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21	Abstract

complex interplay between cell cycle and life cycle progression. Evidence from previous studies indicated that Cdc2 related kinase 3 (CRK3) in complex with the cyclin CYC6 is a functional homologue of the major cell cycle regulator CDK1, yet definitive genetic evidence for an essential role in parasite proliferation is lacking. To address this, we have implemented an inducible gene deletion system based on a dimerised Cre recombinase (diCre) to target CRK3 and elucidate its role in the cell cycle of *L. mexicana*. Induction of diCre activity in promastigotes with rapamycin resulted in efficient deletion of floxed *CRK3*, resulting in G2/M growth arrest. Co-expression of a *CRK3* transgene during rapamycin-induced deletion of *CRK3* resulted in complementation of growth, whereas expression of an active site *CRK3*^{T178E} mutant did not, showing that protein kinase activity is crucial for CRK3 function. Inducible deletion of *CRK3* in stationary phase promastigotes resulted in attenuated growth in

mice, thereby confirming CRK3 as a useful therapeutic target and diCre as a valuable new

Leishmania mexicana has a large family of cyclin-dependent kinases (CDKs) that reflect the

Introduction

tool for analysing essential genes in *Leishmania*.

The leishmaniases, diseases caused by protozoan parasites of the genus *Leishmania*, have diverse clinical manifestations dependent on the species and host immune response. Leishmaniasis is a substantial public health issue, causing an estimated 40,000 deaths annually and approximately 0.2 to 0.4 and 0.7 to 1.2 million visceral and cutaneous manifestations of the disease respectively (Alvar *et al.*, 2012). Existing drug therapies are problematic due to high treatment costs, toxicity and undesirable administration routes, making the development of novel and effective drug therapies to expand the current repertoire crucial. Phenotypic strategies to identify drug targets in the mammalian infective amastigote life cycle stage are of particular importance for drug discovery programs.

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As unicellular organisms, Leishmania depend on stringent control of cellular division to propagate and maintain infection. Protein kinases elicit pronounced effects on the Leishmania cell cycle by regulation of cell signalling pathways, and a number of protein kinases have been identified that are essential for promastigote viability (Wang et al., 2005; Dacher et al., 2014). The cyclin-dependent kinases (CDK) are of particularly interest due to their pivotal roles as cell cycle regulators. The use of CDK inhibitors in cancer therapy (Cicenas and Valius, 2011; Knapp and Sundström, 2014) and the relative expansion of this protein family in Leishmania relative to other unicellular organisms distinguishes them as suitable drug targets. In particular, the CDK related kinase CRK3 has been demonstrated as being important for regulation of the L. mexicana promastigote cell cycle by existing genetic manipulation techniques and cell cycle arrest following treatment with CDK inhibitors (Grant et al. 1998; Hassan et al. 2001; Grant et al. 2004). Recombinant protein kinase activity assays (Gomes et al., 2010) and yeast recovery mutants (Wang et al., 1998) have provided further validation of CRK3 as a drug target, leading to the identification and synthesis of a number of CRK3 inhibitors (Grant et al., 2004; Cleghorn et al., 2011; Walker et al., 2011; Goyal et al., 2014; Řezníčková et al., 2015). Regulation of CRK3 expression in L. mexicana is desirable to further assess its function in both procyclic promastigote and amastigote life cycle stages, however, no system exists for conditional deletion of essential genes. Recent application of plasmid shuffle methodology has addressed this issue by enabling the generation of partial null mutants to further study essentiality and important residues within coding sequences (Morales et al., 2010; Dacher et al., 2014), however the gene is not deleted and this prevents phenotyping of a *null* mutant.

To address this limitation, we have implemented a rapamycin-inducible gene deletion system using a dimerised Cre recombinase (diCre) (Jullien *et al.*, 2003; Collins *et al.*, 2013; Andenmatten *et al.*, 2013) to target *CRK3* and elucidate its role in the cell cycle of *L. mexicana*. *L. mexicana* is generally diploid (Rogers *et al.*, 2011) and both *CRK3* alleles were replaced with a 'floxed' *CRK3* open reading frame and the diCre coding sequence through promastigote transfection and homologous recombination. This system was used to conditionally delete *CRK3* during promastigote growth and so prove that CRK3 mediates the transition through G2/M. Induced loss of *CRK3* was complemented by expression of a *CRK3* transgene but not by expression of an inactive site (T178E) *CRK3* mutant, showing that protein kinase activity is crucial for CRK3 function. Significantly, conditional deletion of *CRK3* in stationary phase promastigotes and subsequent attenuation during murine infection demonstrates that CRK3 activity is essential for establishing infection. This system represents a new method to directly assess whether a gene is essential to parasite viability and provides novel insight into the function of essential genes in *Leishmania*.

Results

DiCre activity is tightly regulated in L. mexicana promastigotes and amastigotes

To test the activity of diCre in L. mexicana promastigotes, a reporter cell line was generated by integration of a loxP-flanked GFP into the ribosomal locus: [SSU GFP^{Flox}]. This cell line was transfected with a diCre construct containing the two dimerizable Cre recombinase subunits with the homologous flanks of crk3 to generate the heterozygous line $(\Delta crk3::DICRE/CRK3 [SSU GFP^{Flox}])$. Integration of the diCre construct at the CRK3 locus was confirmed by PCR analysis (Fig. S1A). No effect on the growth of SSU GFP^{Flox} or $\Delta crk3::DICRE/CRK3$ [SSU GFP^{Flox}] was observed in the presence of the dimerization ligand, rapamycin, up to the highest dose of 250 nM (Fig. S1B). GFP excision following incubation with increasing concentrations of rapamycin was investigated by PCR using specific primers flanking GFP. A single 1.45 kb PCR product, the floxed GFP fragment, was detected in the absence of rapamycin, whilst a 0.69 kb PCR product, representing the excised locus, was detected following rapamycin treatment only (Fig. 1A), indicating tight regulation of diCre activity. $\Delta crk3::DICRE/CRK3$ [SSU GFP^{Flox}] and [SSU GFP^{Flox}] promastigotes grown for 5 days in the presence or absence of increasing concentrations of rapamycin were analysed by flow cytometry to measure levels of GFP expression (Fig. 1B). Treatment of $\Delta crk3$::DICRE/CRK3 [SSU GFP^{Flox}] promastigotes with greater than 5 nM rapamycin resulted in substantial loss of GFP expression compared with the untreated controls, whilst GFP expression in [SSU GFP^{Flox}] was the same following growth in all concentrations of rapamycin. GFP loss in $\Delta crk3$::DICRE/CRK3 [SSU GFP^{Flox}] promastigotes grown in the presence or absence of 100 nM rapamycin for 5 days was further assessed by Western blotting of total protein extracts using anti-GFP antibody (Fig. 1C). Rapamycin treated promastigotes had considerably reduced GFP compared with the untreated controls, thereby demonstrating that gene loss results in reduced protein expression. These data also demonstrate that expression of diCre from the CRK3 locus is sufficient to efficiently excise the GFP transgene at rapamycin concentrations above 5 nM, and that no background diCre activity can be detected in the absence of ligand. 100 nM rapamycin was chosen as the optimum concentration to induce diCre activity in promastigotes whilst having no effect on in vitro cell growth.

To test diCre functionality in amastigotes, infectious promastigotes of the experimental line Δ*crk3::DICRE/CRK3* [*SSU GFP*^{Flox}] were inoculated into BALB/c footpads and amastigotes purified from the resulting lesion. *Ex vivo* amastigotes retained high levels of green fluorescence and were incubated with rapamycin for 24 hrs in Schneider's medium prior to infection of bone-marrow derived macrophages. Efficient excision of *GFP*^{Flox} was detected by PCR amplification of a 0.69 kb fragment representative of *GFP* loss in all rapamycin treated samples (Fig. 1D) and GFP⁻ (non-fluorescent) amastigotes were observed by comparing images obtained through fluorescence live cell imaging (Fig. S1C). Residual GFP⁺ amastigotes were still visible by microscopy (Fig. S1C) and could be detected by flow cytometry (Fig. S1D); this was possibly due to the slow replication rate of amastigotes leading to a low rate of GFP turnover. These data demonstrate inducible diCre activity in amastigotes.

