmids generated for this study are available from the corresponding author upon request.

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519

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679

680 **Figure legends**

681 **Fig. 1 | Conditional destabilization identified *P. falciparum* CRK4 as essential for**
682 **asexual blood-stage development. a**, Schematic of the *P. falciparum* blood-stage cycle
683 indicating peak transcript times of kinases targeted with a destabilization domain; purple,
684 approximate onset of DNA replication; blue, approximate duration of schizogony. **b**, Single
685 crossover 3' replacement strategy to fuse candidate kinases with a hemagglutinin (HA) and a
686 destabilization domain (DD) tag, which destabilizes kinases in absence of Shield-1. Note,
687 PKAc, ABCk2, and PF3D7_0420100 were refractory to HA-DD fusion. **c**, Effect of
688 conditional kinase destabilization on parasite growth over two cycles (\pm SD); PMR, parasite
689 multiplication rate. **d**, Western blots demonstrate conditional destabilization of *Pf*CRK4 in
690 two genetic backgrounds; the level of knockdown is indicated below; H3, histone H3.

691 **Fig. 2 | Nuclear localized *Pf*CRK4 is essential for trophozoite-to-schizont transition and**
692 **DNA replication. a**, Domain structure of *Pf*CRK4; blue, kinase domain; yellow, kinase
693 domain insertions relative to human CDK2; red, predicted nuclear localization signal (NLS);
694 asterisk, active site residues; numbers, first and last amino acid (AA). **b**, Immunofluorescence
695 detection of *Pf*CRK4-HA-DD in blood stage parasites (representative of three biological
696 replicates). **c**, Light microscopy of May-Grünwald-Giemsa-stained *Pf*CRK4-depleted
697 parasites (D10 parent, representative of two biological replicates and results with P2G12-
698 *Pf*CRK4-HA-DD). **d**, Ultrastructure of *Pf*CRK4-depleted parasites (D10 parent,
699 representative of $n \geq 25$ cells per condition); arrowhead, spindle pole body; n, nucleus; Rh,
700 rhoptry. **e**, Nuclear development in *Pf*CRK4-depleted parasites (D10 parent, representative of
701 three biological replicates and results with P2G12-*Pf*CRK4-HA-DD). **f**, Immunofluorescence
702 detection of spindle structures in *Pf*CRK4-depleted parasites (P2G12 parent, representative of
703 two biological replicates). **g**, Quantification of centriolar plaques by immunofluorescence in

704 *PfCRK4*-depleted parasites (D10 parent; *p*-value, Chi-Square test; representative of two
705 biological replicates and results with P2G12-*PfCRK4*-HA-DD). **h**, DNA content of *PfCRK4*-
706 depleted parasites from both parental lines; mean \pm SD of triplicates (representative of three
707 biological replicates); ring stage DNA content defined as 1 C. **i**, Proliferation of parasites
708 when *PfCRK4* was re-stabilized from 38 hpi or 48 hpi of the first cycle onwards, mean \pm SD
709 (representative of two biological replicates and results with P2G12-*PfCRK4*-HA-DD).

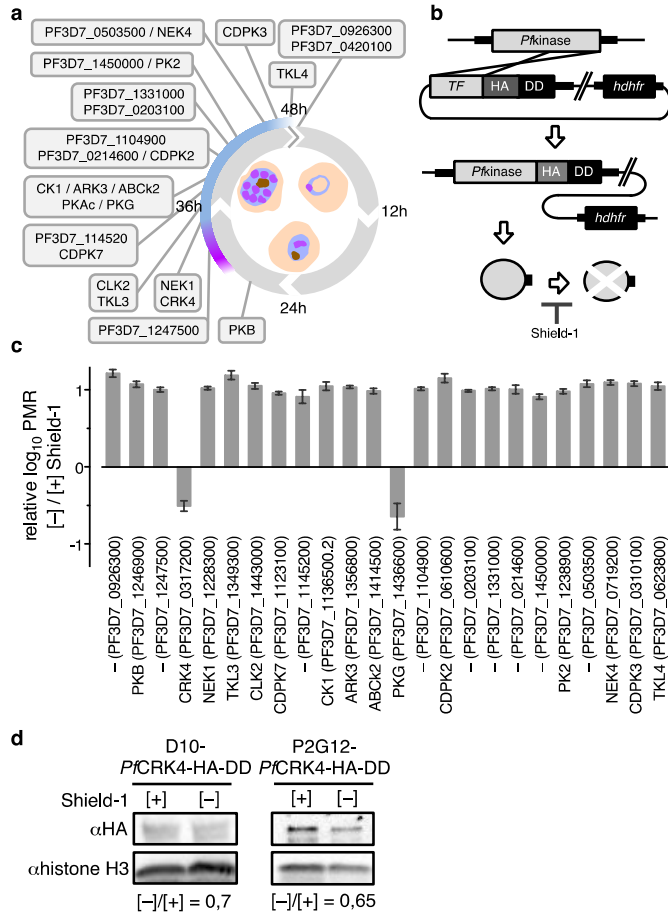
710 **Fig. 3 | *PfCRK4* regulates S phase.** **a**, Schematic illustrating the sampling time points for
711 phosphoproteomic profiling. **b**, Differential phosphorylation in D10-*PfCRK4*-HA-DD
712 parasites [-] Shield-1 relative to [+] Shield-1; 29 hpi in duplicate, 37 hpi in triplicate; *p*-
713 values by Student's *t*-test; grey-shaded regions indicate >2 -fold changes in phosphorylation
714 and *p*-values of <0.05 , number of peptides and respective proteins are indicated. **c**, GO term
715 enrichment analysis of proteins with ≥ 2 -fold reduced phosphorylation (*p*-values <0.05); *p*-
716 values, Modified Fisher's Exact Test. Box: *P. falciparum* homologues of *S. cerevisiae* factors
717 required for origin of replication activation *in vitro* with reduced phosphorylation at 37 hpi in
718 *PfCRK4*-depleted parasites; Pol, DNA polymerase; MCM, mini chromosome maintenance
719 complex; TOP2, topoisomerase 2; RPA, replication protein A; ORC, origin recognition
720 complex. **d**, Percentage of shared GO terms of proteins with ≥ 2 -fold reduced phosphorylation
721 (*p*-values <0.05) upon *PfCRK4*-depletion and of substrate sets of human and *S. cerevisiae*
722 (*Sc*) kinases.

