1	EXP1 is required for organization of the intraerythrocytic malaria parasite vacuole
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16	Running title: Functional analysis of Plasmodium falciparum EXP1
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20	
21	
22	Abstract word count: 250
23	
24	
25	Importance word count: 149
26	

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27 Abstract

28 Intraerythrocytic malaria parasites reside within a parasitophorous vacuole membrane (PVM) 29 that closely overlays the parasite plasma membrane (PPM) and constitutes the barrier between 30 parasite and host compartments. The PVM is the site of several essential transport activities but 31 the basis for organization of this membrane system is unknown. We utilized the second-32 generation promiscuous biotin ligase BioID2 fused to EXP2 or HSP101 to probe the content of 33 the PVM, identifying known and novel candidate PVM proteins. Among the best represented 34 hits were members of a group of single-pass integral membrane proteins that constitute a major 35 component of the PVM proteome but whose function remains unclear. We investigated the 36 function of EXP1, the longest known member of this group, by adapting a CRISPR/Cpf1 37 genome editing system to install the TetR-DOZI-aptamers system for conditional translational 38 control. EXP1 knockdown was essential for intraerythrocytic development and accompanied by 39 profound changes in vacuole ultrastructure, including increased separation of the PVM and 40 PPM and formation of abnormal membrane structures in the enlarged vacuole lumen. While 41 previous in vitro studies indicated EXP1 possesses glutathione S-transferase activity, a mutant 42 version of EXP1 lacking a residue important for this activity in vitro still provides substantial 43 rescue of endogenous exp1 knockdown in vivo. Intriguingly, while activity of the Plasmodium 44 translocon of exported proteins was not impacted by depletion of EXP1, the distribution of the 45 translocon pore-forming protein EXP2 was substantially altered. Collectively, our results reveal 46 a novel PVM defect that indicates a critical role for EXP1 in maintaining proper PVM 47 organization.

48

49 Importance

Like other obligate intracellular apicomplexans, blood-stage malaria parasites reside within a
membrane-bound compartment inside the erythrocyte known as the parasitophorous vacuole.
Although the vacuole is the site of several transport activities essential to parasite survival, little

53 is known about its organization. To explore vacuole biology, we adopted recently developed 54 proteomic (BioID2) and genetic (CRISPR/Cpf1) tools for use in Plasmodium falciparum, which 55 allowed us to query the function of the prototypical vacuole membrane protein EXP1. 56 Knockdown of EXP1 showed that a previously reported glutathione S-transferase activity cannot 57 fully account for the essential function(s) of EXP1 and revealed a novel role for this protein in 58 maintaining normal vacuole morphology and PVM protein arrangement. Our results provide new 59 insight into vacuole organization and illustrate the power of BioID2 and Cpf1 (which utilizes a T-60 rich PAM uniquely suited to the *P. falciparum* genome) for proximity protein identification and 61 genome editing in *P. falciparum*.

62

63 Introduction

64 The obligate intracellular malaria parasite *Plasmodium falciparum* resides within a 65 parasitophorous vacuole (PV) established during host cell invasion which constitutes the 66 principal barrier between the parasite and its host cell (1). During the parasite blood stage, 67 several essential transport activities at this membrane enable host erythrocyte subversion and 68 parasite growth (2). This includes the uptake of large amounts of host cytosol through a process 69 involving a double-membrane invagination of the parasitophorous vacuole membrane (PVM) 70 and parasite plasma membrane (PPM) known as the cytostome (3). Catabolism of hemoglobin, 71 the principal component of this ingested material, provides free amino acids for parasite 72 metabolism and opens space within the host compartment for parasite expansion (4-6). The 73 vacuole is also a key trafficking site for the export of hundreds of parasite proteins which are 74 first secreted into the PV and then translocated across the PVM by the *Plasmodium* translocon 75 of exported proteins (PTEX) (7-9). The PTEX core complex is composed of the HSP101 AAA+ 76 ATPase chaperone, which unfolds exported cargo and passes it through an oligomeric pore in 77 the PVM formed by a second PTEX protein, EXP2 (10). A third component, PTEX150, serves to 78 couple HSP101 to the pore. While proteins must be unfolded to pass through the EXP2 pore

(11), small molecules up to 1,400 Da can pass through this channel, allowing for nutrient uptakeand waste exchange (12-14).

81 Aside from EXP2, the known *P. falciparum* PVM proteome is comprised mainly of a 82 group of single-pass integral membrane proteins oriented with C-terminus facing the host 83 cytosol and N-terminus in the PV lumen (15-17). This group consists of the prototypical PVM 84 protein EXP1 and the early transcribed membrane proteins (ETRAMPs, also known as small 85 exported proteins (SEPs) in *P. berghei*) and contains some of the most highly expressed genes 86 in the blood stage (17-19). The function of these proteins is largely unknown although EXP1 has 87 been shown to possess glutathione S-transferase (GST) in vitro, which has been proposed to 88 support detoxification of hematin released by hemoglobin catabolism (20). 89 One of the most striking features of the PV is the intimate proximity of the PVM and 90 PPM, which is maintained until a very late stage of parasite development when PVM rounding 91 occurs just prior to egress (2, 21). Additional lateral organization of this compartment is 92 suggested by the formation of distinct oligomeric arrays of EXP1 and ETRAMPs in the PVM (22) 93 and by the non-uniform distribution of PTEX components as well as PV-targeted exported and 94 non-exported fluorescent fusion reporter proteins, which have been shown to display a punctate 95 distribution in the PV described as a "necklace of beads" (7, 23-27). Visualized by 96 immunofluorescence, this punctate arrangement is most prominent in the early ring stage of 97 parasite development and resolves into a more homogenous distribution in the trophozoite and 98 schizont stage, particularly for EXP2 (26). While these observations point to a highly 99 coordinated membrane system, the basis for PV/PVM organization is unknown. 100 Here, we applied the second generation BioID2 proximity ligase system fused to multiple 101 components of PTEX to probe the protein content of the PV and PVM, revealing known vacuole 102 proteins as well as novel PV/PVM candidates. To interrogate the function of select hits, we 103 adapted a CRISPR/Cpf1 system that recognizes a T-rich protospacer adjustment motif, greatly 104 expanding the repertoire of guide RNA targets available for editing the *P. falciparum* genome.

105	Prompted by the high representation of EXP1 in our BioID2 datasets, we used Cpf1 editing to
106	generate an EXP1 conditional knockdown mutant to explore its function. Depletion of EXP1
107	produced a lethal defect that was largely rescued by a mutant version of EXP1 shown to be
108	defective in GST activity in vitro, suggesting that this activity does not fully account for EXP1
109	function in vivo. Instead, EXP1 knockdown resulted in dramatic changes to PV morphology and
110	PVM protein organization, revealing a novel role in maintaining proper order within the PV.
111	
112	Materials and Methods
113	
114	Parasite Culture
115	Deidentified, IRB-exempt red blood cells (RBCs) were obtained from the American National Red
116	Cross. <i>P. falciparum</i> NF54 ^{attB} and derivatives were maintained under 5% O_2 , 5% CO_2 , and 90%
117	N_2 at 2% hematocrit in RPMI 1640 supplemented with 27 mM sodium bicarbonate, 11 mM
118	glucose, 0.37 mM hypoxanthine, 10 μ g/ml gentamicin ad 0.5% Albumax I (Gibco).
119	
120	Plasmids and genetic modification of <i>P. falciparum</i>
121	Cloning was carried out with Infusion (Clontech) or NEBuilder HiFi (NEB) unless noted
122	otherwise. Primer and synthetic gene sequences are given in Table S1. To generate a BioID2
123	fusion to the endogenous EXP2 C-terminus, the coding sequence of the Aquifex aeolicus biotin
124	ligase with an R40G mutation (BioID2) bearing a 3' 3xHA epitope tag was amplified with primers
125	P1/P2 from plasmid MCS-BioID2-HA (28) (Addgene #74224) and inserted between AvrII and
126	Eagl in plasmid pyPM2GT-EXP2-mNeonGreen (29), replacing the AvrII site with an Nhel site
127	and resulting in the plasmid pyPM2GT-EXP2-BioID2-3xHA. This plasmid was linearized at the
128	AfIII site between the 3' and 5' homology flanks and co-transfected with pUF-Cas9-EXP2-CT-
129	gRNA (14) into NF54 ^{attB} . Selection was applied with 2 μ M DSM1 (30) 24 hours post-transfection.

A clonal line was isolated by limiting dilution after parasites returned from selection and
 designated NF54^{attB}::EXP2-BioID2-3xHA.

132 For fusion of BioID2 to endogenous HSP101, a flank assembly targeting the 3' end of 133 hsp101 was amplified from plasmid pPM2GT-HSP101-3xFlag (14) with primers P3/P4 and 134 inserted between XhoI and NheI in pyPM2GT-EXP2-BioID2-3xHA, resulting in plasmid 135 pvPM2GT-HSP101-BioID2-3xHA. This plasmid was linearized at AfIII, co-transfected with plasmid pAIO-HSP101-CT-gRNA (14) into NF54^{attB}, selected with 2 µM DSM1 and cloned upon 136 137 returning from selection, resulting in the line NF54^{attB}::HSP101-BioID2-3xHA. 138 For Cas9-mediated editing of the exp1 locus, the pAIO (31) plasmid was first simplified 139 by removing the vDHODH-2A fusion to Cas9-NLS-FLAG using a QuikChange Lightning Multi 140 Site Directed Mutagenesis kit (Agilent) and the primer P5, resulting in the plasmid pAIO2. The 141 BtgZI site in the pre-sgRNA cassette was then replaced with AfIII by inserting the annealed 142 sense and antisense oligo pair P6/P7 into BtgZI, resulting in the plasmid pAIO3. Three Cas9 143 targets were chosen at the 3' end of exp1 (GACGACAACAACCTCGTAAG, 144 AGGTTGTTGTCGTCACCTTG and AGTGTTCAGTGCCACTTACG) and each guide RNA 145 (gRNA) seed sequence was synthesized as a sense and anti-sense oligo pair (P8/P9, P10/P11, 146 and P12/P13, respectively). Each oligo pair was annealed and inserted into the AfIII site of 147 pAIO3 to yield the plasmids pAIO3-EXP1-CT-gRNA1, pAIO3-EXP1-CT-gRNA2 and pAIO3-148 EXP1-CT-gRNA3.

For Cpf1 editing, AsCpf1 and LbCpf1 were PCR amplified from plasmids pcDNA3.1hAsCfp1 and pcDNA3.1-hLbCfp1 (Addgene #69982 and #69988) (32) with primer pairs P14/15 and P16/15, respectively, and inserted into pAIO3 between BamHI and XhoI. The Cas9 pregRNA cassette was then replaced with a pre-gRNA cassette with the appropriate direct repeat region for AsCpf1 or LbCpf1 using a QuikChange Lightning Multi Site Directed Mutagenesis kit and the primers P17 or P18, resulting in plasmids pAIO-AsCpf1 and pAIO-LbCpf1, respectively. For Cpf1-mediated editing of the *exp1* locus, two Cpf1 targets were chosen at the 3' end of *exp1* 156 (CAGCTGTTTAGTGTTCAGTGCCAC and GTGTTCAGTGCCACTTACGAGGTT) and each 157 gRNA seed sequence was synthesized as a sense and anti-sense oligo pair for AsCpf1 158 (P19/P20 and P21/P22, respectively) and LbCpf1 (P23/P24 and P25/P26, respectively). Each 159 oligo pair was annealed and inserted into the AfIII site of pAIO-AsCpf1 or pAIO-LbCpf1 to yield 160 the plasmids pAIO-AsCpf1-EXP1-CT-gRNA1, pAIO-AsCpf1-EXP1-CT-gRNA2, pAIO-LbCpf1-161 EXP1-CT-gRNA1 and pAIO-LbCpf1-EXP1-CT-gRNA2. To generate selectable Cpf1 plasmids, 162 the AfIII site between the yDHODH and Cas9 expression cassettes in the plasmid pUF-Cas9-163 pre-sgRNA (14) was replaced with an Xmal site using QuikChange Lightning Multi Site Directed 164 Mutagenesis kit and the primer P27. The Cpf1 coding sequence with the PbDT 3' UTR and 165 adjacent pre-gRNA cassette was then amplified from pAIO-AsCpf1 or pAIO-LbCpf1 using 166 primers P28/P29 or P28/P30 and inserted between XhoI and NotI, resulting in the plasmids 167 pUF-AsCpf1-pre-gRNA and pUF-LbCpf1-pre-gRNA, respectively. Oligo pairs P21/P22 and 168 P25/P26 were annealed and inserted into the AfIII site in the pre-gRNA cassette of these 169 vectors to yield pUF-AsCpf1-EXP1-CT-gRNA2 and pUF-LbCpf1-EXP1-CT-gRNA2, respectively. 170 For fusion of 3xHA-GFP11 to EXP1, a 5' homology flank (up to, but not including, the 171 stop codon) was amplified from NF54^{attB} genomic DNA using the primers P31/P32. As the *exp1* 172 coding sequence did not allow for synonymous mutations in the protospacer adjustment motifs 173 of the three Cas9 gRNAs, several synonymous mutations were incorporated in the seed 174 sequences of these target sites. A 3' homology flank (beginning 11 bp downstream of the stop 175 codon) was amplified using the primers P33/P34 and the flank amplicons were assembled in a 176 second PCR reactions using the primers P32/P33 and inserted between XhoI and AvrII in 177 pyPM2GT-EXP2-3xHA-GFP11 (14), resulting in the plasmid pyPM2GT-EXP1-3xHA-GFP11. 178 This plasmid was linearized at the AfIII site between the 3' and 5' homology flanks and co-179 transfected into NF54^{attB} with the above Cas9 and Cpf1 plasmids designed to target *exp1*. Selection was applied with DSM1 and the expected integration was confirmed by diagnostic 180 181 PCR using primers P35/P36.

For generation of the ETRAMP10.2^{apt} and ETRAMP5^{apt} lines bearing a 3xHA fusion, a 182 183 Cpf1 gRNA target was chosen just upstream of the etramp10.2 and etramp5 stop codons (TGACTCTTGGTGTGGTACTTCTTC and GGTTCTTCGGTTTTGACTTCGTCT, respectively) 184 185 and the gRNA seed sequences were synthesized as the sense and anti-sense primer pairs 186 P37/P38 and P39/P40, which were annealed and inserted into the AfIII site of the plasmid pAIO-187 AsCpf1, resulting in the plasmids pAIO-AsCpf1-ETRAMP10.2-gRNA1 and pAIO-AsCpf1-188 ETRAMP5-gRNA1. A modified version of the plasmid pMG75 (33) was generated by first 189 replacing the AvrII site with an Ncol site using a QuikChange Lightning Multi Site Directed 190 Mutagenesis kit and the primer P41. The BirA*-3xHA coding sequence from plasmid pBirA*-191 3xHA-LIC-DHFR (34) was then amplified with primers P42/P43 and inserted between 192 BstEll/Aatll and the 2xattP sequence was subsequently removed by using QuikChange 193 Lightning Multi Site Directed Mutagenesis kit and the primer P44, resulting in the plasmid 194 pMG75∆attP-BirA*-3xHA. To target the 3' end of *etramp10.2* or *etramp5*, a 5' homology flank (up to but not including the stop codon) was amplified from NF54^{attB} genomic DNA using primers 195 196 pairs P45/P46 or P47/P48, incorporating synonymous mutations in the seed sequence and 197 protospacer adjustment motif of the gRNA target sites within each etramp coding sequence. A 198 3' homology flank (beginning 89 or 257 bp downstream of the stop codon, respectively) was 199 amplified using the primer pairs P49/P50 or P51/P52. The corresponding 5' and 3' flank 200 amplicons were assembled in a second PCR reaction using the primer pairs P46/P49 and 201 P48/P51 and inserted between AscI/AvrII in pMG75∆attP-BirA*-3xHA, removing the BirA* 202 sequence and resulting in the plasmids pMG75dattP-ETRAMP10.2-3xHA and pMG75dattP-203 ETRAMP5-3xHA. These plasmids were linearized at the AfIII site and co-transfected with the corresponding pAsCpf1-gRNA plasmid into NF54^{attB} parasites. Cultures were maintained with 204 205 1µM aTc from the time of transfection and selection with 2.5 µg/ml Blasticidin-S was applied 24 206 hours post-transfection. After returning from selection, parasites were cloned and proper

integration at the 3' end of *etramp10.2* or *etramp5* was confirmed by PCR with primer pairs
P53/P36 or P54/P36, respectively.