Inducible deletion of CRK3 in L. mexicana promastigotes

The functional and efficient levels of diCre-mediated excision of *GFP* underpinned the development of a system for conditional deletion of essential genes. Gateway recombineering was used to flank appropriate diCre and loxP expression constructs with gene-specific, homologous flanks (Fig. S2). Plasmids were generated by this method to replace the two alleles of *CRK3*, an essential gene in *L. mexicana* (Hassan *et al.*, 2001) (Fig S3A). The first allele of *CRK3* was replaced with *DICRE* (Δ*crk3*::*DICRE/CRK3*) and the second allele of

CRK3 was subsequently replaced with a floxed C-terminal GFP-tagged CRK3 version ($\Delta crk3$:: $DICRE/\Delta crk3$:: $CRK3^{Flox}$; Figs. 2A and S3B). In addition, an mCherry red fluorescent protein coding sequence was incorporated downstream from the floxed CRK3-GFP to facilitate flow cytometry and microscopy analysis. Transfection resulted in multiple clones with the expected genetic modifications, as confirmed by PCR analysis (Fig. S3B).

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The growth of promastigotes from two $\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ clones were assessed following diCre-mediated excision induced with 100 nM rapamycin (Fig. 2B). Cells were counted over the course of 5 days, revealing a pronounced growth defect and reduction in cell number in rapamycin treated cells compared with uninduced controls. PCR analysis of promastigates grown in the presence or absence of 100 nM rapamycin for 24 h and 48 h confirmed efficient loss of the CRK3 gene (Fig. 2C) by the amplification of a single 1.36 kb DNA fragment for both rapamycin treated clones. The retention of the 3.4 kb amplicon containing the CRK3 gene in both untreated clones is evidence that no background diCre activity can be detected in the absence of rapamycin. To test for loss of the CRK3-GFP protein, total protein extracts of clone 2 promastigotes grown for 96 h in the presence or absence of 100 nM rapamycin were analysed by Western blot analysis with anti-GFP antibody (Fig. 2D) Very low levels of protein were detected in the treated promastigotes compared to the untreated cells, confirming that the conditional gene loss leads to reduced protein levels. Treatment with 100 nM rapamycin did not result in any noticeable effect on L. mexicana promastigote growth (Fig. S1B), however the pronounced growth arrest arising from loss of the essential gene could possibly result in cellular stress that synergises with rapamycin. These data show that this is a viable genetic manipulation strategy and that loss of CRK3 resulted in growth arrest and reduced cell numbers, both phenotypes consistent with loss of an essential gene.

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Cell cycle analysis of CRK3-deficient promastigotes.

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Previous attempts to impair CRK3 function in *Leishmania* by treatment with protein kinase inhibitors may have resulted in off-target effects (Grant *et al.*, 2004; Reichwald *et al.*, 2008; Cleghorn *et al.*, 2011; Jorda *et al.*, 2011; Efstathiou *et al.*, 2014; Řezníčková *et al.*, 2015). Here the utilisation of diCre mediated gene deletion enabled the effect of CRK3 depletion on the cell cycle to be investigated. Firstly, microscopic analysis of the cells at 96 h post-induction showed an accumulation of large, aberrant cells with altered organelle homeostasis

as evidenced by the presence of cells with multiple flagella (Fig. 3A). DAPI labelling of such multi-flagellated cells to visualise cellular DNA revealed the presence of enlarged nuclei indicative of an arrest in mitosis. Interestingly, cells were also observed that lacked a nucleus but retained the kinetoplast ('zoids'), a cell cycle defect observed previously by the treatment of promastigotes with CDK inhibitors (Grant et al., 2004). Secondly, flow cytometry was performed to determine the overall DNA content of Δcrk3::DICRE/Δcrk3::CRK3^{Flox} promastigotes grown in the presence or absence of 100 nM rapamycin for 72 and 96 h (Fig. 3B). This analysis showed that conditional deletion of CRK3 resulted in the accumulation of cells with 4C DNA content, associated with cell cycle arrest at G2/M, whilst an increasing population of cells with DNA content <1C indicates the accumulation of zoids. Finally, to assess the rate of cell death occurring in CRK3-deficient cells a viability assay was performed on promastigotes after growth in the presence or absence of 100 nM rapamycin for 72 h (Figs. 3C & S4). After 72 h the proportion of propidium iodide positive cells (PI⁺) was around 40% indicating a high level of cell death, which likely resulted from the accumulation of anucleated zoids at this time point. Flow cytometry analysis of cell size (using forward scatter) was in agreement with the microscopy analysis and showed that CRK3 deficient cells were substantially larger than cells retaining the gene (Fig. S4). Taken together, these data provide evidence that CRK3 plays an essential role in regulating mitosis in replicating promastigotes.

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Active CRK3 is required for cell cycle progression in promastigotes

192 We demonstrated that diCre could be used to efficiently delete a floxed copy of CRK3, so we 193 exploited the efficiency of this system to further study gene function through 194 complementation. Such a system was established by expressing a histidine-tagged CRK3 (CRK3his) (Hassan et al., 2001) transgene in $\Delta crk3$:DICRE/ $\Delta crk3$::CRK3^{Flox} promastigotes. 195 196 No significant difference in growth was noted in the presence or absence of rapamycin over a 197 5 day period (Fig. 4A). Efficient excision of floxed CRK3 in the induced culture was 198 confirmed by PCR amplification of the diagnostic 1.36 kb fragment by 24 h post-treatment 199 with 100 nM rapamycin (Fig. 4B). The proliferation of promastigotes, despite loss of floxed CRK3, indicates CRK3 transgene complementation in the induced $\Delta crk3$ cell line. Previous 200 studies have shown that recombinant L. mexicana CRK3^{T178E} protein lacks H1 kinase activity 201 (Gomes et al., 2010) and an L. major CRK3^{T178E} mutant fails to complement a cdc2-33(ts) 202 203 yeast mutant (Wang et al., 1998). To test whether active CRK3 is required for cell growth,

we exploited this complementation approach by generation of the cell line Δcrk3::DICRE/Δcrk3::CRK3^{Flox} [SSU CRK3^{T178E}] expressing a T-loop residue mutated version of CRK3 from the ribosomal locus. Growth curves indicate that expression of the CRK3^{T178E} transgene failed to complement the loss of CRK3^{Flox} following induction with rapamycin (Fig. 4A, B) thereby demonstrating that CRK3^{T178E} cannot rescue loss of active CRK3. The overall growth rate of both complementation mutants was reduced relative to the parental line (Table 1) and may explain the growth arrest at 72 h following excision of CRK3 in $\triangle crk3::DICRE/\triangle crk3::CRK3^{Flox}$ [SSU CRK3^{T178E}] compared with a more rapid onset of growth arrest in the parental line (Fig. 2B). These data show that active CRK3 is required for parasite growth. The CRK3 deficient cells were analysed by flow cytometry and fluorescence microscopy showing that $\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ [SSU CRK3^{T178E}] cells were blocked in G2/M (Fig. 4C) and were multi-nucleate and aberrant (Fig. 4D). These data are in agreement with the phenotype observed following excision of CRK3 in wild-type cells (Fig. 3A & B), thereby indicating the importance of the T-loop in regulating CRK3 activity. Based on these results, we conclude that transgene complementation can be used to confirm the specificity of conditional deletion of essential genes and also to probe the function of genes following mutagenesis.