723 **Fig. 4 | *PfCRK4* is essential throughout schizogony and critical for transmission.** **a**, DNA
724 content analysis of D10-*PfCRK4*-HA-DD parasites when *PfCRK4* was present or depleted
725 from 32 hpi, or 36 hpi onwards; 1 C, ring stage DNA content; asterisks, time of Shield-1
726 removal (representative of two biological replicates and results with P2G12-*PfCRK4*-HA-
727 DD). Note, second cycle ring-stage parasites re-appear at 48 hpi when *PfCRK4* is present. **b**,

728 DAPI-stained nuclei of parasites from Fig. 4a (representative of $n \geq 70$ cells per condition). **c**,
729 Proliferation of cultures from Fig. 4a, mean \pm SD of triplicates. **d**, Immunofluorescence
730 detects *PfCRK4*-HA-DD in gametocytes (P2G12 parent, representative of two biological
731 replicates). **e**, Depletion of *PfCRK4* during gametocytogenesis diminished oocyst numbers on
732 *An. gambiae* female midguts; bars, mean oocyst numbers; ns, not significant; **, $p \leq 0.01$
733 (Kruskal-Wallis test). **f**, Ultrastructure of mosquito-stage parasites from infections with
734 *PfCRK4*-depleted gametocytes (P2G12 parent, representative of \geq ten cells from independent
735 infection experiments); n, nucleus; ER, endoplasmic reticulum; m, mitochondria; c, capsule.
736