209 For generation of a combined TetR-DOZI-aptamers (TDA) and DiCre knockdown system 210 to target *exp1*, the tandem NLS-FKBP12-Cre19-59 and NLS-FRB-Cre60-343 cassettes from 211 plasmid pDiCre (35) were amplified with primers P55/P56 and inserted into the Ascl site of 212 plasmid pSN054 (36), a pJAZZ-based plasmid containing the TDA elements. This plasmid also 213 contains *loxP* sites immediately upstream of the 5' aptamer and immediately downstream of the 214 TetR-DOZI cassette. The latter *loxP* site is adjacent to the AscI site and was removing during 215 insertion of the DiCre cassette. To target *exp1*, 5' and 3' homology flanks immediately upstream 216 and downstream of the exp1 start and stop codons were amplified with the primer pairs 217 P57/P58 and P59/P60 and inserted sequentially at Fsel (5' flank) and between I-Ceul/I-Scel (3' 218 flank). A promoterless mruby3 coding sequence with the hsp86 3' UTR was amplified from 219 plasmid pLN-HSP101-SP-mRuby3 (21) using primers P61/P62, adding a loxP site immediately 220 before the mruby3 start codon, and inserted at I-Ceul. Finally, the exp1 coding sequence 221 (without introns) was recoded, synthesized as a gene block (IDT) and PCR amplified with 222 primers P63/P64, adding a 3' 3xHA tag, and inserted at AsiSI, resulting in the plasmid pEXP1^{apt}. This plasmid was co-transfected with pAIO-LbCpf1-EXP1-CT-gRNA1 into NF54^{attB}::HSP101-223 224 3xFLAG (14). Parasites were maintained with 1µM aTc from the time of transfection and 2.5 225 µg/ml Blasticidin-S was applied 24 hours later. A clonal line was isolated by limiting dilution after 226 parasites returned from selection and designated EXP1^{apt}. Integration at the 5' and 3' ends of 227 the *exp1* locus was evaluated by PCR with primer pairs P65/P36 and P66/P67, respectively. 228 Excision by DiCre was monitored with the primers P65/P68.

For complementation of EXP1^{apt} parasites, the re-coded *exp1* coding sequence (without introns) was PCR amplified from the synthesized gene block with primers P69/P70 and inserted between XhoI and NheI in pyEOE-attP-EXP2-3xMYC (14), resulting in the plasmid pyEOE-attP-EXP1-WT-3xMYC. The *exp1* codon 70 in this plasmid was then changed from AGA to ACA 233 using a QuikChange Lightning Multi Site Directed Mutagenesis kit and the primer P71, resulting 234 in the plasmid pyEOE-attP-EXP1-R70T-3xMYC. Finally, the mNeonGreen coding sequence 235 was amplified from pyPM2GT-EXP2-mNeonGreen (29) with primers P72/P73 and inserted at 236 Nhel in pyEOE-attP-EXP1-WT-3xMYC, resulting in the plasmid pyEOE-attP-EXP1-237 mNeonGreen-3xMYC. Complementing plasmids were co-transfected with pINT (37) into 238 EXP1^{apt} parasites to facilitate integration into the *attB* site on chromosome 6 and selection with 239 2 µM DSM1 was applied 24 hours post-transfection (in addition to 2.5 µg/ml Blasticidin-S and 240 1µM aTc for maintenance of endogenous exp1 control by the TetR-DOZI-aptamers system). 241 Parasites were cloned when they returned from selection and expression of EXP1 second 242 copies was confirmed by Western blot. 243 For targeting mRuby3 to the PV, the exp2 promoter and signal peptide were amplified 244 from NF54^{attB} genomic DNA with primers P74/P75 and inserted between AatII and Nhel in 245 plasmid pyEOE-attP-EXP2-3xMYC (14), replacing the hsp86 promoter and exp2 coding 246 sequence. The mRuby3 coding sequence was then PCR amplified from pLN-HSP101-SP-247 mRuby3 (21) using primers P76/P77 and inserted between Nhel and Eagl, replacing the 3xMYC 248 sequence and resulting in the plasmid pyEOE-attP-EXP2-5'UTR-SP-mRuby3. This plasmid was co-transfected with pINT (37) into EXP1^{apt} parasites, selection was applied 24 hours post-249 250 transfection with 2µM DSM1 and parasites were cloned upon returning from selection. To monitor EXP2 by live fluorescence in EXP1^{apt}, an endogenous mNeonGreen fusion to 251 252 EXP2 was generated by co-transfecting EXP1^{apt} with plasmids pyPM2GT-EXP2-mNeonGreen 253 (29) (linearized at AfIII) and pUF-Cas9-EXP2-CT-gRNA (14). Selection was applied 24 hours 254 post-transfection with 2µM DSM1 and parasites were cloned upon returning from selection. 255 256 Proximity labeling and mass spectrometry

257 Parasites were synchronized to an ~8 hour window by treatment with 5% w/v D-sorbitol and 200
258 μM exogenous biotin was added in the ring stage. After 18 hours, trophozoite and schizont-

infected RBCs were purified on an LD column mounted on a QuadroMACs magnetic separator
(Miltenyi Biotech), washed in PBS to remove residual biotin and lysed in RIPA buffer containing
protease inhibitors. Lysates were briefly sonicated and insoluble material and hemozoin was
cleared by centrifugation before loading onto streptavidin magnetic beads (Pierce). After rotating
at 4°C overnight, beads were washed 5X with RIPA followed by 5X washes in 50 mM Tris 6.8
containing 8M urea.

265 Protein samples were reduced and alkylated using 5mM Tris (2-carboxyethyl) phosphine and 10mM iodoacetamide, respectively, and then enzymatically digested by the sequential 266 267 addition of trypsin and lys-C proteases as described (38, 39). The digested peptides were 268 desalted using Pierce C18 tips (Thermo Fisher Scientific), dried and resuspended in 5% formic 269 acid. Approximately 1 µg of digested peptides were loaded onto a 25 cm long, 75 um inner 270 diameter fused silica capillary packed in-house with bulk C18 reversed phase resin (1.9 um, 100A 271 pores, Dr. Maisch GmbH). The 140-minute water-acetonitrile gradient was delivered using a 272 Dionex Ultimate 3000 UHPLC system (Thermo Fisher Scientific) at a flow rate of 200 nl/min 273 (Buffer A: water with 3% DMSO and 0.1% formic acid and Buffer B: acetonitrile with 3% DMSO 274 and 0.1% formic acid). Eluted peptides were subsequently ionized by the application of a distal 275 2.2kv and introduced into the Orbitrap Fusion Lumos mass spectrometer (Thermo Fisher 276 Scientific) and analyzed by tandem mass spectrometry (MS/MS). Data was acquired using a Data-Dependent Acquisition (DDA) method consisting of a full MS1 scan (Resolution = 120.000) 277 278 followed by sequential MS2 scans (Resolution = 15,000) to utilize the remainder of the 3 second 279 cycle time.

Data analysis was accomplished using the Integrated Proteomics pipeline 2 (Integrated Proteomics Applications, San Diego, CA). Data was searched against the protein database from *Plasmodium falciparum* 3D7 downloaded from UniprotKB (10,826 entries) on October 2013. MS/MS spectra searched using the ProLuCID algorithm followed by filtering of peptide-tospectrum matches (PSMs) by DTASelect using a decoy database-estimated false discovery rate
of <1%.

286

287 Antibodies

288 The following antibodies were used for immunofluorescence assays (IFA) and western blot

289 (WB) analysis at the indicated dilutions: mouse anti-HA monoclonal antibody HA.11 (Covance;

290 1:500 WB); rabbit polyclonal anti-HA SG77 (ThermoFisher; 1:500 IFA and WB); mouse anti-

Flag monoclonal antibody clone M2 (Sigma; 1:300 IFA); mouse anti-EXP1 monoclonal antibody

292 (20) (1:500 WB); mouse anti-EXP2 monoclonal antibody clone 7.7 (40) (1:500 IFA and WB);

rabbit polyclonal anti-SBP1 (41) (1:500 IFA); mouse anti-cMYC monoclonal antibody 9E10

294 (ThermoFisher; 1:166 IFA and WB); rabbit polyclonal anti-*Plasmodium* Aldolase ab207494

295 (Abcam; 1:500 WB).

296

297 Western blot

Western blots were carried out as previously described (14) and imaging with an Odyssey
infrared imaging system (Li-COR Biosciences). Biotinylated proteins were detected with IRDye
800-conjugated streptavidin used at 1:200. Signal quantification was performed with Image

302

301

303 Parasite growth assays

Studio software (Li-COR Biosciences).

304 Parasite cultures were washed five times to remove aTc, then plated at 5% parasitemia

305 (percentage of total RBCs infected) with or without 1µM aTc in triplicate. Every 24 hours, media

306 was changed and parasitemia was measured by flow cytometry on an Attune NxT

307 (ThermoFisher) by nucleic-acid staining with PBS containing 0.8 µg/ml acridine orange.

308 Subculture (1:1) was performed each day parasitemia exceeded 10% and half as often for -aTc

309 cultures within that group containing parasitemia less than 1%. Cumulative parasitemia was

back-calculated based on subculture schedule, data were log2 transformed and fitted to a linearequation to determine slope using Prism (Graphpad).

312

313 Transmission electron microscopy

EXP1^{apt} parasites were extensively washed to remove aTc, replated and allowed to develop 48 314 hours with or without aTc along with the parental NF54^{attB}:HSP101-3xFLAG line. Trophozoite-315 316 and schizont-infected RBCs were purified on an LD column mounted on a QuadroMACs 317 magnetic separator and fixed in 100 mM sodium cacodylate buffer, pH 7.2 containing 2% 318 paraformaldehyde and 2.5% glutaraldehyde (Polysciences) for 1 hour at room temperature for 319 ultrastructural analyses. Samples were washed in sodium cacodylate buffer at room 320 temperature and post-fixed in 1% osmium tetroxide (Polysciences) for 1 hour. Samples were 321 then extensively rinsed in dH20 before en bloc staining with 1% aqueous uranyl acetate (Ted 322 Pella) for 1 h. Following several rinses in water, samples were dehydrated in a graded series of 323 ethanol and embedded in Eponate 12 resin (Ted Pella). Sections (95 nm thick) were cut with a Leica Ultracut UCT ultramicrotome (Leica Microsystems), stained with uranyl acetate and lead 324 325 citrate, and viewed on a JEOL 1200 EX transmission electron microscope (JEOL USA) 326 equipped with an AMT 8-megapixel digital camera and AMT Image Capture Engine V602 327 software (Advanced Microscopy Techniques). For quantification of abnormalities, 100 328 trophozoites and 100 segmented schizonts were scored in each of two independent replicates. 329

330 Immunofluorescence imaging

331 For IFAs, cells were fixed with a mixture of cold 90% acetone and 10% methanol for 2 minutes,

except for export assays where fixation was carried out with room temperature 100% acetone

for 2 minutes, and processed as described (9). For detection of biotinylated proteins,

334 streptavidin-conjugated Alexa Fluor 594 (ThermoFisher) was included with secondary

antibodies at 1:200. Images were collected on an Axio Observer 7 equipped with an Axiocam

336 702 mono camera and Zen 2.6 Pro software (Zeiss) using the same exposure times for all337 images across sample groups and experimental replicates.

338

339 **Quantification of protein export**

340 IFA analysis of protein export was performed as described (14) except that image quantification 341 was carried out using the Image Analysis module in Zen 2.6 Pro (Zeiss). The border of each 342 single-infected RBC was traced using the DIC channel as a reference and the PVM was marked 343 using the "Segment by Global Thresholding" tool for the HSP101-3xFLAG-488 channel (low and 344 high thresholds set at 600 and 16,383 respectively and the Fill Holes option enabled). The 345 signal corresponding to exported SBP1 was determined by removing any SBP1 signal within the 346 PVM from the total SBP1 signal in each cell. Individual Maurer's clefts were identified using the 347 "Dynamic Thresholding" tool for the SBP1-594 channel (smoothing set to 7, threshold set to -348 500, minimum area set to 10 and the Watersheds option enabled with count set to 1) and 349 removing puncta within the PVM boundary from the total SBP1 puncta within each cell. 350

351 Live-fluorescent imaging of EXP2-mNeonGreen distribution

352 Parasite cultures were washed 5 times to remove aTc, then plated with or without 1µM aTc and 353 cultured for 48 hours. Trophozoite and schizont-infected RBCs were magnet purified and 354 stained with 2.5 µM BODIPY TR Ceramide (Thermo Fisher) for 15 minutes at 37°C, washed 355 once with media and immediately imaged. For each replicate, 40 single-infected RBCs were 356 selected moving top to bottom and left to right in each field to avoid bias. For each infected 357 RBC, the circumference of the PVM was traced with the profile tool in Zen 2.6 Pro using the 358 BODIPY TR Ceramide signal at the parasite periphery as a guide with the green channel turned 359 off to blind the experimenter to the mNeonGreen (mNG) signal. The mNG fluorescent intensity 360 along each PVM trace was then collected, mNG signal and distance along the trace were 361 normalized and data were analyzed with custom R scripts (ver. 3.5.2). In one approach, peaks

362	were designated as mNG fluorescent intensity that exceeded the mean of the minimum and
363	maximum intensity for at least 3 measured points before falling below this threshold. In an
364	alternative approach, minimum PVM circumference containing indicated amounts of EXP2-mNG
365	fluorescent signal was determined as the shortest distance along the PVM trace that returned
366	the designated portion of total fluorescent signal throughout the trace. Means from three
367	independent experiments were fitted to a smooth line with JMP Pro (ver. 14.2.0).
368	
369	Live-fluorescent imaging of PV-mRuby3
370	Parasite cultures were washed 5 times to remove aTc, then plated with or without $1\mu M$ aTc and
371	cultured for 48 hours. Trophozoite and schizont-infected RBCs were magnet purified and
372	stained with 1μ g/ml Hoechst 33342 trihydrochloride trihydrate (ThermoFisher) for 15 min at
373	37°C, washed once with media and immediately imaged.
374	
375	Data Availability
376	Custom R scripts used in this study are available at https://github.com/tnessel/Beck-Lab
377	
378	Results
379	
380	Identification of proteins at the luminal face of the PVM with BioID2
381	To probe the protein content of the PV/PVM, we attempted to fuse the BioID proximity
382	labeling system to the endogenous EXP2 protein. However, repeated attempts to generate an
383	EXP2-BioID fusion with a verified CRISPR/Cas9 strategy for editing the 3' end of exp2 were
384	unsuccessful. Endogenous EXP2 can tolerate a monomeric NeonGreen (mNG, 27 kDa) fusion
385	without an obvious fitness cost (29); thus the bulkier size (35 kDa) or enzymatic activity of BioID
386	may interfere with EXP2 trafficking or essential functions. To test the former possibility, we next

387 attempted fusion with the second generation BioID2 derived from Aquifex aeolicus (Figure 1A). 388 BioID2 lacks the DNA-binding domain present in the E. coli-derived BioID, resulting in a smaller 389 size (27 kDa) that has been shown to reduce trafficking defects relative to BioID fusions (28). 390 Parasites with an endogenous EXP2-BioID2 fusion were easily obtained and displayed robust 391 biotinylation activity at the PVM (Figure S1A and Figure 1B), suggesting the additional ~9 kDa 392 BioID DNA-binding domain is not compatible with EXP2 function. As an additional probe, we 393 also generated parasites with an endogenous HSP101-BioID2 fusion (Figure 1A,B). As EXP2 394 appears to be expressed at a stoichiometrically higher level than HSP101 and a fraction of 395 PTEX-independent EXP2 exists (14, 24, 26), we reasoned that these two fusions would similarly 396 position BioID2 within PTEX proximal to the luminal face of the PVM but might allow distinct 397 proximity labeling by the PTEX-independent fraction of EXP2 that could provide clues to its dual 398 function or organization in the PVM.