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CRK3 is essential for in vivo infection of murine hosts

The lack of a conditional system to regulate expression of essential genes is a major obstacle for *in vivo* studies of essentiality, with such studies having crucial applications for drug target validation. To address this we tested if CRK3 activity is essential for survival of the parasite over the course of in vivo infection. Monitoring infection by detection of the light signal emitted from bioluminescent Leishmania using an in vivo imaging system (IVIS) is an established, longitudinal and non-invasive method to correlate signal with pathogen load (Lang et al., 2005; Lecoeur et al., 2007; Talmi-Frank et al., 2012; Vasquez et al., 2015). To assess the outcome of CRK3 loss on the proliferation of L. mexicana in vivo, bioluminescent lines were generated by transfection of L. mexicana wild-type and $\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ promastigotes with a ribosomal integration construct encoding red-shifted firefly luciferase, Ppy RE9H (Branchini et al., 2010; McLatchie et al., 2013). Both lines were bioluminescent as determined by luciferase expression assays on logarithmic stage promastigotes. The resulting $\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ [SSU RE9H] cell line produced 5 fold higher bioluminescence compared with the wild-type [SSU RE9H]

control (Fig. S4). Footpad bioluminescence detected with an *in vivo* imaging system (IVIS) correlated well with parasite burden in mice infected with L. mexicana expressing Ppy RE9H (Fig. 5A; y = 4.8 + 0.43x, $R^2 = 0.743$ and p < 0.0001). The slope of the linear regression line (0.43) revealed smaller increases in bioluminescence with increasing parasite burden. This may be related to tissue absorbance of light in vivo or limited substrate availability with increasing numbers of amastigotes within the lesion. Nevertheless, these data show that parasite burdens can be predicted from bioluminescence and that IVIS could be used for the non-invasive monitoring of parasite growth in mice over 10 weeks of infection. Following treatment of $\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ stationary phase promastigotes with rapamycin for 24 h the amplification of a 1.36 kb fragment (Fig. 5B) indicated that the majority of parasites had successfully excised floxed CRK3. The presence of small amounts of a 3.4 kb amplicon corresponding to the intact floxed CRK3 gene, however, also suggested that some parasites had retained the gene. These stationary phase $\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ [SSU RE9H] promastigotes either rapamycin treated (+ Rap) or not treated (- Rap) were then inoculated into the footpads of BALB/c mice. The in vivo bioluminescence in footpads of mice infected with the rapamycin-treated $\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ [SSU RE9H] was significantly reduced compared to the uninduced control by 5 weeks post-infection (p <0.001) and this continued up to 9 weeks post-infection (p<0.005) (Fig. 5C, D). From 5 to 9 weeks the bioluminescence from footpads infected with rapamycin-treated parasites increased 100-fold and was likely due to the proliferation of parasites that had not responded to rapamycin treatment and persisted in the lesion. To investigate this possibility, viable amastigotes were purified from the lesions of four mice at 10 weeks post-infection and analysed for the presence of CRK3^{Flox} by PCR after a single round of in vitro culture (Fig. 5E). A 3.4 kb PCR product containing CRK3 was amplified from all samples, indicating the persistence of parasites that had escaped diCRE mediated excision of CRK3.

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The ability of CRK3 deficient promastigotes to establish infection was further assessed by measuring footpad sizes at weekly intervals (Fig. 5F). The footpad sizes of mice infected with either untreated or rapamycin-treated $\Delta crk3$:: $DICRE/\Delta crk3$:: $CRK3^{Flox}$ [SSU RE9H] parasites were similarly low until about 4 weeks post-infection. Subsequently, footpads containing untreated parasites increased steadily over the course of infection, whilst those infected with rapamycin-treated $\Delta crk3$:: $DICRE/\Delta crk3$:: $CRK3^{Flox}$ [SSU RE9H] remained low until 9 weeks post infection. Comparison of the bioluminescence and lesion sizes suggest that there is a delay in lesion development despite parasite proliferation and that the lesions only increase

significantly when parasite load reaches a certain level (equating to bioluminescence $\approx 10^7$ photons/sec); in the case of the untreated parasites this occurred from about 5 weeks while for rapamycin-treated parasites this level of parasite burden had still not been reached by 9 weeks. Altogether these data show that loss of active CRK3 impairs the establishment of infection *in vivo*, and that a later resurgence of parasites likely results from a small population of cells which previously escaped *CRK3* conditional deletion.

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Discussion

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We have developed an inducible system for the genetic manipulation of essential genes in Leishmania. Inducible diCre was used to demonstrate the requirement for CRK3 activity in the regulation of mitosis. A distinct growth defect was observed 48 h after induced deletion of CRK3 (Fig. 2) resulting in cells arrested in G2/M, as well as an accumulation of zoids and eventually a population of enlarged, multi-flagellated cells (Fig. 3). This phenotype was rescued by expression of a CRK3 transgene from the ribosomal locus, confirming that loss of CRK3 caused mitotic arrest (Fig. 4). Arrest in G2/M and the accumulation of zoids have previously been reported following incubation of L. mexicana promastigotes with the CRK3 inhibitors flavopiridol (Hassan et al., 2001) and indirubin (Grant et al. 2004), showing correlation between genetic and chemical downregulation of CRK3 activity. In Trypanosoma brucei RNAi knockdown of the syntenic orthologue of CRK3 in the procyclic form also results in G2/M arrest and zoid formation (Tu and Wang, 2004), with the accumulation of such aberrant cells explained by the lack of a checkpoint controlling exit from mitosis and entry in cytokinesis (Ploubidou et al., 1999; Hammarton et al., 2003). Inducible deletion of CRK3 indicates that this checkpoint is also absent in L. mexicana promastigotes, resulting in impairment of mitotic progression, followed by re-initiation of G1 in the absence of cytokinesis. It appears that these abnormal cells can eventually undergo cytokinesis; however the daughter cell lacks a nucleus and is often multi-flagellated (see bi-flagellated zoid in Fig 3A), whilst the high levels of cell death occurring 72 h after gene loss show that such progeny are not viable.

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CRK3 is active at different stages in the cell cycle by forming complexes with cyclin partners such as CYC6 and CYCA, therefore CRK3 deletion could impact the cell cycle at multiple stages. RNAi of the CYC6 in *T. brucei* procyclic forms results in growth arrest within 48 h of induction and the accumulation of zoids and cells in G2/M (Hammarton *et al.*, 2003). A

similar phenotype was found in this study with CRK3 inducible deletion, suggesting that the CRK3:CYC6 complex is involved in regulation of mitosis (Walker et al., 2011). Less is known about the activity of CRK3:CYCA. Protein expression assays of L. donovani CYC1 (the functional orthologue of CYCA) demonstrates an increased abundance during S-phase (Banerjee et al., 2006) coupled with histone phosphorylation by an active CRK3:CYC1 complex (Maity et al., 2011), which is suggestive of S-phase kinase activity. Active, recombinant L. mexicana CRK3:CYCA has also been engineered, with phosphorylation of the T-loop residue T178 by the CDK activating kinase (CAK) Civ-1 increasing activity (Gomes et al., 2010). The T178 residue is essential for CRK3 activity as T178E mutagenesis inhibits functional rescue in S. pombe (Wang 1998) and ablates kinase activity in recombinant CRK3^{T178E}:CYCA (Gomes et al., 2010). The necessity of T178 was tested directly in this study, with excision of floxed CRK3 in the $\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ [SSU CRK3^{T178E}] line leading to cell cycle arrest in G2/M and zoid formation. The growth rate of this line and $\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ [SSU CRK3] were reduced when compared to Δcrk3::DICRE/Δcrk3::CRK3^{Flox} (Table 1), indicative of generally reduced growth rate when expressing a transgene. Episomal complementation with CRK3 did not result in an observable growth defect (Hassan et al., 2001), but this may result from the modulation of the number of episomal copies, as has been observed previously following complementation of the essential MCA gene (Ambit et al., 2008). Integration into the 18s rRNA locus results in consistently high levels of expression (Misslitz et al., 2000) leading to non-physiological levels of CRK3 and subsequent CRK3:CYC6 activity at potentially inappropriate stages of the life cycle.