737 **Figures**

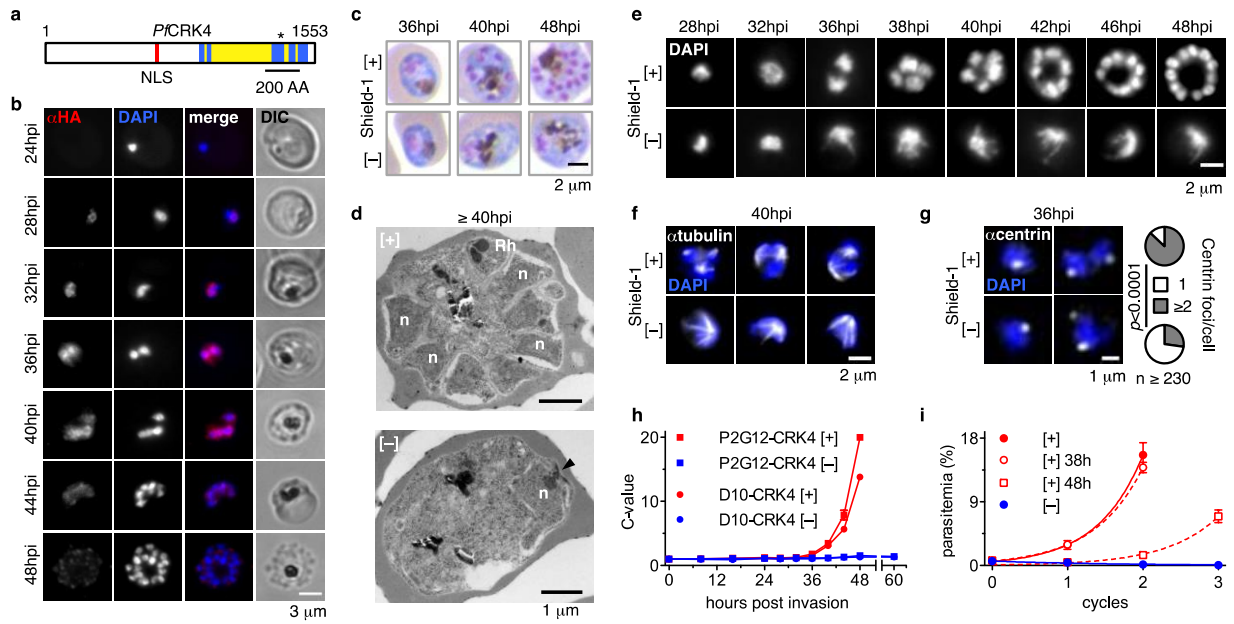
738 **Fig. 1**



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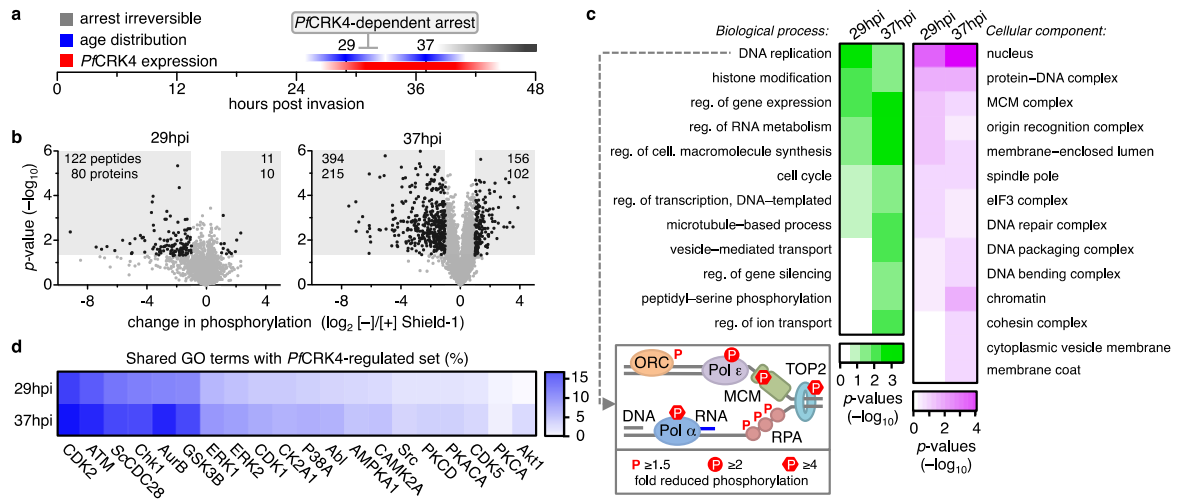
741 **Fig. 2**



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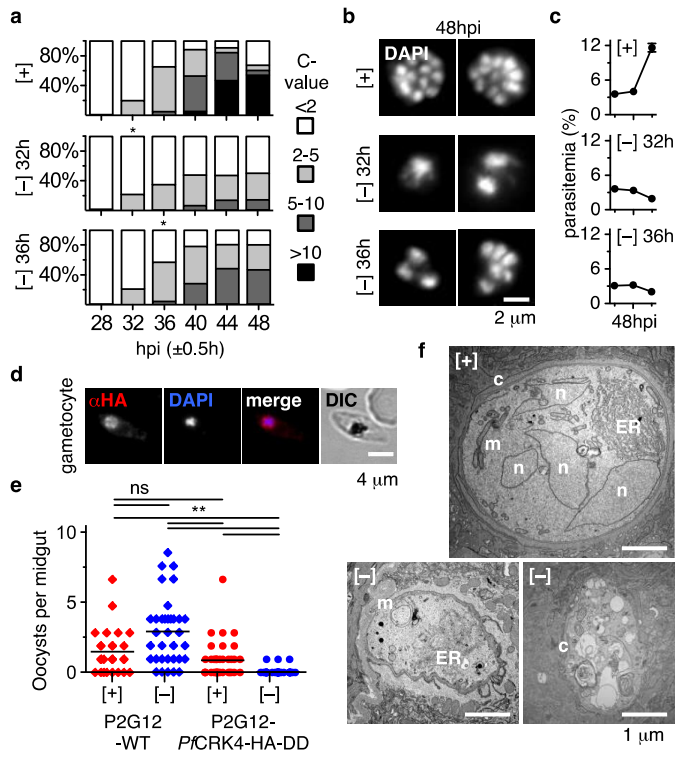
744 **Fig. 3**



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747 **Fig. 4**



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