399 Western blot of EXP2-BioID2 and HSP101-BioID2 lysates showed extensive protein 400 biotinylation relative to parental controls and supplementation of cultures with 200 µM 401 exogenous biotin for 18 hours produced an approximately 2-fold increase in streptavidin signal 402 over parallel cultures maintained in standard RPMI containing 820 nM biotin (Figure 1C). 403 Overall biotinylation levels were higher in EXP2-BioID2 lines which may reflect the higher level 404 of EXP2 expression relative to HSP101 (14). Although streptavidin-labeled banding patterns 405 were distinct between the EXP2-BioID2 and HSP101-BioID2 lysates, both lines showed several 406 strongly labeled bands migrating at molecular weights consistent with other PTEX components 407 (Figure 1C, arrowheads). As expected, the most strongly labeled band in each lysate 408 corresponded with the BioID2-3xHA fusion. Notably, a band migrating at ~23 kDa displayed 409 prominent labeling in both lines (Figure 1C, arrow).

To identify labeled proteins, synchronized parasite cultures were supplemented with 200
 µM biotin in the ring stage and allowed to develop for 18 hours. Parasite-infected RBCs were
 magnetically purified before lysis, streptavidin chromatography and subsequent analysis by

413 mass spectrometry to identify biotinylated proteins. Mass spectrometry datasets from two independent experiments performed with EXP2-BioID2 or HSP101-BioID2 identified ~10-fold 414 415 more proteins than untagged parental controls and were substantially overlapping and highly 416 enriched for known PV/PVM proteins (Figure 1D and Table S2). The highest-ranking proteins by 417 normalized spectral abundance factor (NSAF) identified in both EXP2-BioID2 and HSP101-418 BioID2 included protein export machinery (PTEX and EPIC (42) complexes) and other proteins 419 known to reside in the PV lumen, exported proteins and the PVM membrane proteins EXP1 and 420 ETRAMP10.2 (Table 1). Beyond the top ~10% of each dataset, many additional known PV/PVM 421 and exported proteins were identified (Table S2). 422 EXP1 and ETRAMP10.2, both single-pass PVM membrane proteins with similar 423 expression timing to EXP2 (Figure S1B), ranked particularly high across all BioID2 experiments. 424 When Western blots of lysates from EXP2-BioID2 and HSP101-BioID2 were probed with an

425 anti-EXP1 monoclonal antibody, the signal coincided with the prominent ~23 kDa band labeled

426 by streptavidin (EXP1 is known to migrate at 23 kDa, higher than its predicated molecular

427 weight of 14.7 kDa following signal peptide cleavage (43)), consistent with the high level of

428 EXP1 representation in the BioID2 proteomics (Figure 1E). ETRAMP5, another member of this

429 group, was also well represented (though not consistently among the top hits). The functions of

430 ETRAMP10.2, ETRAMP5 and EXP1 are not known. Given their high level of representation

431 implying particular abundance and/or an intimate proximity to EXP2/PTEX, we focused on

432 functional characterization of these three proteins.

433

434 Efficient genome editing in *P. falciparum* with Cpf1/Cas12a

435 CRISPR/Cas9 genome editing technology has been rapidly adapted for manipulation of
436 *P. falciparum*. A limitation of Cas9-mediated editing is the requirement for a "GG" in the
437 protospacer adjustment motif (PAM), as such sites are comparatively rare given the high A+T
438 content of the *P. falciparum* genome (80.6%) (44). The type II, class V CRISPR endonuclease

Cpf1 (CRISPR from *Prevotella* and *Francisella*, also known as Cas12a) has recently emerged
as a favorable alternative to Cas9 with an inherently lower level of off-target cleavage (45, 46).
In contrast to Cas9, Cpf1 does not require a trans activating CRISPR (tracr) RNA, introduces a
staggered double-strand break in target DNA that is distal from the PAM and utilizes a T-rich
PAM, making it uniquely suited to editing the *P. falciparum* genome (32).

444 Cpf1 from Acidaminococcus sp. BV3L6 and Lachnospiraceae bacterium ND2006 445 (AsCpf1 and LbCpf1, respectively) have been shown to mediate efficient genome editing in 446 mammalian cells (32) and we tested both of these enzymes for their genome editing capacity in 447 P. falciparum relative to a Cas9 editing system we previously developed (14, 31). The 3' region 448 of exp1 was found to contain attractive Cas9 and Cpf1 target sites located in close proximity. 449 providing an opportunity for initial Cpf1 testing (Figure 2A). We generated plasmids for 450 expression of AsCpf1 or LbCpf1 and corresponding gRNAs to target two sites with Cpf1 near 451 the exp1 stop codon. In parallel, we generated constructs for targeting partially overlapping or 452 immediately adjacent sites with Cas9 (Figure 2A). These markerless Cas9 and Cpf1 editing 453 plasmids were co-transfected with a donor plasmid bearing a yDHODH cassette and flanks 454 designed to repair the intended double-strand breaks by double homologous recombination to introduce a 3xHA-GFP11 tag at the 3' end of exp1 (Figure 2A). In each case, parasites returned 455 456 from DSM1 selection in about 21 days with the expected edit, as gauged by diagnostic PCR and 457 Western blot (Figure 2B). To provide a tool for marker-free editing, we additionally generated 458 AsCpf1 and LbCpf1 vectors with a yDHODH cassette to allow for selection in parasites and 459 showed that these vectors also facilitated editing of the *exp1* locus (Figure S2A).

Encouraged by these results, we next applied the AsCpf1 system to edit the *etramp10.2* and *etramp5* loci to insert a 3xHA-epitope and simultaneously install the TetR-DOZI-aptamers (TDA) system for conditional translational repression (Figure S2B-D) (31, 33). Editing of the 3' end of both loci was also successful (Figure S2C,D), however both ETRAMP10.2^{apt} and ETRAMP5^{apt} parasite lines were found to a have truncated 3' aptamer arrays (reduced from 10X to 6X, data not shown). Partial truncation of 3' aptamers rapidly diminishes translational control
(31) and anhydrotetracycline (aTc) washout did not result in measurable knockdown (Figure
S2C,D), thus ETRAMP10.2 and ETRAMP5 were not pursued further in this study. Collectively,
these results indicate Cpf1 editing of the *P. falciparum* genome is equally efficient to Cas9 with
Cpf1 offering many additional gRNA targets owing to its T-rich PAM requirement.

470

471 Lethal EXP1 knockdown with a dual aptamer strategy

472 To guery EXP1 function in PV biology, we employed a dual aptamer TDA strategy using 473 a linear plasmid system to replace the endogenous exp1 coding sequence in an NF54^{attB} 474 parasite line bearing a 3xFLAG tag on the endogenous *hsp101* gene (10, 14). This was 475 accomplished by LbCpf1 editing to introduce an aptamer just upstream of the start codon and a 476 10X aptamer array just downstream of the stop codon (Figure 3A and Figure S3A,B). 477 Installation of 5' aptamers has been shown to reduced baseline expression even in the presence of aTc (14) and the resulting EXP1^{apt} parasites showed 87.5±7.2% reduction in EXP1 478 479 expression and a growth defect relative to the parent line (Figure 3B,C). Removal of aTc further reduced EXP1 levels (99.3±0.4% knockdown relative to parent by probing with anti-EXP1 and 480 481 65.9±3.1% relative to EXP1^{apt} +aTc by probing with anti-HA) and resulted in a complete block in 482 parasite growth, indicating EXP1 is required for intraerythrocytic development (Figure 3B.C). 483 As a complement to the titratable translational control afforded by the TDA system and 484 to simultaneously provide a parallel option for conditional knockout, we also engineered the 485 donor plasmid to place loxP sites around the recoded exp1 gene and inserted cassettes for 486 expression of the rapamycin-inducible dimerizable Cre recombinase (DiCre) downstream of the 487 modified *exp1* locus (Figure S3A). Treatment of EXP1^{apt} cultures with rapamycin induced the 488 expected excision event as gauged by diagnostic PCR but produced only modest impact on 489 parasite growth (Figure S2C-E). Consistent with this, the non-excised locus remained readily

490 detectable in these cultures even when parasites were grown for several days with rapamycin,

491 indicating inefficient excision of *exp1* that was unsuitable for functional analysis (Figure S2E).

492 The reason for the poor excision efficiency by DiCre in EXP1^{apt} was not explored further and the

493 robust translational knockdown achieved by TDA was used for the remainder of the study.

494

495 **EXP1 functional constraints**

496 While a 3xHA-GFP11 tag (6.2 kDa) could be efficiently fused to the endogenous EXP1 497 C-terminus (Figure 2), multiple attempts using the same editing strategy failed to generate an 498 endogenous fusion to mNG or BioID2, suggesting the essential function of EXP1 is perturbed by 499 bulky C-terminal fusions (data not shown). To directly test this possibility, we complemented the EXP1^{apt} parasites with a second copy of EXP1 bearing either a 3xMYC (4.5 kDa) or mNG-500 501 3xMYC (31.5 kDa) fusion (Figure 4A). While both versions of EXP1 were similar expressed, 502 only the 3xMYC fusion was able to rescue parasite growth upon aTc removal (percent growth 503 rate in -aTc relative to +aTc control was 7.28±20% in mNG-3xMYC compared with 67.63±7.36% in WT-3xMYC and 7.76±2.14% in uncomplemented EXP1^{apt}), indicating that introduction of a 504 505 bulky C-terminal fusion does indeed ablate EXP1 function (Figure 4B,C). Similar to EXP1-506 3xMYC, EXP1-mNG-3xMYC still localized at the parasite periphery with endogenous EXP1-507 3xHA but tended to show a more dispersed distribution possibly indicating perturbations in 508 trafficking (Figure 4D).

509 EXP1 has previously been reported to possess GST activity based on an *in silco* 510 functional prediction approach and *in vitro* biochemical analysis (20). This study identified the 511 arginine at position 70 in EXP1 as critical for GST activity *in vitro* and our functional 512 complementation system provided the opportunity to test the importance of this residue *in vivo* 513 by complementing EXP1^{apt} parasites with an EXP1-R70T-3xMYC mutant (Figure 4A,B). Similar 514 to the wild-type second copy, EXP1-R70T-3xMYC colocalized with endogenous EXP1-3xHA at the PVM (Figure 4D). To our surprise, the R70T mutant also provided substantial rescue upon
knockdown of endogenous EXP1 (percent growth rate in -aTc relative to +aTc control was
46.21±6.05% in R70T-3xMYC compared with 67.63±7.36% in WT-3xMYC), suggesting that
GST activity cannot fully explain the *in vivo* function of EXP1 (Figure 4C).

519

520 Depletion of EXP1 results in late cycle arrest and PV/PVM morphological abnormalities While EXP1 expression peaks about midway through the intraerythrocytic development 521 522 cycle (Figure S1B), EXP1 has been localized to merozoite dense granules and is thus expected to be delivered to the PVM immediately following RBC invasion and accumulate to peak levels 523 524 later in the cycle (47). To determine the impact of EXP1 knockdown on parasite development 525 from the point of invasion on, trophozoites (~32-42 hours post invasion) were magnet-purified 526 from synchronized cultures, washed of aTc to prevent new synthesis of EXP1 during dense 527 granule formation at the terminal stages of schizogony and used to initiate cultures with fresh 528 uninfected RBCs. New ring formation was not impacted relative to controls maintained with aTc. 529 indicating EXP1 is not important at this early stage; rather cultures developed normally until a 530 late stage when parasites arrested predominantly as trophozoites and schizonts and failed to 531 complete the cycle (Figure 5A). We compared these developmental defects with those following inactivation of protein export and PVM channel activity in EXP2^{apt}, an EXP2 conditional mutant 532 533 that we previously generated using the same dual aptamer TDA approach (14). EXP2^{apt} 534 parasites grown without aTc arrested at an earlier trophozoite stage that was distinct from 535 EXP1-depleted parasites, suggesting EXP2-dependent transport activities are not impacted by 536 loss of EXP1 (Figure 5B).

537 To directly test this, we monitored protein export beyond the PVM in EXP1^{apt} and 538 EXP2^{apt} parasites using the above experimental design of removing aTc from synchronized, 539 purified trophozoites and then analyzing export by IFA at the midpoint of the following cycle. 540 While EXP2 knockdown results in a robust block in export of SBP1 as previously reported (14), no defect in SBP1 export was observed following knockdown of EXP1 (Figure 5C,D). Rather,
SBP1 export was more robust in the absence of aTc as measured by mean fluorescent intensity
in the infected RBC compartment or the number of SBP1-positive Mauer's clefts (punctate
structures beyond the PVM) per infected RBC (Figure 5C). While the basis for the apparent
increase in exported SBP1 signal is unclear, these results clearly indicate EXP1 is not required
for protein export.

547 To better understand the impact of EXP1 depletion in late stage parasites, we next 548 examined parasite ultrastructure by transmission electron microscopy (TEM). As seen in the 549 parental line and EXP1^{apt} grown with aTc, the PVM normally tightly overlays the PPM (Figure 6A,B). In contrast, EXP1^{apt} parasites grown 48 hours without aTc displayed striking changes in 550 551 PV morphology with increased separation between the PVM and PPM in both trophozoites and 552 segmented schizonts (Figure 6C-E). Curiously, the lumen of these enlarged PVs often 553 contained additional membrane-enclosed structures filled with host cytosol (Figure 6D, double 554 arrowheads). Additionally, membrane structures in the host cell cytosol external to the PVM 555 (Figure 6B, enlarged region) which likely represent portions of the tubulovesicular network 556 (TVN) were observed with increasing frequency upon EXP1 depletion. Quantification of 557 morphological abnormalities observed by TEM in two independent experiments is shown in 558 Table 2.

In EXP1^{apt} parasites that displayed increased separation between PVM and PPM, the enlarged PV lumen often showed equivalent density with the host cytosol, possibly indicating a broken vacuole (Figure 6C,E). To ascertain if PVM integrity was compromised following EXP1 knockdown, we fused a signal peptide to the fluorescent protein mRuby3 to target it to the PV and expressed it under the control of the *exp2* promoter in the EXP1^{apt} line. Parasites depleted of EXP1 continued to show concentrated mRuby3 signal at the cell periphery, indicating the PVM is not compromised, despite altered PVM ultrastructure (Figure 6F).

566

567 EXP1 is required for proper organization of EXP2 in the PVM

568 To further investigate alterations in the PV, we evaluated the impact of EXP1 knockdown 569 on other PVM proteins. As expected from the observation that protein export remains 570 operational in EXP1^{apt}, depletion of EXP1 did not substantially alter EXP2 levels (Figure 7A). To monitor EXP2 distribution in EXP1^{apt} parasites, we introduced a C-terminal mNG fusion on the 571 572 endogenous copy of EXP2 and imaged parasites at a late stage corresponding with 573 developmental arrest. Live fluorescent analysis of a parental EXP2-mNG control line with an 574 unmodified *exp1* locus (29) showed a punctate distribution at early stages that often resolved 575 into several larger patches in trophozoites and schizonts (Figure 7B). In contrast, EXP2 distribution was substantially altered in EXP1^{apt}::EXP2-mNG parasites, often concentrating into 576 577 one or two discrete points along the PVM (Figure 7B). To guantify this altered localization of 578 EXP2-mNG, BODIPY-TR-Ceramide labeling was used as a guide to trace the PVM and collect 579 EXP2-mNG signal along the PVM circumference (Figure 7B). Analysis of these PVM traces 580 showed that the number of discrete EXP2 signal patches around the PVM was significantly reduced in EXP1^{apt} parasites with EXP2 signal concentrated into a smaller proportion of the 581 582 PVM circumference (Figure 7C,D). These results show that although EXP2 function is 583 preserved following depletion of EXP1, its organization in the PVM is drastically altered.