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The reduced growth rate of promastigotes overexpressing CRK3^{T178E} is likely due to a partial dominant negative effect, whereby inactive CRK3^{T178E} binds endogenous CYC6 leading to impaired protein kinase activity even in the presence of active CRK3. This reduced growth rate may explain both the cell cycle arrest at 72 h in the [SSU CRK3^{T178E}] complemented line (Fig. 4A) compared to arrest at 48 hours in $\Delta crk3$::DICRE/ $\Delta crk3$::CRK3^{Flox} (Fig. 2B) and additionally the lower proportion of zoids when analysed by flow cytometry (Fig. 4C). The accumulation in G2/M suggests that mutation ablates CRK3:CYC6 activity, rather than CRK3:CYCA, where an increase of cells in G1/S might be anticipated. Both induced and uninduced $\Delta crk3$::DICRE/ $\Delta crk3$::CRK3^{Flox} [SSU CRK3^{T178E}] have dramatically reduced flagellum length and are immotile (Fig. 4D).. The reduced size of the flagellum and a growth defect are similar phenotypes to those observed in cell lines deficient in ATG5, a key

component of the autophagic pathway (Williams *et al.*, 2012). This is likely a result of their impaired ability to salvage material through the autophagic pathway, imparting selection on the parasites to reduce energy through flagellum regression. The partial dominant negative effect of CRK3^{T178E} may also result in metabolic stress in these cells leading to the phenotype observed. The importance of T178 as an active site residue for regulating progression through G2/M implicates upstream modifiers of this residue as essential regulators of the *L. mexicana* cell cycle. In mammalian cells CDK7 acts as a CAK to regulate CDK1 by phosphorylation at this T-loop residue, yet no CAK homologues have been identified in the *Leishmania* genome (Gomes *et al.*, 2010). The identification of potential post-transcriptional modifiers of the CRK3 T-loop residue that act in an analogous fashion to CDK7 would therefore yield promising targets for drug discovery. The phenotype of the induced cell line shows the importance of the T-loop residue for CRK3 activity and mitotic function within the cell, endorsing this complementation assay as a rational approach for active site investigation.

The assessment of gene essentiality for amastigote viability is an important approach in the context of drug target validation as this life cycle stage is the pathologically significant form. The recent utilisation of plasmid shuffle has facilitated the study of Leishmania genes involved in life cycle differentiation and essentiality both in amastigote and promastigote forms by the generation of partial *null* mutants (Morales et al., 2010; Dacher et al., 2014). Retention of an episomal gene in a null mutant cell line after murine infection is a useful approach to assess that gene as necessary to amastigote survival in vivo (Wiese, 1998). Despite such elegant utilisation of reverse genetic methods to probe gene function, no method exists for the generation of conditional null mutants during in vivo infection. Our study does not address this lack directly due to the sensitivity of amastigotes to rapamycin, however as diCre activity remains high in stationary-phase promastigotes CRK3 was efficiently excised (Fig. 5B) to probe the subsequent infectivity of CRK3-deficient promastigotes. By tracking the progression of infection with reporter parasites expressing the highly sensitive red-shifted luciferase (Branchini et al., 2010; McLatchie et al., 2013) and by footpad size measurement, we demonstrate that the CRK3-deficient L. mexicana are unable to proliferate in their mammalian host (Figs. 5C, 5D and 5F). Importantly, the wild-type line expressing luciferase grows normally in mice following rapamycin treatment, which indicates that lack of growth of the CRK3-deficient mutant is not a result of the drug treatment. The average light intensities emitted from footpads infected with the wild-type [SSU RE9H) line and those from footpads infected with the $\triangle crk3::DICRE/\triangle crk3::CRK3^{Flox}$ [SSU RE9H] line retaining floxed *CRK3* are at similar levels throughout infection, yet mean footpad size is larger in wild-type [SSU RE9H] infected mice after 3 weeks post infection; such disagreement may be a result of the 5 fold lower signal intensity of the wild-type [SSU RE9H] compared with the experimental line (Fig. S5) and therefore an overall higher burden of the wild-type line is likely masked by a reduced bioluminescent signal intensity.

Interestingly, parasite burden as measured by total flux remains consistently above the background intensity (dashed line, Fig. 5D) in those footpads infected with the *CRK3*-deficient line, suggestive of the survival of a low number of bioluminescent parasites. The outgrowth of these parasites was observed through an increased bioluminescence signal at 9 weeks post infection compared with 5 weeks (Fig. 5C and 5D). Purification and PCR analysis of these parasites shows they retained the floxed *CRK3* (Fig. 5E) and that the persistence of signal and subsequent increase are a result of incomplete excision of floxed *CRK3* during the 24 h incubation with rapamycin. These data further demonstrate the essentiality of *CRK3* activity for establishing infection.

This is the first time an essential gene in promastigotes has been studied in vivo by conditional deletion, representing a useful tool to probe gene function. We are validating the feasibility of conditional gene deletion ex vivo and in vivo using rapamycin and non-immunoinhibitory rapamycin analogues ('rapalogs'), with such work being useful for the future of drug target validation. DiCre activity has been demonstrated in vivo (Jullien et al., 2007), however rapamycin treatment may be a limitation due to influence on the host immune response and on amastigote proliferation. Our attempts to study the effect of CRK3^{Flox} deletion in lesion-derived amastigotes grown in axenic culture medium was problematic due to reduced proliferation of both experimental and wild-type L. mexicana at the relatively low dose of 50 nM rapamycin, therefore the use of rapalogs would be a rational approach for induction of diCre activity if they have reduced binding affinity for Leishmania TORs (Madeira da Silva and Beverley, 2010). A second generation diCre is currently in development and may present an alternative method for inducible gene deletion in vivo. In diCre2, each subunit is fused to mutant FKBP domains that are dimerised by the rapalog AP20187, which is amenable to in vivo use (Collins et al., 2013). Such a system could be applied for use in Leishmania and would complement our existing floxed gene replacement approach.

407 In conclusion we have developed a highly efficient inducible gene deletion system that when used with transgene complementation allows for the first time the function of essential 408 409 Leishmania genes to be elucidated. We have applied this approach to show that CRK3 is 410 required for promastigote progression through mitosis, with gene deletion mutants showing a 411 G2/M arrest and an accumulation of zoids, indicative of a lack of a cell cycle checkpoint in 412 cytokinesis. Inducible deletion of CRK3 in stationary phase promastigotes attenuates 413 infection in a murine host, providing further genetic validation of CRK3 as a potential drug 414 target (Grant et al. 1998; Hassan et al. 2001; Grant et al. 2004; Gomes et al. 2010; Walker et 415 al. 2011). Our diCre method provides a powerful tool for analysing genes essential for 416 promastigote proliferation and to the study of the differentiation of promastigotes to 417 amastigotes.

418

419

Figure legends

420

- 421 **Fig. 1.** Validation of inducible diCre in *L. mexicana*: conditional deletion of *GFP* in
- 422 promastigotes and amastigotes.
- 423 A. Gene excision analysed by PCR amplification. Schematic (lower) shows the SSU GFP^{Flox}
- locus and the recombination event expected after treatment with rapamycin (Rap). (upper)
- 425 PCR amplification with oligonucleotides 4287 and 4288 from experimental
- 426 $(\Delta crk3::DICRE/CRK3 [SSU GFP^{Flox}])$ and control [SSU GFP^{Flox}] promastigates at 5 days
- post-treatment with different concentrations of rapamycin.
- 428 B. Flow cytometry assessment of GFP intensity of experimental and control promastigotes
- incubated in the presence or absence of rapamycin for 5 days.
- 430 C. Western blotting analysis with anti-GFP and anti-EF1α loading control antibodies of
- protein extracted from experimental promastigotes grown for 5 days in the presence or
- absence of 100 nM rapamycin.
- D. PCR analysis of GFP^{Flox} loss (as described in A) in amastigates after 24 h rapamycin
- treatment (0 1000 nM), followed by 120 h infection in bone-marrow derived macrophages.
- 435 Lane 2 contains a 1 kb+ DNA ladder.

- 437 **Fig. 2.** Generation of a *CRK3* conditional deletion cell line.
- 438 A. Schematic showing the replacement of endogenous *CRK3* to generate
- 439 $\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$. One allele contains a *loxP* flanked *CRK3-GFP* coding

- sequence with mCherry red-fluorescent protein cassette (RFP) and puromycin drug selectable
- marker (*PAC*). The other allele contains genes encoding both diCre subunits (*CRE59*,
- 442 *CRE60*) each linked with rapamycin binding domains (not shown: *FKBP12 and FRB*
- respectively) and a blasticidin resistance cassette (BSD). Each construct was flanked with 500
- bp arms of homology (light grey) by Gateway recombination to facilitate integration at the
- 445 CRK3 locus. All coding sequences are flanked by regulatory elements (dark grey). L.
- 446 mexicana parasites were transfected sequentially with the diCre construct and floxed CRK3
- to confer resistance to blasticidin and puromycin antibiotics respectively.
- B. Clones 2 and 8 promastigotes were seeded at a density of 5 x 10⁵ cells ml⁻¹ and grown in
- the presence or absence (+/-) of 100 nM rapamycin for 5 days. Cell density was determined
- 450 by counting at 24 h intervals and mean \pm SD of triplicate values was plotted.
- 451 C. (lower) A schematic representation of the floxed CRK3 locus after excision. PCR
- amplification shows the primers binding upstream of the 5' CRK3 homologous flank and
- within the *PAC* cassette. (upper) PCR amplification of clones 2 and 8 at 24 h and 48 h +/-
- 454 100 nM rapamycin treatment was conducted and the resulting amplicons resolved on an
- agarose gel.
- D. Western blotting analysis with anti-GFP and anti-EF1α loading control antibodies of
- protein extracted from experimental clone 2 promastigotes grown for 4 days in the presence
- 458 or absence of 100 nM rapamycin.
- 459
- 460 **Fig. 3.** Analysis of *CRK3* deficient promastigotes.
- A. Representative images of cells grown in the absence (top) or presence (bottom two rows)
- of 100 nM rapamycin for 96 h. Promastigotes (clone 2) were stained with DAPI to observe
- and kinetoplast content alongside mCherry expression by fluorescence microscopy.
- 464 Scale bar represents 5μm.
- B. (upper) DNA content analysis of clone 2 promastigotes at 72 and 96 h post treatment.
- 466 Cells were fixed with methanol and stained with propidium iodide for flow cytometry
- analysis of 100,000 cells to examine nuclear content. Arrows indicate the positions of cells in
- 468 G₁ phase (2C), in G₂/M (4C) and low DNA content associated with increased incidence of
- 469 <1C zoids. (lower) Graphical representation of the DNA content of each population based on
- 470 the flow cytometry plots.
- 471 C. The viability of cells grown in the absence (-) or presence (+) of 100 nM rapamycin for 72
- h. Promastigotes (clone 2) were incubated with 5 µg ml⁻¹ propidium iodide (PI) for 15 min

- and analysed by flow cytometry. A heat lysed (HL) control in which half the sample was
- lysed by incubation at 70°C for 3 min was included to enable an appropriate live / dead gate
- 475 to be drawn. Numbers represent the percentage of cells assessed as PI positive (PI+) based on
- 476 the HL control. Data shown are the means of 3 technical replicates, data are representative of
- 477 2 independent experiments.

478

- 479 **Fig. 4.** *CRK3* wild type and active site mutant complementation assays.
- 480 A. Wild type complemented ($\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ [SSU CRK3], left graph) and
- mutant complemented ($\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ [SSU CRK3^{T178E}], right graph) cell
- lines were seeded as promastigotes at 1 x 10⁵ cells ml⁻¹ and grown +/- 100 nM rapamycin for
- 483 5 days. Cell density was determined by counting at 24 h intervals and the mean ± SD of
- 484 triplicate values was plotted.
- 485 B. The resulting amplicons generated by PCR amplification of each cell line at 24 and 48 h
- 486 after growth +/- 100 nM rapamycin.
- 487 C. (left) DNA content analysis of Δcrk3::DICRE/Δcrk3::CRK3^{Flox} [SSU CRK3^{T178E}]
- promastigotes after methanol fixation and staining with propidium iodide for flow cytometry
- analysis (100,000 cells) to examine nuclear content. Arrows indicate the positions of cells in
- 490 G₁ phase (2C), in G₂ (4C) and low DNA content associated with increased incidence of <2C
- zoids. (right) Graphical representation of the DNA content of each population based on the
- 492 flow cytometry analysis.
- 493 D. Representative images of $\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ [SSU CRK3^{T178E}] promastigates
- grown in the absence (top) or presence (bottom two rows) of 100 nM rapamycin for 96 h.
- 495 Parasites were stained with DAPI to detect nuclear and kinetoplast DNA by fluorescence
- 496 microscopy. Scale bar represents 5μm.

- 498 **Fig. 5.** *CRK3* conditional deletion in stationary phase promastigotes and *in vivo* infection.
- 499 A. Correlation between in vivo bioluminescence (total flux in photons per second) and
- parasite burdens from the same infected footpads. BALB/c mice were infected with L.
- 501 mexicana WT or Ppy RE9H-expressing stationary phase promastigotes and imaged weekly
- using an *in vivo* imaging system (IVIS). At 2, 4, 6 and 8 weeks post-infection mice were
- sacrificed after imaging and parasite burdens in infected footpads determined using limiting
- dilution assays. Each point shows the total flux and parasite burden from the footpad in one

- mouse (n = 3-4 mice per time point). Linear regression line and R^2 was calculated from the
- log transformed data.
- B. PCR amplification of the floxed CRK3 locus of $\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ [SSU
- 508 RE9H] stationary phase promastigotes after incubation in the presence (+) or absence (-) of 1
- 509 μM rapamycin for 24 h.
- 510 C. Control (-) or 24 h rapamycin-treated (+) stationary phase promastigotes were inoculated
- 511 into the footpads of BALB/c mice. The total flux (photons/sec) emitted from the infected
- 512 footpad region of interest (ROI) was quantified weekly.
- D. The total flux measured from infected footpads was plotted over 9 weeks of infection.
- Data shown represent the mean flux and SD from groups of four mice. The dotted line
- 515 indicates the average background flux emitted from uninfected footpads measured 1 week
- 516 post infection (n=12). A significant difference in the mean total flux emitted between the
- 517 footpads of mice infected with untreated and rapamycin-induced parasites was observed at 5
- and 9 weeks post infection (2-way ANOVA, ***P=<0.001;**P=<0.005).
- 519 E. PCR amplification of the floxed CRK3 locus of Δcrk3::DICRE/Δcrk3::CRK3^{Flox} [SSU
- 520 RE9H] + Rap after purification of amastigotes from the footpads of 10-week infected mice.
- 521 Cells were propagated *in vitro* to obtain sufficient genomic DNA for PCR analysis.
- F. Footpad sizes were recorded by weekly caliper measurement. Data shown represent the
- mean footpad size and SD from groups of four mice (Unpaired t-test *P=<0.05).

524525

- **Table 1.** Comparisons of the growth rates of conditional *CRK3* deletion lines measured
- 527 during logarithmic growth.

528

- **Experimental Procedures**
- 530 Ethics statement
- Animal studies were carried out under UK Home Office regulations (Project licence PPL
- 532 60/4442).
- 533 Parasite culture and transfection
- 534 Leishmania mexicana (MNYC/BZ/62/M379) promastigotes were cultured at 25°C
- in HOMEM supplemented with 10% heat inactivated foetal calf serum (HI-FCS) and 1%

- penicillin/streptomycin (PEN/STREP). Amastigotes were cultured in Schneider's Insect
- Medium supplemented with 20% HI-FCS, 1% PEN/STREP and 15µg mL⁻¹ Hemin at pH5.5.
- 538 Mid-log phase L. mexicana promastigotes were transfected with 10µg of digested DNA by
- 539 electroporation using the Nucleofector system with the Human T-Cell kit (Lonza) as
- described previously (Castanys-Muñoz et al., 2012). Transgenic cell lines were grown in the
- presence of appropriate antibiotics at the following concentrations: G418 50 μg mL⁻¹,
- 542 blasticidin 10 μg mL⁻¹ and puromycin 10 μg mL⁻¹ (InvivoGen).
- 543 Construct design and development
- A full list and descriptions of all primers (Table S1) and plasmids (Table S2) used in this
- study are available. To produce a diCre expression vector, the diCre coding sequences Cre59-
- 546 FKBP12 and Cre60-FRB were each flanked by actin and β-tubulin sequences in array with
- 547 blasticidin resistance cassette flanked by *DHFR-TS* regulatory elements. The sequence was
- 548 synthesised and sub-cloned into the pDONR221 vector (GenScript). The backbone of the
- loxP vector containing the loxP sites flanking a multiple cloning site and other restriction
- enzyme regions flanked by regulatory elements was synthesised (GenScript). The PAC,
- 551 mCherry and CRK3-GFP cassettes were inserted by enzymatic restriction digest mediated
- ligation, and subsequently sub-cloned into pDONR221. Addition of CRK3 homology
- flanking homology was performed by MultiSite Gateway 3-fragment vector construction
- 554 (Invitrogen) as per manufacturers' guidelines. Briefly, flanks were amplified by PCR by
- 555 Phusion polymerase (New England BioLabs) using oligonucleotides conferring attB
- recombination sites to the amplicons. Subsequent BP reactions inserted the flanks into
- appropriate pDONR vectors containing attL sites for site-specific recombination. An LR
- 558 reaction resulted in the flanking of diCre and loxP vectors into a pDEST vector for
- transfection. Finally, complementation plasmids were generated by inserting the CRK3,
- 560 CRK3^{T178E} and RE9H genes (Branchini et al., 2010; McLatchie et al., 2013) into a modified
- version of pGL631 (Misslitz et al., 2000) containing a G418r cassette for SSU integration
- construct by *Xho*I & *Not*I restriction enzyme digestion and ligation.
- 563 Induction of diCre mediated gene deletion
- All experiments were conducted using cells in the early to mid log stage of exponential
- growth (between 1-5 x 10⁶ cells mL⁻¹) with the exception of the stationary phase inducible