584

585 Discussion

Promiscuous biotin ligases have emerged as powerful tools for proximity-based protein identification in live cells (48). To date, the original *E. coli* BirA* (BioID) has been employed by four studies in *Plasmodium spp.*, including two that targeted BioID to the PV (49-52). Khosh-Naucke and colleagues targeted a signal peptide-GFP-BirA* fusion to the *P. falciparum* PV lumen (50) while Schnider and colleagues utilized a second copy of EXP1 with a C-terminal BirA* fusion to probe the PV in *P. berghei* (51). In the present study, we present the first use of BioID2 in *Plasmodium spp.*, which we fused to the endogenous copy of EXP2 or HSP101 to

identify proteins at the luminal face of the PVM. The ability to fuse mNG or BioID2 but not BioID 593 594 to EXP2 indicates EXP2 will not tolerate larger bulky fusions, possibly due to defects in 595 trafficking, oligomerization or assembly with other PTEX components, demonstrating that the 596 smaller size of BioID2 offers advantages over the original BioID system. We have also adapted 597 a Cpf1 editing system for use in P. falciparum which facilitated successful genome editing with 598 the same efficiency as our Cas9 editing system. Since the discovery of its T-rich PAM 599 requirements, Cpf1 has been suggested to possess unique promise for manipulation of the A+T 600 rich genomes of *Plasmodium spp.* (32). To our knowledge, this is the first report of Cpf1 use in 601 malaria parasites, realizing that potential of greatly expanding the repertoire of gRNA targets 602 available to manipulate the *P. falciparum* genome.

603 While we expect the EXP2-BioID2 and HSP101-BioID2 datasets to contain novel PV 604 and PVM proteins (Figure S1C) and are currently working to validate these candidates, we have 605 focused in the present study on functional analysis of EXP1, one of the highest-ranking proteins 606 in our proteomics. The study of EXP1 dates back more than 35 years when a monoclonal 607 antibody shown to label the PVM (40, 53, 54) was subsequently found to detect an antigen in 608 the EXP1 C-terminus (43). However, despite being one of the earliest discovered PVM proteins, 609 EXP1 function has remained obscure. Previous failed attempts to disrupt the exp1 gene in P. 610 falciparum (55) and P. berghei (56) indicated a critical role for EXP1 during the blood stage. 611 EXP1 is also expressed and localized to the PVM in the parasite liver stage (57) and a C-612 terminal region of *P. berghei* EXP1 has been shown to interact with rodent and human 613 Apolipoprotein H (56). While this interaction is important for *P. berghei* development within 614 hepatocytes, it could not be recapitulated in yeast two-hybrid assays with *P. falciparum* EXP1, 615 which displays low sequence identity with P. berghei EXP1 in the C-terminal region critical for 616 interaction with Apolipoprotein H, possibly indicating lineage-specific EXP1 adaptions. 617 In the present study, we found that blood stage function of *P. falciparum* EXP1 is 618 compromised by a C-terminal fusion to mNG. The topology of EXP1 orients the C-terminus on

619 the host cytosolic face of the PVM while the N-terminus is positioned within the PV lumen (15, 620 16). As EXP1 bearing a bulkier mDHFR-GFP C-terminal fusion is still inserted into the PVM with 621 proper topology (58), the EXP1-mNG defect seems unlikely to result from a perturbation in 622 trafficking. The mNG fusion may interfere with interactions that enable EXP1 to organize into 623 oligomeric arrays (22) as the C-terminus of P. berghei SEP/ETRAMP family members have 624 been shown to be important for oligomer formation (19). However, the entire C-terminal domain 625 of *P. berghei* EXP1 has recently been shown to be dispensable during the blood stage 626 (although it is important in both the mosquito vector and in the hepatocyte where truncated 627 EXP1 fails to properly traffic to the PVM, illustrating distinct EXP1 functional roles and trafficking 628 constraints between parasite life stages) (59). The inability to similarly truncate the EXP1 N-629 terminus in *P. berghei* implies its essential blood stage function occurs within the PV lumen (59). 630 An *in silico* approach to discovery of gene function through analysis of gene 631 relationships over large evolutionary distance suggested EXP1 may be a membrane GST (20). 632 Subsequent in vitro experiments found recombinant EXP1 to possess GST activity, particularly 633 toward hematin, leading to a model where EXP1 provides protection from the oxidative stress 634 that results from catabolism of hemoglobin within endocytic vesicles and the digestive vacuole 635 (20). While enzymes involved in hemoglobin degradation such as plasmepsin aspartic 636 proteases do traffic through the PV to reach the digestive vacuole, their PV residence is 637 transient with principal localization observed in the digestive vacuole (60). In contrast, the vast 638 majority of EXP1 is localized at the PVM, suggesting this is the principal site of function. Here, 639 we found that a version of EXP1 bearing an R70T mutation, which reduced recombinant EXP1 640 GST activity more than fivefold in vitro, can rescue parasite growth ~70% as well as the wild 641 type protein upon knockdown of endogenous EXP1 (Figure 4C). These results strongly suggest 642 that the proposed GST activity does not fully account for the essential blood stage function of P. 643 falciparum EXP1.

Known essential functions that manifest defects at the PVM include inactivation of 644 645 protein export and small molecule transport following knockdown of PTEX components (8, 9, 646 14, 61) and RON3 (62) or a block in egress following knockdown of key players in the protease 647 cascade that mediates PVM destruction at the end of the cycle (36, 63, 64). Parasites depleted 648 of EXP1 do not display an export defect and develop to a late stage where they arrest mainly as 649 trophozoites, suggesting EXP1 does not function directly in egress (Figure 5). Arrested 650 parasites produced additional membrane structures in the host cytosol, likely reflecting 651 alterations in the TVN, an extension of the PVM thought to function in nutrient acquisition to 652 which EXP1 is partially localized (65-67). Parasites depleted of EXP1 also displayed an 653 enlarged PV lumen that was often filled with hemoglobin-containing membrane-bound 654 structures (Figure 6D). These structures are reminiscent of TVN enclosures of host cytosol (68-655 71) except that they are present within the PVM, and may thus reflect abnormalities in TVN 656 formation. Alternatively, these structures might result from irregularities of the central cavity, a 657 poorly understood structure open to the host cytosol and often observed as an indentation at the 658 parasite periphery (72, 73). Notably, cytostomes remained visible in EXP1 knockdown parasites 659 despite these alterations (Figure 6C,D, arrowheads). Defects in uptake of host cell cytosol 660 would be expected to manifest in the terminal pathway compartment, the digestive vacuole, as 661 observed with a VPS45 conditional mutant that prevents delivery of endocytosed material to the 662 digestive vacuole, severely limiting hemoglobin degradation (74). However, digestive vacuoles 663 appeared largely normal and hemozoin crystal formation was not grossly altered following EXP1 664 knockdown (Figure 5A and 6), suggesting uptake of host cell cytosol is not critically impacted. 665 Reduction of EXP1 levels also produced a striking change in EXP2 distribution with 666 EXP2 often concentrated to one or two discrete points along the PVM (Figure 7). EXP2 forms a 667 dual functional pore in the PVM that is required for small molecule transport and effector protein 668 translocation (10, 14). Although we did not directly evaluate small molecule transport at the 669 PVM, the fact that EXP2 pore function in protein export within PTEX is not impaired and that

670 EXP1 knockdown parasites do not arrest at an earlier developmental stage suggests that EXP2-671 dependent small molecule transport is unlikely to be critically impacted.

672 A striking feature of the PVM is its intimate apposition to the PPM but the basis for 673 maintaining this proximity is not known. When protein export is disabled by inactivating PTEX 674 components, tubular distensions form and project into the host cell (14). This PV swelling 675 appears to result from accumulation of blocked exported proteins within the PV but occurs at 676 discrete points, with PVM-PPM anchoring still largely maintained (14). In contrast, the more 677 uniform separation between PVM and PPM observed upon EXP1 knockdown may indicate a 678 role for EXP1 in maintaining this intimate membrane connection, loss of which might result in 679 broad disorganization of PVM patterning that is somehow tied to the connection with the PPM. 680 Developmental arrest in late stage parasites may reflect a critical, though currently unclear, 681 requirement for maintenance of this connection for proper completion of schizogony. In 682 conclusion, our results call into question the importance of GST activity for EXP1 function and 683 reveal a critical requirement for EXP1 in maintaining proper PVM/TVN organization.

684

685 Acknowledgements

This work was supported by National Institutes of Health grant HL133453 to J.R.B. The funders
had no role in study design, data collection and interpretation, or the decision to submit the work
for publication.

We thank J. Aguiar for the EXP1 antibody, J. McBride, D. Cavanaugh and EMRR for the EXP2
antibody, C. Braun-Breton for the SBP1 antibody, W. Beatty, K. Hausmann and the WUSTL
Molecular Microbiology Imaging Facility for assistance with electron microscopy, B. Vaupel for
assistance with molecular cloning and E. Istvan and A. Polino for assistance with parasite
culture.

694

695 Author contributions

696 T.N., J.M.B., D.E.G. and J.R.B. conceived and designed the experiments. T.N.,	, J.M.B., S.	.R.,
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Y.J. and J.R.B. performed the experiments. T.N., J.M.B., S.R., Y.J., J.A.W., D.E.G. and J.R.B.

analyzed the data. J.R.B. oversaw the project and wrote the manuscript. All authors discussedand edited the manuscript.

700

701 Figures Legends

702

703 Figure 1: BioID2 identifies the protein contents of the PV/PVM. (A) Schematic showing C-

terminal BioID2-3xHA fusion to the endogenous *exp2* and *hsp101* genes. (B)

705 Immunofluorescence assay of parental, EXP2-BioID2 and HSP101-BioID2 parasites grown in

706 media supplemented with 200 μM biotin. Scale bars are 5 μm. (C) Western blot of parental,

707 EXP2-BioID2 and HSP101-BioID2 lines grown in regular RPMI (which contains 820 nM biotin,

designated "-") or 18 hours in RPMI supplemented with 200 µM biotin (designated "+").

709 Molecular weights after signal peptide cleavage are predicted to be 61.2 kDa for EXP2-BioID2-

710 3xHA and 130 kDa for HSP101-BioID2-3xHA. Arrowheads from top to bottom indicate bands

expected to correspond to untagged PTEX150 and HSP101. Arrow indicates prominent band at

~23 kDa. (D) Ven diagram summarizing overlapping and distinct proteins detected in EXP2-

713 BioID2 and HSP101-BioID2 datasets. All proteins identified in untagged negative controls were

removed from the BioID2 datasets and remaining proteins that were present in both

independent replicates of EXP2-BioID2 or HSP101-BioID2 were used to generate the diagram.

716 (E) Western blot as in (C) probed with anti-EXP1 to show correspondence with prominent ~23

kDa band. Molecular weight after signal peptide cleavage is predicted to be 14.7 kDa for EXP1.

Note that EXP1 is observed to migrate at a higher molecular weight than predicted.

719

Figure 2: Genome editing of the *exp1* locus with Cpf1. (A) Schematic showing strategy for
 double homologous recombination repair of double-strand breaks mediated by Cas9 or Cpf1 at

722 the 3' end of exp1. Individual gRNA target sequences are shown in the zoomed view in the dashed box. Exon sequences are shown in uppercase while intron and UTR sequences are 723 724 shown in lowercase. 3' UTR, 3' untranslated region; yDHODH, yeast dihydroorotate 725 dehydrogenase; PAM, protospacer adjustment motif. (B) Diagnostic PCR with primers indicated 726 in the schematic and Western blot showing successful integration at the 3' end of exp1 727 mediated by either Cas9 or Cpf1. Aldolase serves as a loading control. Molecular weight after 728 signal peptide cleavage is predicted to be 20.9 kDa for EXP1-3xHA-GFP11. Note that EXP1 729 and derivative fusions are observed to migrate at a higher molecular weight than predicted.

730

731 Figure 3: Lethal EXP1 knockdown with a dual aptamer strategy. (A) Schematic of modified exp1 locus and TDA control of translation in EXP1^{apt} parasites. 2A, Thosea asigna virus 2A skip 732 733 peptide; BSD, blasticidin-S deaminase. (B) Western blot of parental parasite line grown without 734 aTc and EXP1^{apt} parasites grown 48 hours with or without aTc. Aldolase serves as a loading 735 control. Molecular weights after signal peptide cleavage are predicted to be 14.7 kDa for EXP1 736 and 18 kDa for EXP1-3xHA. Note that EXP1 and derivative fusions are observed to migrate at a higher molecular weight than predicted. Results are representative of three independent 737 experiments. (C) Growth analysis of parental and EXP1^{apt} parasites with or without aTc. Results 738 739 from one experiment with three technical replicates plotted are shown. The scatter plot shows 740 the slope of the line fitted to the mean of log2-transformed parasitemias for each of three 741 independent experiments. Error bars indicate SEM. P values were determined by an unpaired, 742 two-sided Student's t-test.

743

Figure 4: EXP1 *in vivo* function is ablated by a bulky C-terminal fusion but not by
mutation of a residue important for GST activity *in vitro*. (A) Schematic showing
complementing versions of EXP1 introduced into EXP1^{apt}. (B) Western blot of parent, EXP1^{apt}
and complemented lines. Two independently complemented lines were generated with each