566 gene deletion. Between 1nM to 1µM rapamycin (Abcam) was administered by inoculation 567 into the cell culture medium from a 100 µM working stock. 568 Conditional gene deletion analysis Tag polymerase (NEB) was used to PCR amplify the regions surrounding GFP^{Flox} and 569 $\textit{CRK3}^{Flox}$ using primers shown in Table S1 and a T_A calculated using an online T_m calculator 570 (New England BioLabs) and 30 cycles for amplification. 571 572 Western Blot Analysis For western blotting analysis, either 1 x 10⁷ cells were loaded per lane or equal 573 concentrations of protein extract as quantified by Bradford assay of a 10% NuPAGE Bis-Tris 574 575 gel (Invitrogen) in MOPS running buffer and transferred onto Hybond-C nitrocellulose 576 membranes (GE Healthcare). Primary antibodies against GFP were used to detect GFP and 577 CRK3-GFP expression at 1:1000 whilst anti-EF1α was used as a loading control at 1:5000. 578 Membranes were washed three times in TBST, incubating for 10 min each time, before 579 incubation with horse radish peroxidase (HRP)-conjugated secondary rabbit and mouse 580 antibodies at 1:5000 dilution for 1 h at room temperature. After washing three times in TBST, 581 the membrane was treated with an ECL (enhanced chemiluminescence) kit (SuperSignal 582 West Pico Chemoluminescent Substrate, Pierce) according to manufacturer's instructions and 583 then exposed on Kodak photographic film. 584 Infection of mice 585 BALB/c mice were purchased from Charles River (MA., USA) and infected in the right footpad with 2×10^6 stationary-phase *L. mexicana* promastigotes in $1 \times PBS$. Lesion size was 586 monitored weekly and $\Delta crk3$::DICRE/CRK3 [SSU GFP^{Flox}] amastigates were purified before 587 588 the lesions reached a thickness of 5mm. 589 Purification of lesion derived amastigotes Lesion derived $\triangle crk3::DICRE/CRK3$ [SSU GFP^{Flox}] amastigotes were purified by 590 591 homogenising the extracted lesion in 1xPBS and passing the solution through a 20 µm cell 592 strainer. Amastigotes were pelleted by centrifugation at 2,000 g for 10 mins, followed by resuspension in culture medium. To prevent cells from clumping together and ensure accurate 593

cell counting, amastigote cultures were first centrifuged at 2,000 g for 10 mins and the

595 supernatant removed to leave the pellet in 500uL volume. The pellet was re-suspended in this 596 volume by gentle syringing through a blunt 16G needle and the single cell suspension added 597 back to the culture medium. Cell counting was performed by mixing the homogenised culture 598 1:1 with Trypan blue and cell counting with a Haemocytometer (Neubauer). 599 Macrophage differentiation and amastigote infection 600 Non-differentiated monocytes were extracted from the femurs and tibia of BALB/c mice by 601 dissection to remove the bones. RPMI 1640 medium was used to wash the bone marrow out 602 of the intact bones by syringing with a 25G needle. Extracted cells were quantified by dilution in Trypan blue (1:1) and counting with a haemocytometer. Monocytes were seeded at 603 5×10^5 cells ml⁻¹ in M Φ Medium (DMEM + L-Glut + 20%FCS + 1% P/S + 30% L-Cell M) in 604 8 ml volumes in Petri dishes and incubated at 37°C with 5% CO₂ for 3 days to induce 605 606 differentiation to monocyte-derived macrophage. After this period the medium was replaced 607 and by day 5 the cells were removed from the dishes using a cell scraper with ice-cold RPMI 1640. Bone marrow derived macrophage were adhered at a concentration of 5 x 10⁵ cells ml⁻¹ 608 609 overnight in DMEM medium with 10 % HIFCS at 37°C in 5% CO₂ onto 8-chamber tissue 610 culture slides (LAB-TEK) for microscopic analysis or 12 well plates for DNA extraction and 611 flow cytometry analysis. Macrophages were then infected at a ratio of 5 parasites per macrophage with lesion-derived $\Delta crk3::DICRE/CRK3$ [SSU GFP^{Flox}] amastigotes, which had 612 613 been previously grown in axenic medium in the presence or absence of rapamycin for 24 h. 614 Wells were washed at 24 h post infection to remove extracellular parasites and media 615 replenished with DMEM/10% HIFCS. Cells were removed from the plates for DNA 616 extraction and flow cytometry analysis by gentle scraping in ice cold RPMI at the 120 h end 617 time point. 618 Fluorescence microscopy analysis 619 For imaging, 2 x 10⁶ parasites were washed in 1 x PBS, re-suspended in Fluoromount-G (SouthernBiotech) DAPI infused mounting medium and mounted on glass slides for analysis. 620 621 Parasite morphology was observed by DIC and mCherry fluorescent imaging, and DNA 622 content observed by DAPI fluorescent imaging using a Delta Vision core (Image Solutions) 623 inverted microscope equipped with mCherry and DAPI filter sets. Images were processed 624 using Photoshop CS (Adobe) image software. GFP expression of intracellular amastigotes 625 was assessed by fluorescent microscopy. Cells were imaged between 24 and 120 h after 626 infection in the DeltaVision Core environmental chamber at 37°C and 5 % CO₂ upon 627 incubation in 1 x PBS infused with DAPI. 628 DNA content and GFP expression analysis by flow cytometry 629 Parasites were prepared for DNA content analysis as described previously (Paul Hassan et 630 al., 2001) with the exceptions that a MacsQuant flow cytometer was used to analyse 100,000 631 cells per sample. Cell distribution was modelled using FlowJo software (Tree Star). For 632 determining GFP expression of promastigotes and amastigotes by flow cytometry analysis, 633 live cells were washed twice in 1xPBS and passed through a nitex mesh prior to acquisition. 634 Viability assay Log-phase promastigotes were seeded at 5 x 10⁵ cells ml⁻¹ and grown in the presence or 635 absence of 100 nM rapamycin. At 72 h post treatment 1 x 10⁷ cells were washed once with 1 636 x PBS and incubated with 5ug ml⁻¹ propidium iodide (PI) for 15 minutes at room temperature 637 in the dark. A heat lysed (HL) control in which half the sample was lysed by incubation at 638 639 70°C for 3 min was included to enable an appropriate live / dead gate to be drawn. Cells were 640 washed with 1 x PBS and used to acquire 100,000 events per group by flow cytometry using 641 a MacsQuant flow cytometer. 642 In vivo *imaging* 643 For imaging, mice were anaesthetised with 4.0% isofluorane/1.5 L O₂ per minute and inoculated by subcutaneous injection with 200µl D-luciferin (15 mg ml⁻¹ in Mg/Ca-free 644 645 Dulbecco's modified PBS). Light emission was recorded 10 minutes after inoculation using 646 an IVIS Spectrum bioluminescence imaging system (PerkinElmer). Imaging was performed 647 with an open emission filter, for 30-60 second exposures, large binning, and 1 f/stop, and captured with a charge-coupled device (CCD) camera. The absolute unit of photon emission 648 was given as radiance (photons /second/cm²/steradian). Images were analysed using Living 649 650 Image Software (PerkinElmer) and regions of interest (ROI) of equal size were selected over 651 the infected footpads to quantify the amount of photon emission as total photon flux in 652 photons per second (photons/sec). 653 Statistical analysis

Statistical analysis was performed using GraphPad Prism 5. The analysis of significance of

- 655 the data was performed by 2-way ANOVA when comparing data from induced (+Rap) and
- uninduced (-Rap) $\Delta crk3::DICRE/\Delta crk3::CRK3^{Flox}$ [SSU RE9H] infections and by paired t-test
- when comparing footpad sizes.