748	construct and are designated A or B. Aldolase serves as a loading control. Molecular weights
749	after signal peptide cleavage are predicted to be 18 kDa for EXP1-3xHA, 19.2 kDa for EXP1-
750	WT-3xMYC and EXP1-R70T-3xMYC and 46.2 kDa for EXP1-mNG-3xMYC. Note that EXP1 and
751	derivative fusions are observed to migrate at a higher molecular weight than predicted. (C)
752	Growth analysis of complemented EXP1 ^{apt} lines with or without aTc. The scatter plot shows the
753	slope of the line fitted to the mean of log2-transformed parasitemias for four (EXP1-WT-
754	3xMYC), five (EXP1-R70T-3xMYC) or six (EXP1-mNG-3xMYC) independent experiments.
755	Means from independently generated lines complemented with the same version of EXP1 were
756	pooled and are distinguished by different symbols (circles and triangles). Error bars indicate
757	SEM. P values were determined by an unpaired, two-sided Student's t-test. (D)
758	Immunofluorescence assay of EXP1 ^{apt} complemented lines. Scale bars are 5 μ m.
759	
760	Figure 5: Depletion of EXP1 results in late cycle arrest but does not impact protein
700	
761	export. (A-B) Quantification of parasite stages of development from Giemsa-stained thin
761	export. (A-B) Quantification of parasite stages of development from Giemsa-stained thin
761 762	export. (A-B) Quantification of parasite stages of development from Giemsa-stained thin smears of (A) EXP1 ^{apt} and (B) EXP2 ^{apt} parasites synchronized to a 10-hour window and grown
761 762 763	export. (A-B) Quantification of parasite stages of development from Giemsa-stained thin smears of (A) EXP1 ^{apt} and (B) EXP2 ^{apt} parasites synchronized to a 10-hour window and grown with or without aTc. Time 0 hours indicates the point of aTc removal at which purified late
761 762 763 764	export. (A-B) Quantification of parasite stages of development from Giemsa-stained thin smears of (A) EXP1 ^{apt} and (B) EXP2 ^{apt} parasites synchronized to a 10-hour window and grown with or without aTc. Time 0 hours indicates the point of aTc removal at which purified late trophozoites (~32-42 hours post-invasion) were mixed with fresh uninfected RBCs. Results from
761 762 763 764 765	export. (A-B) Quantification of parasite stages of development from Giemsa-stained thin smears of (A) EXP1 ^{apt} and (B) EXP2 ^{apt} parasites synchronized to a 10-hour window and grown with or without aTc. Time 0 hours indicates the point of aTc removal at which purified late trophozoites (~32-42 hours post-invasion) were mixed with fresh uninfected RBCs. Results from one experiment performed in technical triplicate are shown and are representative of two
761 762 763 764 765 766	export. (A-B) Quantification of parasite stages of development from Giemsa-stained thin smears of (A) EXP1 ^{apt} and (B) EXP2 ^{apt} parasites synchronized to a 10-hour window and grown with or without aTc. Time 0 hours indicates the point of aTc removal at which purified late trophozoites (~32-42 hours post-invasion) were mixed with fresh uninfected RBCs. Results from one experiment performed in technical triplicate are shown and are representative of two independent experiments. Error bars indicate SD. Representative images of the majority
761 762 763 764 765 766 767	export. (A-B) Quantification of parasite stages of development from Giemsa-stained thin smears of (A) EXP1 ^{apt} and (B) EXP2 ^{apt} parasites synchronized to a 10-hour window and grown with or without aTc. Time 0 hours indicates the point of aTc removal at which purified late trophozoites (~32-42 hours post-invasion) were mixed with fresh uninfected RBCs. Results from one experiment performed in technical triplicate are shown and are representative of two independent experiments. Error bars indicate SD. Representative images of the majority parasite population at each time point are shown. Scale bars are 2 µm. (C-D)
761 762 763 764 765 766 767 768	export. (A-B) Quantification of parasite stages of development from Giemsa-stained thin smears of (A) EXP1 ^{apt} and (B) EXP2 ^{apt} parasites synchronized to a 10-hour window and grown with or without aTc. Time 0 hours indicates the point of aTc removal at which purified late trophozoites (~32-42 hours post-invasion) were mixed with fresh uninfected RBCs. Results from one experiment performed in technical triplicate are shown and are representative of two independent experiments. Error bars indicate SD. Representative images of the majority parasite population at each time point are shown. Scale bars are 2 µm. (C-D) Immunofluorescence assay showing SBP1, which is exported to the Maurer's clefts, in (C)
761 762 763 764 765 766 767 768 769	export. (A-B) Quantification of parasite stages of development from Giemsa-stained thin smears of (A) EXP1 ^{apt} and (B) EXP2 ^{apt} parasites synchronized to a 10-hour window and grown with or without aTc. Time 0 hours indicates the point of aTc removal at which purified late trophozoites (~32-42 hours post-invasion) were mixed with fresh uninfected RBCs. Results from one experiment performed in technical triplicate are shown and are representative of two independent experiments. Error bars indicate SD. Representative images of the majority parasite population at each time point are shown. Scale bars are 2 µm. (C-D) Immunofluorescence assay showing SBP1, which is exported to the Maurer's clefts, in (C) EXP1 ^{apt} and (D) EXP2 ^{apt} parasites. Quantification of SBP1 export beyond the PVM (both
761 762 763 764 765 766 767 768 769 770	export. (A-B) Quantification of parasite stages of development from Giemsa-stained thin smears of (A) EXP1 ^{apt} and (B) EXP2 ^{apt} parasites synchronized to a 10-hour window and grown with or without aTc. Time 0 hours indicates the point of aTc removal at which purified late trophozoites (~32-42 hours post-invasion) were mixed with fresh uninfected RBCs. Results from one experiment performed in technical triplicate are shown and are representative of two independent experiments. Error bars indicate SD. Representative images of the majority parasite population at each time point are shown. Scale bars are 2 μm. (C-D) Immunofluorescence assay showing SBP1, which is exported to the Maurer's clefts, in (C) EXP1 ^{apt} and (D) EXP2 ^{apt} parasites. Quantification of SBP1 export beyond the PVM (both EXP1 ^{apt} and EXP2 ^{apt} contain a 3xFLAG tag on HSP101 which was used as a marker for the
761 762 763 764 765 766 767 768 769 770 771	export. (A-B) Quantification of parasite stages of development from Giemsa-stained thin smears of (A) EXP1 ^{apt} and (B) EXP2 ^{apt} parasites synchronized to a 10-hour window and grown with or without aTc. Time 0 hours indicates the point of aTc removal at which purified late trophozoites (~32-42 hours post-invasion) were mixed with fresh uninfected RBCs. Results from one experiment performed in technical triplicate are shown and are representative of two independent experiments. Error bars indicate SD. Representative images of the majority parasite population at each time point are shown. Scale bars are 2 μm. (C-D) Immunofluorescence assay showing SBP1, which is exported to the Maurer's clefts, in (C) EXP1 ^{apt} and (D) EXP2 ^{apt} parasites. Quantification of SBP1 export beyond the PVM (both EXP1 ^{apt} and EXP2 ^{apt} contain a 3xFLAG tag on HSP101 which was used as a marker for the PVM) is shown as MFI of SBP1 within the host compartment or as the number of SBP1-positive

corresponding DIC image. Data are pooled from three independent experiments and *n* is the
number of individual parasite-infected RBCs. Boxes and whiskers delineate 25th-75th and 10th90th percentiles, respectively. P values were determined by an unpaired, two-sided Student's ttest. MFI, mean fluorescence intensity; AU, arbitrary units. Scale bars are 2 µm.

779 Figure 6: Depletion of EXP1 results in PV/PVM morphological abnormalities. (A-E) TEM 780 visualization of parasite ultrastructure in parent and EXP1^{apt} parasites grown 48 hours with or 781 without aTc. Images are shown of (A-D) trophozoites and (E) segmented schizonts. Dashed 782 boxes indicate enlarged areas shown to the right. Arrows indicate PVM and PPM. Arrowheads 783 indicate cytostomes. Double arrowheads indicate abnormal membrane-enclosed structures 784 filled with host cytosol in the PV lumen. Results are representative of two independent 785 experiments. Quantification of morphological abnormalities is shown in Table 2. Scale bars are 786 500 nm. (F) Live fluorescence imaging of magnet-purified EXP1^{apt}::PV-mRuby3 parasites grown 787 48 hours with or without aTc. The dashed lines indicate the boundary of infected RBC traced 788 from the corresponding DIC image. Results are representative of three independent 789 experiments. Scale bars are 5 µm.

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791 Figure 7: Depletion of EXP1 alters EXP2 distribution in the PVM. (A) Western blot of parental parasite line grown without aTc and EXP1^{apt} parasites grown 48 hours with or without 792 793 aTc. Aldolase serves as a loading control. Molecular weights after signal peptide cleavage are 794 predicted to be 30.8 kDa for EXP2 and 18 kDa for EXP1-3xHA. Note that EXP1 and derivative 795 fusions are observed to migrate at a higher molecular weight than predicted. (B) Live fluorescent images of EXP2-mNG and EXP1^{apt}::EXP2-mNG parasites grown 48 hours with or 796 797 without aTc. Merged images include EXP2-mNG signal in green together with DIC and BODIPY 798 TR Ceramide signal in red used to trace the PVM. Scale bars are 2 µm. The corresponding 799 histograms of EXP2-mNG signal along the PVM trace are shown for each cell. Dashed line is

800	the mean of the minimum and maximum EXP2-mNG signal intensity. (C) Quantification of the
801	number of discrete peaks of EXP2-mNG signal exceeding the mean signal threshold as shown
802	in (B) for EXP2-mNG and EXP1 ^{apt} ::EXP2-mNG parasites grown 48 hours with or without aTc.
803	Data are pooled from three independent experiments and n is the number of individual
804	parasites. Boxes and whiskers delineate 25 th -75 th and 10 th -90 th percentiles, respectively. P
805	values were determined by an unpaired, two-sided Student's t-test. (D) Graph showing the
806	minimum distance along the PVM (given as a percent of total PVM length) containing the
807	indicated amounts of EXP2-mNG fluorescent signal (given as percent of total EXP2-mNG signal
808	per trace) in EXP2-mNG and EXP1 ^{apt} ::EXP2-mNG parasites grown 48 hours with or without
809	aTc. Data points are means from three independent experiments fitted to a smooth line.
810	Figure S1: Generation of the EXP2-BioID2 fusion line and expression profiles of select
010	
811	hits. (A) Western blot of parent and EXP2-BioID2-3xHA parasites. Molecular weights after
812	signal peptide cleavage are predicted to be 30.8 kDa for EXP2 and 61.2 kDa for EXP2-BioID2-
813	3xHA. (B) Transcript fold change throughout intraerythrocytic development assessed by
814	transcriptomic analysis of synchronized P. falciparum 3D7 parasites for EXP2, HSP101, EXP1
815	and select ETRAMP family members. Data are from RNAseq analysis by Otto and colleagues
816	(75). Genes with similar expression pattern to exp2 are shown in color. (C) Ven diagram
817	summarizing overlapping and distinct proteins detected in EXP2-BioID2, HSP101-BioID2 and
818	SP-GFP-BirA* datasets (50). Data from Khosh-Naucke et al 2018 was processed in the same
819	way as the BioID2 datasets in this study by pooling SP-GFP-BirA* mass spectrometry datasets
820	obtained from saponin supernatant and pellet fractions and removing all proteins identified in
821	3D7 negative controls.

822

823 Figure S2: Genome editing of exp1, etramp10.2 and etramp5 with Cpf1. (A) Western blot of parent and EXP1-3xHA-GFP11 fusion parasites generated with AsCpf1 or LbCpf1 and gRNA2 824 825 as shown in Figure 2A but expressed from the selectable pUF-AsCpf1 or pUF-LbCpf1 plasmids 826 that contain a yDHODH cassette. Aldolase serves as a loading control. Molecular weight after 827 signal peptide cleavage is predicted to be 20.9 kDa for EXP1-3xHA-GFP11. Note that EXP1 828 and derivative fusions are observed to migrate at a higher molecular weight than predicted. (B) 829 Schematic showing strategy for double homologous recombination repair of double-strand 830 breaks mediated by Cpf1 at the 3' end of the etramp10.2 and etramp5 genes to install a 3xHA 831 fusion and 3' TetR-DOZI-aptamers. 3' UTR, 3' untranslated region; yDHODH, yeast 832 dihydroorotate dehydrogenase. (C,D) Sequence of the 3' end of etramp10.2 or etramp5 with 833 Cpf1 gRNA target indicated. Successful integration mediated by AsCpf1 to generate a 3' fusion 834 to 3xHA is shown by diagnostic PCR with primers indicated in the schematic and by Western 835 blot of ETRAMP10.2^{apt} and ETRAMP5^{apt} parasites grown with or without 1µM aTc for 96 hours. 836 EXP2 serves as a loading control. Molecular weight after signal peptide cleavage is predicted to 837 be 39.6 kDa for ETRAMP10.2-3xHA and 19.6 kDa for ETRAMP5-3xHA. Similar to EXP1, both 838 ETRAMP10.2-3xHA and ETRAMP5-3xHA are observed to migrate at a higher molecular weight 839 than predicated. PAM, protospacer adjustment motif.

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Figure S3: Generation of EXP1^{apt} parasites and analysis of DiCre-mediated *exp1* excision. 841 842 (A) Schematic for strategy used to replace the endogenous exp1 coding sequence with a re-843 coded version of *exp1* with TDA and DiCre elements by Cpf1 editing and double homologous 844 recombination. (B) Diagnostic PCR with primers indicated in the schematic in (A) showing successful integration at the 5' and 3' ends of the *exp1* locus in EXP1^{apt} parasites. The absence 845 846 of the product in EXP1^{apt} using primers P65/P67 is likely due to the very large amplicon size. (C) Schematic of *exp1* locus following excision between *loxP* sites by DiCre. The pEXP1^{apt} plasmid 847 848 was designed so that DiCre excision of the modified locus would place the promoter-less

- 849 *mruby3* coding sequence under the control of the endogenous *exp1* promoter. (D) Growth
- analysis of parental and EXP1^{apt} parasites with or without 1 µM aTc or with and without 100 µM

851 or 250 μM rapamycin. The scatter plot shows the slope of the line fitted to the mean of log2-

- transformed parasitemias for each of three independent experiments. Error bars indicate SEM.
- (E) Diagnostic PCR for *exp1* excision by DiCre with primers indicated in the schematics.
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855 References

- Lingelbach K, Joiner KA. 1998. The parasitophorous vacuole membrane surrounding
 Plasmodium and Toxoplasma: an unusual compartment in infected cells. J Cell Sci 111 (
 Pt 11):1467-75.
- Sherling ES, van Ooij C. 2016. Host cell remodeling by pathogens: the exomembrane
 system in Plasmodium-infected erythrocytes. FEMS Microbiol Rev 40:701-21.
- Aikawa M, Hepler PK, Huff CG, Sprinz H. 1966. The feeding mechanism of avian
 malarial parasites. J Cell Biol 28:355-73.
- Francis SE, Sullivan DJ, Goldberg DE. 1997. Hemoglobin metabolism in the malaria
 parasite Plasmodium falciparum. Annu Rev Microbiol 51:97-123.
- Lew VL, Tiffert T, Ginsburg H. 2003. Excess hemoglobin digestion and the osmotic
 stability of Plasmodium falciparum-infected red blood cells. Blood 101:4189-94.
- 867 6. Hanssen E, Knoechel C, Dearnley M, Dixon MW, Le Gros M, Larabell C, Tilley L. 2012.
 868 Soft X-ray microscopy analysis of cell volume and hemoglobin content in erythrocytes
 869 infected with asexual and sexual stages of Plasmodium falciparum. J Struct Biol
 870 177:224-32.
- de Koning-Ward TF, Gilson PR, Boddey JA, Rug M, Smith BJ, Papenfuss AT, Sanders
 PR, Lundie RJ, Maier AG, Cowman AF, Crabb BS. 2009. A newly discovered protein
 export machine in malaria parasites. Nature 459:945-9.
- 874 8. Elsworth B, Matthews K, Nie CQ, Kalanon M, Charnaud SC, Sanders PR, Chisholm SA,
 875 Counihan NA, Shaw PJ, Pino P, Chan JA, Azevedo MF, Rogerson SJ, Beeson JG,
 876 Crabb BS, Gilson PR, de Koning-Ward TF. 2014. PTEX is an essential nexus for protein
 877 export in malaria parasites. Nature 511:587-91.
- Beck JR, Muralidharan V, Oksman A, Goldberg DE. 2014. PTEX component HSP101
 mediates export of diverse malaria effectors into host erythrocytes. Nature 511:592-5.
- 88010.Ho CM, Beck JR, Lai M, Cui Y, Goldberg DE, Egea PF, Zhou ZH. 2018. Malaria parasite881translocon structure and mechanism of effector export. Nature 561:70-75.
- Behde N, Hinrichs C, Montilla I, Charpian S, Lingelbach K, Przyborski JM. 2009. Protein
 unfolding is an essential requirement for transport across the parasitophorous vacuolar
 membrane of Plasmodium falciparum. Mol Microbiol 71:613-28.
- Besai SA, Krogstad DJ, McCleskey EW. 1993. A nutrient-permeable channel on the intraerythrocytic malaria parasite. Nature 362:643-6.
- Basi SA, Rosenberg RL. 1997. Pore size of the malaria parasite's nutrient channel.
 Proc Natl Acad Sci U S A 94:2045-9.
- 889 14. Garten M, Nasamu AS, Niles JC, Zimmerberg J, Goldberg DE, Beck JR. 2018. EXP2 is
 890 a nutrient-permeable channel in the vacuolar membrane of Plasmodium and is essential
 891 for protein export via PTEX. Nat Microbiol 3:1090-1098.

- Signathan Straight St
- Ansorge I, Benting J, Bhakdi S, Lingelbach K. 1996. Protein sorting in Plasmodium
 falciparum-infected red blood cells permeabilized with the pore-forming protein
 streptolysin O. Biochem J 315 (Pt 1):307-14.
- Spielmann T, Fergusen DJ, Beck HP. 2003. etramps, a new Plasmodium falciparum
 gene family coding for developmentally regulated and highly charged membrane
 proteins located at the parasite-host cell interface. Mol Biol Cell 14:1529-44.
- 901 18. Spielmann T, Beck HP. 2000. Analysis of stage-specific transcription in plasmodium
 902 falciparum reveals a set of genes exclusively transcribed in ring stage parasites. Mol
 903 Biochem Parasitol 111:453-8.
- Surrà C, Pace T, Franke-Fayard BM, Picci L, Bertuccini L, Ponzi M. 2012. Erythrocyte
 remodeling in Plasmodium berghei infection: the contribution of SEP family members.
 Traffic 13:388-99.
- 20. Lisewski AM, Quiros JP, Ng CL, Adikesavan AK, Miura K, Putluri N, Eastman RT,
 Scanfeld D, Regenbogen SJ, Altenhofen L, Llinás M, Sreekumar A, Long C, Fidock DA,
 Lichtarge O. 2014. Supergenomic network compression and the discovery of EXP1 as a
 glutathione transferase inhibited by artesunate. Cell 158:916-928.
- 911 21. Glushakova S, Beck JR, Garten M, Busse BL, Nasamu AS, Tenkova-Heuser T, Heuser
 912 J, Goldberg DE, Zimmerberg J. 2018. Rounding precedes rupture and breakdown of
 913 vacuolar membranes minutes before malaria parasite egress from erythrocytes. Cell
 914 Microbiol 20:e12868.
- 915 22. Spielmann T, Gardiner DL, Beck HP, Trenholme KR, Kemp DJ. 2006. Organization of
 916 ETRAMPs and EXP-1 at the parasite-host cell interface of malaria parasites. Mol
 917 Microbiol 59:779-94.
- 918 23. Wickham ME, Rug M, Ralph SA, Klonis N, McFadden GI, Tilley L, Cowman AF. 2001.
 919 Trafficking and assembly of the cytoadherence complex in Plasmodium falciparum-920 infected human erythrocytes. EMBO J 20:5636-49.
- Riglar DT, Rogers KL, Hanssen E, Turnbull L, Bullen HE, Charnaud SC, Przyborski J,
 Gilson PR, Whitchurch CB, Crabb BS, Baum J, Cowman AF. 2013. Spatial association
 with PTEX complexes defines regions for effector export into Plasmodium falciparuminfected erythrocytes. Nat Commun 4:1415.
- Bullen HE, Charnaud SC, Kalanon M, Riglar DT, Dekiwadia C, Kangwanrangsan N, Torii
 M, Tsuboi T, Baum J, Ralph SA, Cowman AF, de Koning-Ward TF, Crabb BS, Gilson
 PR. 2012. Biosynthesis, localization, and macromolecular arrangement of the
 Plasmodium falciparum translocon of exported proteins (PTEX). J Biol Chem 287:787184.
- 26. Charnaud SC, Jonsdottir TK, Sanders PR, Bullen HE, Dickerman BK, Kouskousis B,
 Palmer CS, Pietrzak HM, Laumaea AE, Erazo AB, McHugh E, Tilley L, Crabb BS, Gilson
 PR. 2018. Spatial organization of protein export in malaria parasite blood stages. Traffic
 19:605-623.
- Adisa A, Rug M, Klonis N, Foley M, Cowman AF, Tilley L. 2003. The signal sequence of
 exported protein-1 directs the green fluorescent protein to the parasitophorous vacuole
 of transfected malaria parasites. J Biol Chem 278:6532-42.
- 837 28. Kim DI, Jensen SC, Noble KA, Kc B, Roux KH, Motamedchaboki K, Roux KJ. 2016. An
 938 improved smaller biotin ligase for BioID proximity labeling. Mol Biol Cell 27:1188-96.
- 939 29. Glushakova S, Busse BL, Garten M, Beck JR, Fairhurst RM, Goldberg DE, Zimmerberg
 940 J. 2017. Exploitation of a newly-identified entry pathway into the malaria parasite-
- 941 infected erythrocyte to inhibit parasite egress. Sci Rep 7:12250.

942 30. Ganesan SM, Morrisev JM, Ke H, Painter HJ, Laroiva K, Phillips MA, Rathod PK, Mather 943 MW, Vaidya AB. 2011. Yeast dihydroorotate dehydrogenase as a new selectable marker 944 for Plasmodium falciparum transfection. Mol Biochem Parasitol 177:29-34. 945 31. Spillman NJ, Beck JR, Ganesan SM, Niles JC, Goldberg DE. 2017. The chaperonin 946 TRiC forms an oligometric complex in the malaria parasite cytosol. Cell Microbiol 19. 947 Zetsche B, Gootenberg JS, Abudayyeh OO, Slaymaker IM, Makarova KS, Essletzbichler 32. 948 P, Volz SE, Joung J, van der Oost J, Regev A, Koonin EV, Zhang F. 2015. Cpf1 is a 949 single RNA-guided endonuclease of a class 2 CRISPR-Cas system. Cell 163:759-71. 950 33. Ganesan SM, Falla A, Goldfless SJ, Nasamu AS, Niles JC. 2016. Synthetic RNA-protein 951 modules integrated with native translation mechanisms to control gene expression in malaria parasites. Nat Commun 7:10727. 952 953 34. Chen AL, Kim EW, Toh JY, Vashisht AA, Rashoff AQ, Van C, Huang AS, Moon AS, Bell 954 HN, Bentolila LA, Wohlschlegel JA, Bradley PJ. 2015. Novel components of the 955 Toxoplasma inner membrane complex revealed by BioID. MBio 6:e02357-14. Yap A. Azevedo MF. Gilson PR. Weiss GE. O'Neill MT. Wilson DW. Crabb BS. Cowman 956 35. 957 AF. 2014. Conditional expression of apical membrane antigen 1 in Plasmodium 958 falciparum shows it is required for erythrocyte invasion by merozoites. Cell Microbiol 959 16:642-56. 960 Nasamu AS, Glushakova S, Russo I, Vaupel B, Oksman A, Kim AS, Fremont DH, Tolia 36. 961 N, Beck JR, Meyers MJ, Niles JC, Zimmerberg J, Goldberg DE. 2017. Plasmepsins IX 962 and X are essential and druggable mediators of malaria parasite egress and invasion. 963 Science 358:518-522. 964 37. Nkrumah LJ, Muhle RA, Moura PA, Ghosh P, Hatfull GF, Jacobs WR, Jr., Fidock DA. 965 2006. Efficient site-specific integration in Plasmodium falciparum chromosomes mediated by mycobacteriophage Bxb1 integrase. Nat Methods 3:615-21. 966 967 38. Wohlschlegel JA. 2009. Identification of SUMO-conjugated proteins and their SUMO 968 attachment sites using proteomic mass spectrometry. Methods Mol Biol 497:33-49. 969 39. Florens L. Carozza MJ. Swanson SK. Fournier M. Coleman MK. Workman JL. 970 Washburn MP. 2006. Analyzing chromatin remodeling complexes using shotgun proteomics and normalized spectral abundance factors. Methods 40:303-11. 971 972 40. Hall R, McBride J, Morgan G, Tait A, Zolg JW, Walliker D, Scaife J. 1983. Antigens of 973 the erythrocytes stages of the human malaria parasite Plasmodium falciparum detected 974 by monoclonal antibodies. Mol Biochem Parasitol 7:247-65. 975 41. Blisnick T, Morales Betoulle ME, Barale JC, Uzureau P, Berry L, Desroses S, Fujioka H, 976 Mattei D, Braun Breton C. 2000. Pfsbp1, a Maurer's cleft Plasmodium falciparum protein, 977 is associated with the erythrocyte skeleton. Mol Biochem Parasitol 111:107-21. 978 42. Batinovic S, McHugh E, Chisholm SA, Matthews K, Liu B, Dumont L, Charnaud SC, 979 Schneider MP, Gilson PR, de Koning-Ward TF, Dixon MWA, Tilley L. 2017. An exported 980 protein-interacting complex involved in the trafficking of virulence determinants in 981 Plasmodium-infected erythrocytes. Nat Commun 8:16044. 982 Hope IA, Mackay M, Hyde JE, Goman M, Scaife J. 1985. The gene for an exported 43. 983 antigen of the malaria parasite Plasmodium falciparum cloned and expressed in 984 Escherichia coli. Nucleic Acids Res 13:369-79. 985 44. Gardner MJ, Hall N, Fung E, White O, Berriman M, Hyman RW, Carlton JM, Pain A, 986 Nelson KE, Bowman S, Paulsen IT, James K, Eisen JA, Rutherford K, Salzberg SL, 987 Craig A, Kyes S, Chan MS, Nene V, Shallom SJ, Suh B, Peterson J, Angiuoli S, Pertea 988 M, Allen J, Selengut J, Haft D, Mather MW, Vaidya AB, Martin DM, Fairlamb AH, 989 Fraunholz MJ, Roos DS, Ralph SA, McFadden GI, Cummings LM, Subramanian GM, 990 Mungall C, Venter JC, Carucci DJ, Hoffman SL, Newbold C, Davis RW, Fraser CM, 991 Barrell B. 2002. Genome sequence of the human malaria parasite Plasmodium 992 falciparum. Nature 419:498-511.

993	45.	Kim D, Kim J, Hur JK, Been KW, Yoon SH, Kim JS. 2016. Genome-wide analysis
994		reveals specificities of Cpf1 endonucleases in human cells. Nat Biotechnol 34:863-8.
995	46.	Kleinstiver BP, Tsai SQ, Prew MS, Nguyen NT, Welch MM, Lopez JM, McCaw ZR,
996		Aryee MJ, Joung JK. 2016. Genome-wide specificities of CRISPR-Cas Cpf1 nucleases
997		in human cells. Nat Biotechnol 34:869-74.
998	47.	Iriko H, Ishino T, Otsuki H, Ito D, Tachibana M, Torii M, Tsuboi T. 2018. Plasmodium
999		falciparum Exported Protein 1 is localized to dense granules in merozoites. Parasitol Int
1000		67:637-639.
1001	48.	Trinkle-Mulcahy L. 2019. Recent advances in proximity-based labeling methods for
1002		interactome mapping. F1000Res 8.
1003	49.	Kehrer J, Frischknecht F, Mair GR. 2016. Proteomic Analysis of the Plasmodium berghei
1004		Gametocyte Egressome and Vesicular bioID of Osmiophilic Body Proteins Identifies
1005		Merozoite TRAP-like Protein (MTRAP) as an Essential Factor for Parasite Transmission.
1006	50	Mol Cell Proteomics 15:2852-62.
1007	50.	Khosh-Naucke M, Becker J, Mesen-Ramirez P, Kiani P, Birnbaum J, Frohlke U,
1008		Jonscher E, Schluter H, Spielmann T. 2017. Identification of novel parasitophorous
1009	F 4	vacuole proteins in P. falciparum parasites using BioID. Int J Med Microbiol.
1010	51.	Schnider CB, Bausch-Fluck D, Bruhlmann F, Heussler VT, Burda PC. 2018. BiolD
1011		Reveals Novel Proteins of the Plasmodium Parasitophorous Vacuole Membrane.
1012	52.	mSphere 3. Bougher M.L. Chech S. Zhang L. Lei A. Jang S.W. Ju A. Zhang S. Wang Y. Balah S.A.
1013 1014	52.	Boucher MJ, Ghosh S, Zhang L, Lal A, Jang SW, Ju A, Zhang S, Wang X, Ralph SA, Zou J, Elias JE, Yeh E. 2018. Integrative proteomics and bioinformatic prediction enable
1014		a high-confidence apicoplast proteome in malaria parasites. PLoS Biol 16:e2005895.
1015	53.	McBride JS, Walliker D, Morgan G. 1982. Antigenic diversity in the human malaria
1010	55.	parasite Plasmodium falciparum. Science 217:254-7.
1017	54.	Hope IA, Hall R, Simmons DL, Hyde JE, Scaife JG. 1984. Evidence for immunological
1010	01.	cross-reaction between sporozoites and blood stages of a human malaria parasite.
1020		Nature 308:191-4.
1021	55.	Maier AG, Rug M, O'Neill MT, Brown M, Chakravorty S, Szestak T, Chesson J, Wu Y,
1022		Hughes K, Coppel RL, Newbold C, Beeson JG, Craig A, Crabb BS, Cowman AF. 2008.
1023		Exported proteins required for virulence and rigidity of Plasmodium falciparum-infected
1024		human erythrocytes. Cell 134:48-61.
1025	56.	Sa ECC, Nyboer B, Heiss K, Sanches-Vaz M, Fontinha D, Wiedtke E, Grimm D,
1026		Przyborski JM, Mota MM, Prudencio M, Mueller AK. 2017. Plasmodium berghei EXP-1
1027		interacts with host Apolipoprotein H during Plasmodium liver-stage development. Proc
1028		Natl Acad Sci U S A 114:E1138-E1147.
1029	57.	Sanchez GI, Rogers WO, Mellouk S, Hoffman SL. 1994. Plasmodium falciparum:
1030		exported protein-1, a blood stage antigen, is expressed in liver stage parasites. Exp
1031		Parasitol 79:59-62.
1032	58.	Tribensky A, Graf AW, Diehl M, Fleck W, Przyborski JM. 2017. Trafficking of PfExp1 to
1033		the parasitophorous vacuolar membrane of Plasmodium falciparum is independent of
1034		protein folding and the PTEX translocon. Cell Microbiol 19.
1035	59.	Wolanin K, Fontinha D, Sanches-Vaz M, Nyboer B, Heiss K, Mueller AK, Prudêncio M.
1036		2019. A crucial role for the C-terminal domain of exported protein 1 during the mosquito
1037	00	and hepatic stages of the Plasmodium berghei life cycle. Cell Microbiol:e13088.
1038	60.	Klemba M, Beatty W, Gluzman I, Goldberg DE. 2004. Trafficking of plasmepsin II to the
1039	04	food vacuole of the malaria parasite Plasmodium falciparum. J Cell Biol 164:47-56.
1040	61.	Charnaud SC, Kumarasingha R, Bullen HE, Crabb BS, Gilson PR. 2018. Knockdown of
1041		the translocon protein EXP2 in Plasmodium falciparum reduces growth and protein
1042		export. PLoS One 13:e0204785.