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664 Conflict of Interest

The authors declare no conflict of interest.

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Figure S1

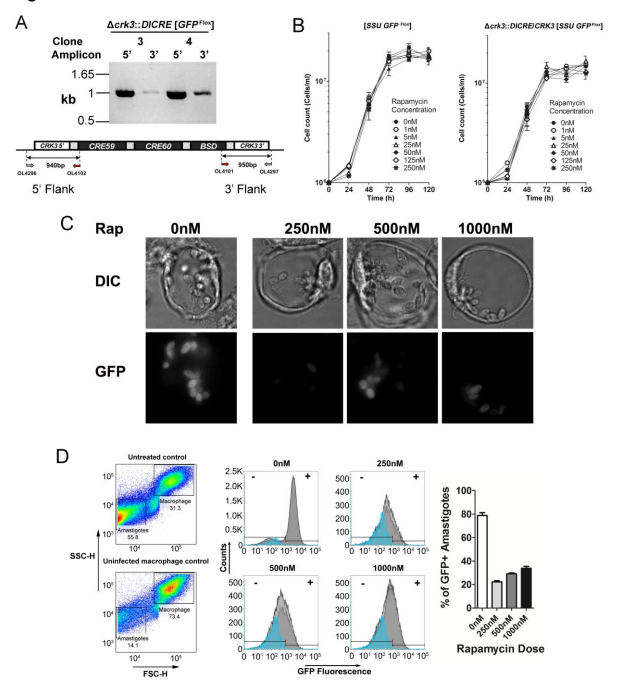


Fig. S1. A. Replacement of a single copy of CRK3 by diCre construct integration into the $[SSU\ GFP\ ^{Flox}]$ cell line was confirmed by PCR amplification of genomic DNA extracted from two clones (3 and 4). Oligonucleotides (OL) that bind outside the integration site (grey arrows) and within the diCre coding sequence (red arrows) were used to amplify 940 bp and 950 bp amplicons. Clone 3 was designated as the experimental line $\Delta crk3::DICRE/CRK3$ $[SSU\ GFP^{Flox}]$.

- B. Experimental $\Delta crk3::DICRE/CRK3$ [SSU GFP^{Flox}] or control [SSU GFP^{Flox}] L. mexicana promastigotes were seeded at $1x10^6$ cells ml⁻¹ and incubated in the presence or absence of between 1 to 250 nM rapamycin. Cell density was determined at 24 hour intervals by cell counting (N=1-3 technical replicates, error SEM).
- C. Representative DIC (upper) and GFP (lower) images from live cell imaging of amastigotes-infected macrophages at 5 days post-infection. GFP expression from live amastigotes was imaged using a Delta Vision core fluorescent microscope.
- D. GFP intensity loss in amastigotes extracted at day 5 post *in vitro* macrophage infection; (left) amastigotes were gated from large, granular macrophage by forward scatter (FSC) for size and side-scatter (SSC) for granularity. (middle) Histograms of amastigote GFP intensity were generated from amastigote gates with retention of GFP expression at >10³ fluorescence intensity based on rapamycin untreated controls. Blue plots represent the amastigote gate plotted from a macrophage only control group to represent background cellular 'debris' as a result of macrophage lysis following sample preparation (left). >20,000 amastigote events were analysed per treatment group based on two biological replicates shown as dark and light grey plots. (right) Retention of GFP signal as a % of amastigote gate displayed as bar graphs for each treatment group (Data represent means ± SEM).

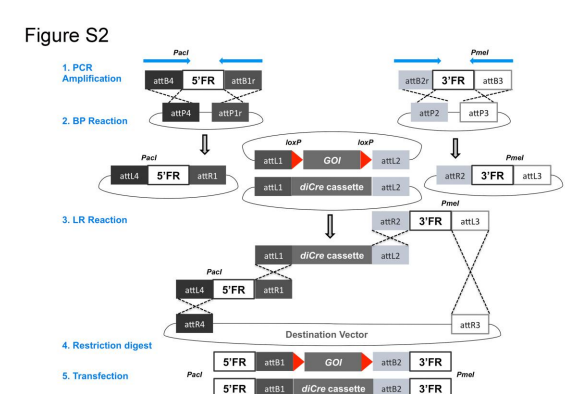


Fig. S2. Pipeline of Gateway-mediated addition of target gene homologous flanks to diCre and loxP vectors. (1) Primers (blue arrows) containing appropriate att sites and 5' PacI or 3'PmeI unique restriction sites amplify a 0.5-1 kb region up- and downstream of the gene. (2) BP clonase catalyses the insertion of these flanks into their appropriate vectors. (3) The resulting 5', 3' and diCre or loxP vectors are recombined into a pDEST vector by LR clonase. (4) The final vector is linearised by PacI and PmeI digest for (5) transfection into L. mexicana. This method enables flanking of both the floxed gene of interest (GOI) expression cassette and diCre expression cassette.

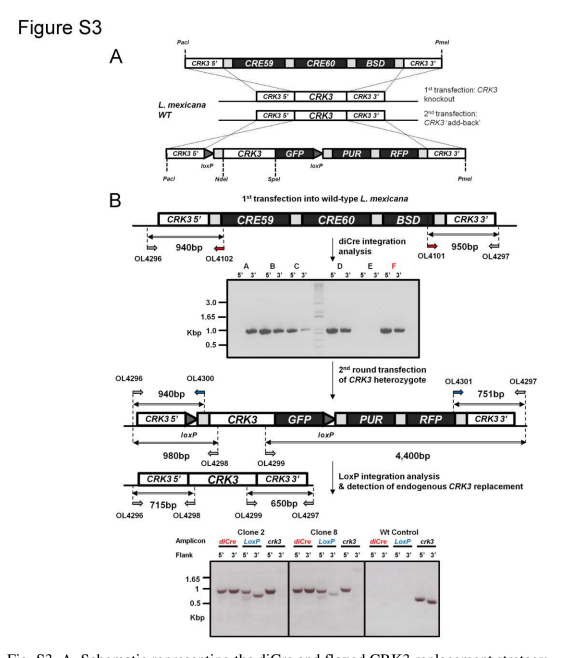


Fig. S3. A. Schematic representing the diCre and floxed CRK3 replacement strategy. Homologous recombination was facilitated by Gateway flanking of both diCre and loxP vectors with ~500 bp of *crk3* 5' and 3' homologous regions to replace both alleles. B. Transfection of wild-type *L. mexicana* with the diCre construct: integration was confirmed by PCR amplification of genomic DNA extracted from six clones with oligonucleotides (OL) binding outside the integration site (grey arrows) and within the diCre coding sequence (blue arrows) to amplify 940 and 950 bp amplicons. A single blasticidin (BSD) resistant clone F with *diCre* integrated at the *crk3* locus was subsequently transfected with the loxP construct to replace the remaining endogenous *crk3* allele with a floxed *CRK3* fused to a 3' *GFP* tag, thereby generating a diCre-mediated conditional deletion line: Δ*crk3*::*DICRE*/

 $\Delta crk3$:: $CRK3^{Flox}$. PCR amplification of genomic DNA extracted from two blasticidin/ puromycin (PUR) double resistant clones (2 and 8) with oligonucleotides binding outside the integration site (grey arrows), within the crk3 coding sequence (grey arrows), within the loxP vector (blue arrows) and diCre sequences (red arrows).