Low LM, Azasi Y, Sherling ES, Garten M, Zimmerberg J, Tsuboi T, Brzostowski J, Mu J. 1043 62. 1044 Blackman MJ, Miller LH. 2019. Deletion of Plasmodium falciparum Protein RON3 Affects 1045 the Functional Translocation of Exported Proteins and Glucose Uptake. MBio 10. 1046 63. Thomas JA, Tan MSY, Bisson C, Borg A, Umrekar TR, Hackett F, Hale VL, Vizcay-1047 Barrena G, Fleck RA, Snijders AP, Saibil HR, Blackman MJ. 2018. A protease cascade regulates release of the human malaria parasite Plasmodium falciparum from host red 1048 1049 blood cells. Nat Microbiol 3:447-455. 1050 64. Pino P, Caldelari R, Mukherjee B, Vahokoski J, Klages N, Maco B, Collins CR, 1051 Blackman MJ, Kursula I, Heussler V, Brochet M, Soldati-Favre D. 2017. A multistage antimalarial targets the plasmepsins IX and X essential for invasion and egress. Science 1052 1053 358:522-528. 1054 65. Elmendorf HG, Haldar K. 1994. Plasmodium falciparum exports the Golgi marker 1055 sphingomyelin synthase into a tubovesicular network in the cytoplasm of mature 1056 erythrocytes. J Cell Biol 124:449-62. Lauer SA, Rathod PK, Ghori N, Haldar K, 1997, A membrane network for nutrient import 1057 66. 1058 in red cells infected with the malaria parasite. Science 276:1122-5. 1059 Behari R, Haldar K. 1994. Plasmodium falciparum: protein localization along a novel, 67. 1060 lipid-rich tubovesicular membrane network in infected erythrocytes. Exp Parasitol 1061 79:250-9. 1062 68. Elford BC, Cowan GM, Ferguson DJ. 1995. Parasite-regulated membrane transport 1063 processes and metabolic control in malaria-infected erythrocytes. Biochem J 308 (Pt 1064 2):361-74. 1065 69. Hanssen E, Hawthorne P, Dixon MW, Trenholme KR, McMillan PJ, Spielmann T, Gardiner DL, Tilley L. 2008. Targeted mutagenesis of the ring-exported protein-1 of 1066 Plasmodium falciparum disrupts the architecture of Maurer's cleft organelles. Mol 1067 1068 Microbiol 69:938-53. 1069 70. Hanssen E, Sougrat R, Frankland S, Deed S, Klonis N, Lippincott-Schwartz J, Tilley L. 1070 2008. Electron tomography of the Maurer's cleft organelles of Plasmodium falciparuminfected ervthrocytes reveals novel structural features. Mol Microbiol 67:703-18. 1071 71. Hanssen E, Carlton P, Deed S, Klonis N, Sedat J, DeRisi J, Tilley L. 2010. Whole cell 1072 1073 imaging reveals novel modular features of the exomembrane system of the malaria 1074 parasite, Plasmodium falciparum. Int J Parasitol 40:123-34. Gruring C, Heiber A, Kruse F, Ungefehr J, Gilberger TW, Spielmann T. 2011. 1075 72. 1076 Development and host cell modifications of Plasmodium falciparum blood stages in four dimensions. Nat Commun 2:165. 1077 1078 73. Abu Bakar N, Klonis N, Hanssen E, Chan C, Tilley L. 2010. Digestive-vacuole genesis 1079 and endocytic processes in the early intraerythrocytic stages of Plasmodium falciparum. 1080 J Cell Sci 123:441-50. 1081 74. Jonscher E, Flemming S, Schmitt M, Sabitzki R, Reichard N, Birnbaum J, Bergmann B, 1082 Höhn K, Spielmann T. 2019. PfVPS45 Is Required for Host Cell Cytosol Uptake by 1083 Malaria Blood Stage Parasites. Cell Host Microbe 25:166-173.e5. 1084 75. Otto TD, Wilinski D, Assefa S, Keane TM, Sarry LR, Bohme U, Lemieux J, Barrell B, 1085 Pain A, Berriman M, Newbold C, Llinas M. 2010. New insights into the blood-stage 1086 transcriptome of Plasmodium falciparum using RNA-Seg. Mol Microbiol 76:12-24. 76. Heiber A, Kruse F, Pick C, Gruring C, Flemming S, Oberli A, Schoeler H, Retzlaff S, 1087 1088 Mesen-Ramirez P, Hiss JA, Kadekoppala M, Hecht L, Holder AA, Gilberger TW, 1089 Spielmann T. 2013. Identification of New PNEPs Indicates a Substantial Non-PEXEL 1090 Exportome and Underpins Common Features in Plasmodium falciparum Protein Export. 1091 PLoS Pathog 9:e1003546. 1092

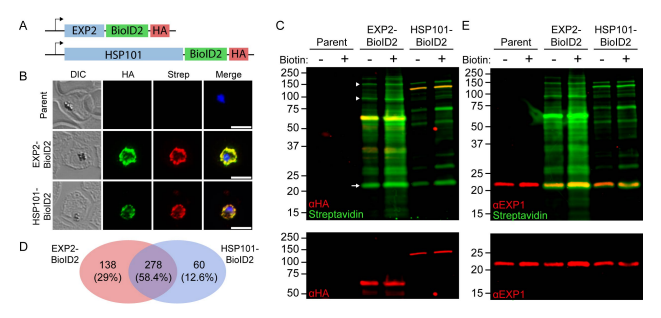


Figure 1: BioID2 identifies the protein contents of the PV/PVM. (A) Schematic showing Cterminal BioID2-3xHA fusion to the endogenous exp2 and hsp101 genes. (B) Immunofluorescence assay of parental, EXP2-BioID2 and HSP101-BioID2 parasites grown in media supplemented with 200 µM biotin. Scale bars are 5 µm. (C) Western blot of parental, EXP2-BioID2 and HSP101-BioID2 lines grown in regular RPMI (which contains 820 nM biotin, designated "-") or 18 hours in RPMI supplemented with 200 µM biotin (designated "+"). Molecular weights after signal peptide cleavage are predicted to be 61.2 kDa for EXP2-BioID2-3xHA and 130 kDa for HSP101-BioID2-3xHA. Arrowheads from top to bottom indicate bands expected to correspond to untagged PTEX150 and HSP101. Arrow indicates prominent band at ~23 kDa. (D) Ven diagram summarizing overlapping and distinct proteins detected in EXP2-BioID2 and HSP101-BioID2 datasets. All proteins identified in untagged negative controls were removed from the BioID2 datasets and remaining proteins that were present in both independent replicates of EXP2-BioID2 or HSP101-BioID2 were used to generate the diagram. (E) Western blot as in (C) probed with anti-EXP1 to show correspondence with prominent ~23 kDa band. Molecular weight after signal peptide cleavage is predicted to be 14.7 kDa for EXP1. Note that EXP1 is observed to migrate at a higher molecular weight than predicted.

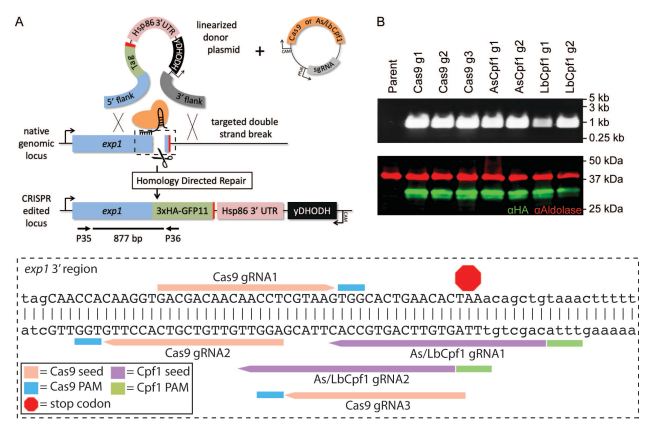


Figure 2: Genome editing of the *exp1* **locus with Cpf1.** (A) Schematic showing strategy for double homologous recombination repair of double-strand breaks mediated by Cas9 or Cpf1 at the 3' end of *exp1*. Individual gRNA target sequences are shown in the zoomed view in the dashed box. Exon sequences are shown in uppercase while intron and UTR sequences are shown in lowercase. 3' UTR, 3' untranslated region; yDHODH, yeast dihydroorotate dehydrogenase; PAM, protospacer adjustment motif. (B) Diagnostic PCR with primers indicated in the schematic and Western blot showing successful integration at the 3' end of *exp1* mediated by either Cas9 or Cpf1. Aldolase serves as a loading control. Molecular weight after signal peptide cleavage is predicted to be 20.9 kDa for EXP1-3xHA-GFP11. Note that EXP1 and derivative fusions are observed to migrate at a higher molecular weight than predicted.

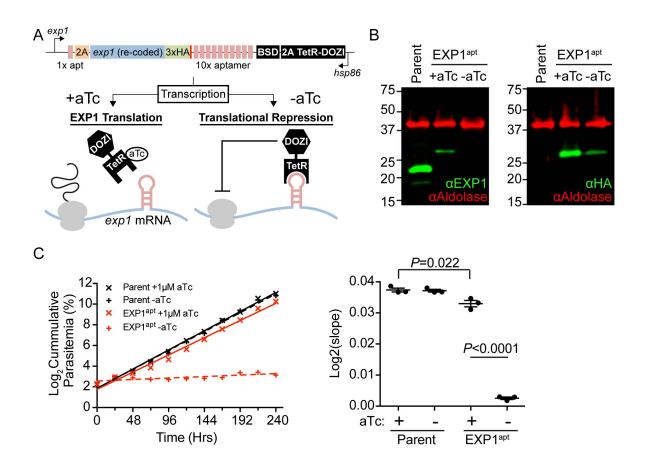


Figure 3: Lethal EXP1 knockdown with a dual aptamer strategy. (A) Schematic of modified *exp1* locus and TDA control of translation in EXP1^{apt} parasites. 2A, *Thosea asigna* virus 2A skip peptide; BSD, blasticidin-S deaminase. (B) Western blot of parental parasite line grown without aTc and EXP1^{apt} parasites grown 48 hours with or without aTc. Aldolase serves as a loading control. Molecular weights after signal peptide cleavage are predicted to be 14.7 kDa for EXP1 and 18 kDa for EXP1-3xHA. Note that EXP1 and derivative fusions are observed to migrate at a higher molecular weight than predicted. Results are representative of three independent experiments. (C) Growth analysis of parental and EXP1^{apt} parasites with or without aTc. Results from one experiment with three technical replicates plotted are shown. The scatter plot shows the slope of the line fitted to the mean of log2-transformed parasitemias for each of three independent experiments. Error bars indicate SEM. P values were determined by an unpaired, two-sided Student's t-test.

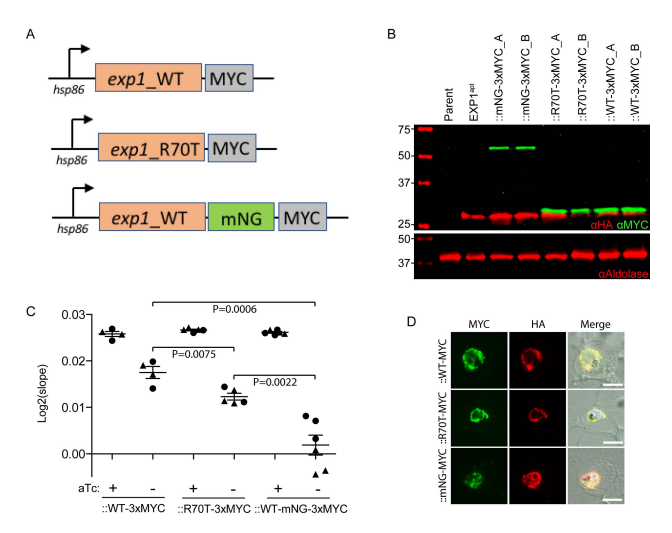


Figure 4: EXP1 *in vivo* function is ablated by a bulky C-terminal fusion but not by mutation of a residue important for GST activity *in vitro*. (A) Schematic showing complementing versions of EXP1 introduced into EXP1^{apt}. (B) Western blot of parent, EXP1^{apt} and complemented lines. Two independently complemented lines were generated with each construct and are designated A or B. Aldolase serves as a loading control. Molecular weights after signal peptide cleavage are predicted to be 18 kDa for EXP1-3xHA, 19.2 kDa for EXP1-WT-3xMYC and EXP1-R70T-3xMYC and 46.2 kDa for EXP1-mNG-3xMYC. Note that EXP1 and derivative fusions are observed to migrate at a higher molecular weight than predicted. (C) Growth analysis of complemented EXP1^{apt} lines with or without aTc. The scatter plot shows the slope of the line fitted to the mean of log2-transformed parasitemias for four (EXP1-WT-3xMYC), five (EXP1-R70T-3xMYC) or six (EXP1-mNG-3xMYC) independent experiments. Means from independently generated lines complemented with the same version of EXP1 were pooled and are distinguished by different symbols (circles and triangles). Error bars indicate SEM. P values were determined by an unpaired, two-sided Student's t-test. (D) Immunofluorescence assay of EXP1^{apt} complemented lines. Scale bars are 5 μm.

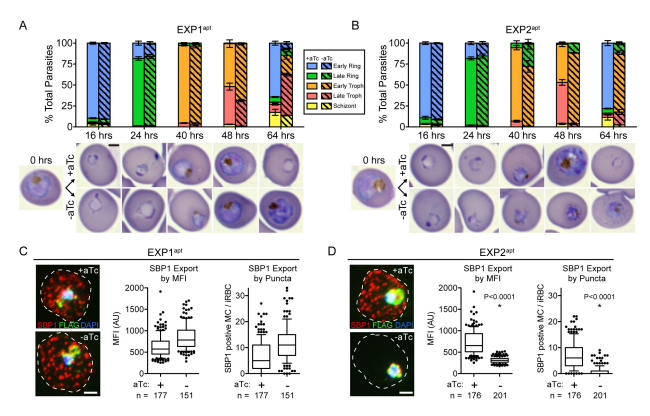


Figure 5: Depletion of EXP1 results in late cycle arrest but does not impact protein export. (A-B) Quantification of parasite stages of development from Giemsa-stained thin smears of (A) EXP1^{apt} and (B) EXP2^{apt} parasites synchronized to a 10-hour window and grown with or without aTc. Time 0 hours indicates the point of aTc removal at which purified late trophozoites (~32-42 hours post-invasion) were mixed with fresh uninfected RBCs. Results from one experiment performed in technical triplicate are shown and are representative of two independent experiments. Error bars indicate SD. Representative images of the majority parasite population at each time point are shown. Scale bars are 2 µm. (C-D) Immunofluorescence assay showing SBP1, which is exported to the Maurer's clefts, in (C) EXP1^{apt} and (D) EXP2^{apt} parasites. Quantification of SBP1 export beyond the PVM (both EXP1^{apt} and EXP2^{apt} contain a 3xFLAG tag on HSP101 which was used as a marker for the PVM) is shown as MFI of SBP1 within the host compartment or as the number of SBP1-positive puncta (Maurer's clefts) within the host compartment. Merged images of SBP1, FLAG and DAPI signal are shown. The dashed line indicates the boundary of the host RBC traced from the corresponding DIC image. Data are pooled from three independent experiments and n is the number of individual parasite-infected RBCs. Boxes and whiskers delineate 25th-75th and 10th-90th percentiles, respectively. P values were determined by an unpaired, two-sided Student's ttest. MFI, mean fluorescence intensity; AU, arbitrary units. Scale bars are 2 µm.

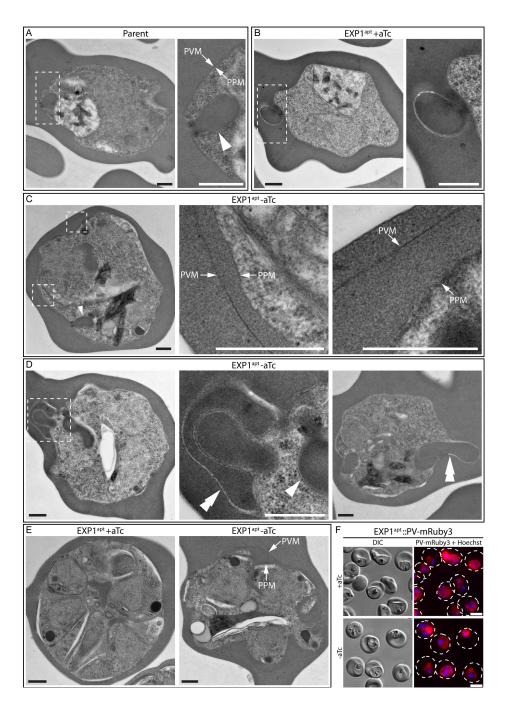


Figure 6: Depletion of EXP1 results in PV/PVM morphological abnormalities. (A-E) TEM visualization of parasite ultrastructure in parent and EXP1^{apt} parasites grown 48 hours with or without aTc. Images are shown of (A-D) trophozoites and (E) segmented schizonts. Dashed boxes indicate enlarged areas shown to the right. Arrows indicate PVM and PPM. Arrowheads indicate cytostomes. Double arrowheads indicate abnormal membrane-enclosed structures filled with host cytosol in the PV lumen. Results are representative of two independent experiments. Quantification of morphological abnormalities is shown in Table 2. Scale bars are 500 nm. (F) Live fluorescence imaging of magnet-purified EXP1^{apt}::PV-mRuby3 parasites grown 48 hours with or without aTc. The dashed lines indicate the boundary of infected RBC traced from the corresponding DIC image. Results are representative of three independent experiments. Scale bars are 5 μm.

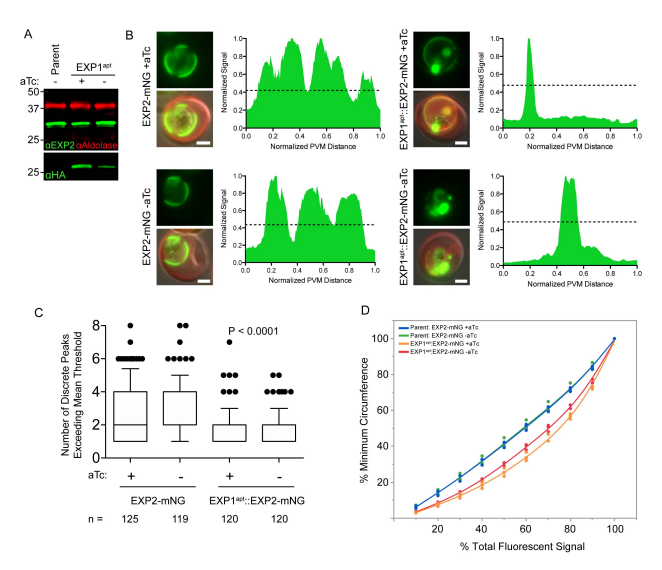


Figure 7: Depletion of EXP1 alters EXP2 distribution in the PVM. (A) Western blot of parental parasite line grown without aTc and EXP1^{apt} parasites grown 48 hours with or without aTc. Aldolase serves as a loading control. Molecular weights after signal peptide cleavage are predicted to be 30.8 kDa for EXP2 and 18 kDa for EXP1-3xHA. Note that EXP1 and derivative fusions are observed to migrate at a higher molecular weight than predicted. (B) Live fluorescent images of EXP2-mNG and EXP1^{apt}::EXP2-mNG parasites grown 48 hours with or without aTc. Merged images include EXP2-mNG signal in green together with DIC and BODIPY TR Ceramide signal in red used to trace the PVM. Scale bars are 2 µm. The corresponding histograms of EXP2-mNG signal along the PVM trace are shown for each cell. Dashed line is the mean of the minimum and maximum EXP2-mNG signal intensity. (C) Quantification of the number of discrete peaks of EXP2-mNG signal exceeding the mean signal threshold as shown in (B) for EXP2-mNG and EXP1^{apt}::EXP2-mNG parasites grown 48 hours with or without aTc. Data are pooled from three independent experiments and n is the number of individual parasites. Boxes and whiskers delineate 25th-75th and 10th-90th percentiles, respectively. P values were determined by an unpaired, two-sided Student's t-test, (D) Graph showing the minimum distance along the PVM (given as a percent of total PVM length) containing the indicated amounts of EXP2-mNG fluorescent signal (given as percent of total EXP2-mNG signal per trace) in EXP2-mNG and EXP1^{apt}::EXP2-mNG parasites grown 48 hours with or without aTc. Data points are means from three independent experiments fitted to a smooth line.

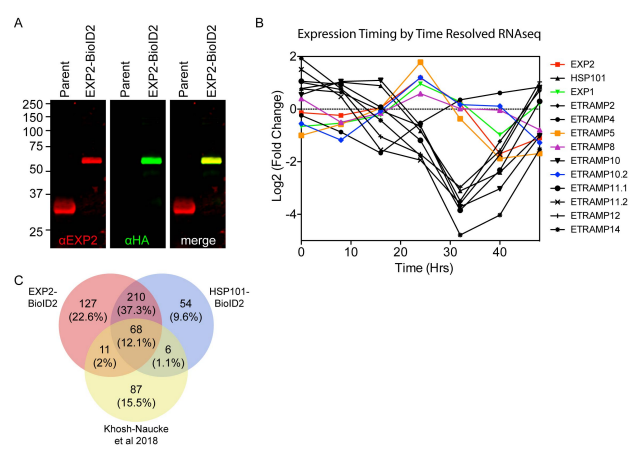


Figure S1: Generation of the EXP2-BioID2 fusion line and expression profiles of select hits. (A) Western blot of parent and EXP2-BioID2-3xHA parasites. Molecular weights after signal peptide cleavage are predicted to be 30.8 kDa for EXP2 and 61.2 kDa for EXP2-BioID2-3xHA. (B) Transcript fold change throughout intraerythrocytic development assessed by transcriptomic analysis of synchronized *P. falciparum* 3D7 parasites for EXP2, HSP101, EXP1 and select ETRAMP family members. Data are from RNAseq analysis by Otto and colleagues (75). Genes with similar expression pattern to *exp2* are shown in color. (C) Ven diagram summarizing overlapping and distinct proteins detected in EXP2-BioID2, HSP101-BioID2 and SP-GFP-BirA* datasets (50). Data from Khosh-Naucke et al 2018 was processed in the same way as the BioID2 datasets in this study by pooling SP-GFP-BirA* mass spectrometry datasets obtained from saponin supernatant and pellet fractions and removing all proteins identified in 3D7 negative controls.

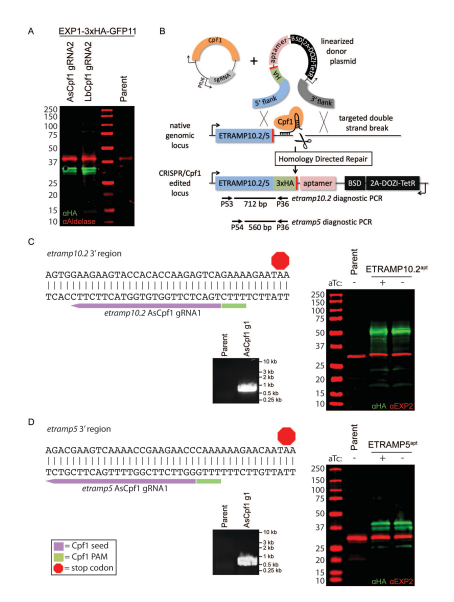


Figure S2: Genome editing of exp1, etramp10.2 and etramp5 with Cpf1. (A) Western blot of parent and EXP1-3xHA-GFP11 fusion parasites generated with AsCpf1 or LbCpf1 and gRNA2 as shown in Figure 2A but expressed from the selectable pUF-AsCpf1 or pUF-LbCpf1 plasmids that contain a vDHODH cassette. Aldolase serves as a loading control. Molecular weight after signal peptide cleavage is predicted to be 20.9 kDa for EXP1-3xHA-GFP11. Note that EXP1 and derivative fusions are observed to migrate at a higher molecular weight than predicted. (B) Schematic showing strategy for double homologous recombination repair of double-strand breaks mediated by Cpf1 at the 3' end of the etramp10.2 and etramp5 genes to install a 3xHA fusion and 3' TetR-DOZI-aptamers. 3' UTR, 3' untranslated region; yDHODH, yeast dihydroorotate dehydrogenase. (C,D) Sequence of the 3' end of etramp10.2 or etramp5 with Cpf1 gRNA target indicated. Successful integration mediated by AsCpf1 to generate a 3' fusion to 3xHA is shown by diagnostic PCR with primers indicated in the schematic and by Western blot of ETRAMP10.2^{apt} and ETRAMP5^{apt} parasites grown with or without 1µM aTc for 96 hours. EXP2 serves as a loading control. Molecular weight after signal peptide cleavage is predicted to be 39.6 kDa for ETRAMP10.2-3xHA and 19.6 kDa for ETRAMP5-3xHA. Similar to EXP1, both ETRAMP10.2-3xHA and ETRAMP5-3xHA are observed to migrate at a higher molecular weight than predicated. PAM, protospacer adjustment motif.

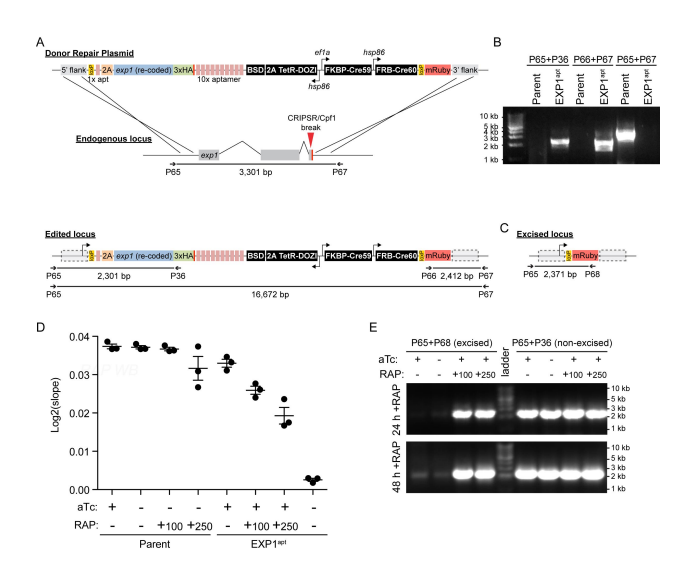


Figure S3: Generation of EXP1^{apt} **parasites and analysis of DiCre-mediated** *exp1* **excision.** (A) Schematic for strategy used to replace the endogenous *exp1* coding sequence with a recoded version of *exp1* with TDA and DiCre elements by Cpf1 editing and double homologous recombination. (B) Diagnostic PCR with primers indicated in the schematic in (A) showing successful integration at the 5' and 3' ends of the *exp1* locus in EXP1^{apt} parasites. The absence of the product in EXP1^{apt} using primers P65/P67 is likely due to the very large amplicon size. (C) Schematic of *exp1* locus following excision between *loxP* sites by DiCre. The pEXP1^{apt} plasmid was designed so that DiCre excision of the modified locus would place the promoter-less *mruby3* coding sequence under the control of the endogenous *exp1* promoter. (D) Growth analysis of parental and EXP1^{apt} parasites with or without 1 μM aTc or with and without 100 μM or 250 μM rapamycin. The scatter plot shows the slope of the line fitted to the mean of log2-transformed parasitemias for each of three independent experiments. Error bars indicate SEM. (E) Diagnostic PCR for *exp1* excision by DiCre with primers indicated in the schematics.

TABLE 1 Top 30 proteins identified in EXP2-BioID2 and HSP101-BioID2 experiments ranked by NSAF score.

	EXP2-BioID2_replicate 1		EXP2-BioID2_replicate 2		HSP1	01-BioID2_replicate 1	HSP101-BioID2_replicate 2		
#	Gene ID	Name or localization	Gene ID	Name or localization	Gene ID	Name or localization	Gene ID	Name or localization	
1	PF3D7_1471100	EXP2	PF3D7_1436300	PTEX150	PF3D7_1436300	PTEX150	PF3D7_1116800	HSP101	
2	PF3D7_1121600	EXP1	PF3D7_1033200	ETRAMP10.2	PF3D7_1116800	HSP101	PF3D7_1436300	PTEX150	
3	PF3D7_1436300	PTEX150	PF3D7_1135400	PV Localization (ref 40)	PF3D7_0721100	None	PF3D7_1212000	TPx(GI)	
4	PF3D7_1135400	PV Localization (ref 40)	PF3D7_1471100	EXP2	PF3D7_1212000	TPx(GI)	PF3D7_0721100	None	
5	PF3D7_1033200	ETRAMP10.2	PF3D7_1024800	EXP3	PF3D7_1033200	ETRAMP10.2	PF3D7_1226900	PV2	
6	PF3D7_1335100	MSP7	PF3D7_1212000	TPx(GI)	PF3D7_1135400	PVLocalization (ref 40)	PF3D7_1121600	EXP1	
7	PF3D7_1226900	PV2	PF3D7_1129100	PV1	PF3D7_0702500	Exported Protein	PF3D7_1108600	ERC	
8	PF3D7_0207500	SERA6	PF3D7_0207500	SERA6	PF3D7_1024800	EXP3	PF3D7_1471100	EXP2	
9	PF3D7_1129100	PV1	PF3D7_1335100	MSP7	PF3D7_1121600	EXP1	PF3D7_1135400	PV Localization (ref 40)	
10	PF3D7_1212000	TPx(GI)	PF3D7_1226900	PV2	PF3D7_1226900	PV2	PF3D7_0801000	PHISTC	
11	PF3D7_0721100	None	PF3D7_1345100	TRX2	PF3D7_1108600	ERC	PF3D7_1033200	ETRAMP10.2	
12	PF3D7_1312700	None	PF3D7_1121600	EXP1	PF3D7_1129100	PV1	PF3D7_1129100	PV1	
13	PF3D7_1116800	HSP101	PF3D7_0532100	ETRAMP5	PF3D7_0801000	PHISTc	PF3D7_1130100	RPL38	
14	PF3D7_1024800	EXP3	PF3D7_1105600	PTEX88	PF3D7_0629200	Partial PV Localization (ref 40)	PF3D7_0702500	Exported Protein	
15	PF3D7_1105600	PTEX88	PF3D7_0612700	PTEX88	PF3D7_1130100	RPL38	PF3D7_0625400	None	
16	PF3D7_0316300.1	PPase	PF3D7_1116700	DPAP1	PF3D7_1471100	EXP2	PF3D7_1105600	PTEX88	
17	PF3D7_1464600	UIS2	PF3D7_0316300.1	PPase	PF3D7_1105600	PTEX88	PF3D7_0202400	PTEF	
18	PF3D7_1228600	MSP9	PF3D7_0721100	None	PF3D7_1460700	RPL27	PF3D7_1404900	None	
19	PF3D7_0316300.2	PPase	PF3D7_1116800	HSP101	PF3D7_0202400	PTEF	PF3D7_0629200	Partial PV Localization (ref 40)	
20	PF3D7_0902800	SERA9	PF3D7_0404900	P41	PF3D7_1104400	Trx-mero	PF3D7_1345100	TRX2	
21	PF3D7_1345100	TRX2	PF3D7_1464600	UIS2	PF3D7_1345100	TRX2	PF3D7_1108700	Pfj2	
22	PF3D7_1419800	GR	PF3D7_0316300.2	PPase	PF3D7_1404900	None	PF3D7_1335100	MSP7	
23	PF3D7_0207800	SERA3	PF3D7_1419800	GR	PF3D7_1108700	Pfj2	PF3D7_1024800	EXP3	
24	PF3D7_0532100	ETRAMP5	PF3D7_1334800	MSRP2	PF3D7_0830400	PV/Exported Localization (ref 76)	PF3D7_1008900	AK1	
25	PF3D7_1420700	P113	PF3D7_1454400	APP	PF3D7_1016400	FIKK10.1	PF3D7_1232100	CPN60	
26	PF3D7_1014100	MSA180	PF3D7_0830400	PV/Exported Localization (ref 76)	PF3D7_1232100	CPN60	PF3D7_0706400	RPL37	
27	PF3D7_0207400	SERA7	PF3D7_0902800	SERA9	PF3D7_1008900	AK1	PF3D7_1104400	Trx-mero	
28	PF3D7_0702500	Exported	PF3D7_0207800	SERA3	PF3D7_1335100	MSP7	PF3D7_1016400	FIKK10.1	
29	PF3D7_0501200	PIESP2	PF3D7_0930300	MSP1	PF3D7_0625400	None	PF3D7_0501200	PIESP2	
30	PF3D7_1419800	GR	PF3D7_0207400	SERA7	PF3D7_1001200	ACBP2	PF3D7_1460700	RPL27	

Dorocito Stago	Mornhological Abnormality	Replicate -	% of Cells Displaying Abnormality		
Parasite Stage	Morphological Abnormality		Parent -aTc	EXP1 ^{apt} +aTc	EXP1 ^{apt} -aTc
	Increased seperation between PVM and PPM	1	0	1	8
	increased seperation between r vivi and r r w	2	2	0	11
Tranhazaitas	Increased seperation between PVM and PPM with abnormal	1	0	2	24
Trophozoites	membrane structures in the PV that contain RBC cytosol	2	0	0	7
	TVN-like membrane features in RBC cytosol	1	4	21	26
		2	11	20	25
	Increased seperation between PVM and PPM	1	0	0	4
	increased seperation between FVW and FFW	2	1	0	10
Segmented	Increased seperation between PVM and PPM with abnormal	1	1	3	15
Schizonts	membrane structures in the PV that contain RBC cytosol	2	1	1	15
	TV/N like membrane features in BBC external	1	2	12	24
	TVN-like membrane features in RBC cytosol	2	8	8	21