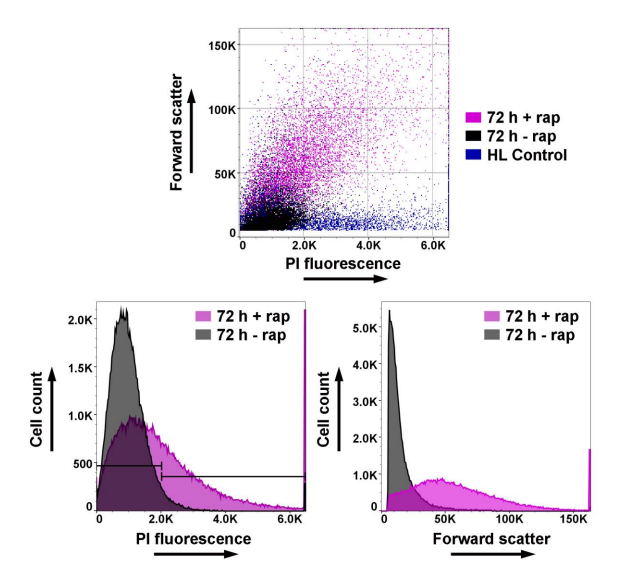


Figure S4. Viability assay of $\Delta crk3$:: $DICRE/\Delta crk3$:: $CRK3^{Flox}$ promastigotes. Cells were grown in the presence or absence of 100 nM rapamycin for 72 h. Live cells were incubated with 5ug ml⁻¹ propidium iodide (PI) for 15 minutes and uptake measured by flow cytometry alongside a heat lysed (HL) control in which half the cells were lysed by incubation at 70°C for 3 min prior to flow cytometry analysis. Top panel shows cell size as measured by forward scatter in the y-axis and cell lysis by increasing PI fluorescence along the x-axis. Bottom left panel shows the gating strategy whereby cells are defined as + or – in PI uptake based on the HL control. Bottom right panel is an analysis of promastigote cell size following incubation in the presence or absence of rapamycin. Results are representative of 2 independent experiments.

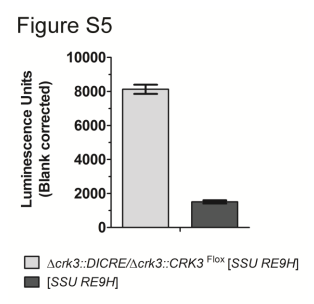


Figure S5. *In vitro* bioluminescence expression assay of experimental and control promastigotes. Promastigotes were assayed during logarithmic growth and luminescence expression data was acquired 30 minutes post luciferin treatment. Error bars represent the SEM of two technical replicates per clone.

Table S1

011	_	Table 31	T
Oligo No.		Description	Sequence
		Gateway cloning of CRK3 homolog	ous flanks
OL4249	F	Amplification of a 5' CRK3 GGGGACAACTTTGTATAGAAA	AAGTTGCCCTTAATTAAAAAGGTAGAGGATGCCGTTTT
OL4250	R	homologous flank with attB4/P1r GGGGACTGCTTTTTTGTACAA	ACTTGCTTGAAATGTTGCAGGGAGAAA
OL4251	F		AAGTGGGGAGTGGAAAAGGCATGACTGAA
OL4252	R	homologous flank with attB2r/B3 GGGGACAACTTTGTATAATAA	AGTTGCGGTTTAAACTTTCCTCCCCAGCACGCACAC
		Generation of CRK3 loxP expression and com	plementation vectors
OL4065	F	Amplification of puromycin resistance cassette from pGL631	GATCCTGCAGCGCGTGGATGTCGCGCAG
OL4066	R		GATCGCTAGCCTAGGCACCGGGCTTGCG
OL4293	F	Amplification of SAS-HASPB-mCherry from pGL1893 to integrate at	GATCCTCGAGAATTGCCCGCTTTCCAT
OL4294	R	reporter site	GATCGCGGCCGCGGGATCCTCAATGATGA
OL4316	F	Amplification of GFP from pGL1773 for integration as N-terminal tag	GATCCATATGATGGTGAGCAAGGGCGAG
OL4317	R	, , , , , , , , , , , , , , , , , , , ,	GATCGGTACCCTTGTACAGCTCGTCCAT
OL4318	F	Amplification of 6xHA integration as N-terminal tag	GATCCATATGTACCCTTACGATGTGCCT
OL4319	R		GATCGGTACCTGCGTAATCGGGCACATC
OL4320	F	Amplification of GFP from pGL1773 for integration as C-terminal tag	GATCACTAGTATGGTGAGCAAGGGCGAG
OL4321	R	7 Amplinoadon of O. F. Holli pozitivi o o milogradon do O tominal lag	GATCTCTAGATCACTTGTACAGCTCGTCCAT
OL4541	F	Amplification of SAS-HASPB-mCherry for insertion via HindIII:	GATCAAGCTTAATTGCCCGCTTTCCATTTCG
OL4542	R	enables the replacement of HASPB-mCherry by Xhol and Notl	GATCGCGGCCGCGGGATCCTCAATGATGATGAT
OL4067	F	Amplification of the CRK3 CDS for insertion into the loxP MCS: no	GATCCATATGTCTTCGTTTGGCCGTGTG
OL4103	R	Stop codon amplified due to C-terminal <i>GFP</i> fusion	GATCATCGATCCAACGAAGGTCGCTGAA
OL4103	F	Amplification of the CRK3 CDS for insertion into the loxP MCS: Stop	GATCACTAGTTCTTCGTTTGGCCGTGTGACC
OL4389	R	codon amplified due to N-terminal <i>GFP</i> fusion	GATCTCTAGACTACCAACGAAGGTCGCTGAA
OL4509	F	Amplification of <i>CRK3-his</i> for insertion into pGL2277 to generate an	CTCGAGATGTCTTCGTTTGGCCGT
	R	18S RNA integration vector for complementation of the floxed <i>CRK3</i>	GCGCCGCCTAATGATGATGATGATGATGCCAACG
OL4592	11	inducible deletion line	AAGGTCGCTGAA
OL4601	F	Mutagenesis primers for T178 mutation to a glutamic acid residue to	GCACACCTACGAGCACGAGGTGG
OL4602	R	create CRK3 T178E	ATGGGCACTTGAAACGCAC
014002	l IX		
		Primers for analysis of vector integration and floxed g	
OL4101	F	Internal forward (BLA) and reverse (FKBP12) primers to detect diCre	CTGGTTATGTGTGGGAGG
OL4102	R	integration into the genome	GATGGTTTCCACCTGCAC
OL4287	F	Upstream and downstream primers to amplify the floxed GFP	GCTCGCGTGTTGAGCC
OL4288	R	fragment to detect gene loss by diCre induction	CATTCGTGGGCTCCAGCT
	F	Primers binding out-with the CRK3 integration site	GATCGTGGGAAGGGGAAG
OL4296	1.8		
OL4297	R	Colored and the Consultation of Management of the Colored States o	GGAAGTCCAAGTAGCGCG
OL4297 OL4298	R	Primers binding the CRK3 gene	GGAAGTCCAAGTAGCGCG GGTCACACGGCCAAACGA
OL4297 OL4298 OL4299	R R F	, Medical Section (1997) Section (1997) → Section (1997) → Section (1997)	GGAAGTCCAAGTAGCGCG GGTCACACGGCCAAACGA GCCAAGGAGGCCCTACAG
OL4297 OL4298 OL4299 OL4300	R R F	Primers binding the loxP vector at the 5' splice acceptor site (SAS)	GGAAGTCCAAGTAGCGCG GGTCACACGGCCAAACGA GCCAAGGAGGCCCTACAG GGTGGACGGCTCAACACA
OL4297 OL4298 OL4299	R R F	, Medical Section (1997) Section (1997) → Section (1997) → Section (1997)	GGAAGTCCAAGTAGCGCG GGTCACACGGCCAAACGA GCCAAGGAGGCCCTACAG
OL4297 OL4298 OL4299 OL4300	R R F	Primers binding the loxP vector at the 5' splice acceptor site (SAS)	GGAAGTCCAAGTAGCGCG GGTCACACGGCCAAACGA GCCAAGGAGGCCCTACAG GGTGGACGGCTCAACACA
OL4297 OL4298 OL4299 OL4300 OL4301 OL4781 OL4782	R R F R	Primers binding the loxP vector at the 5' splice acceptor site (SAS) and 3' poly-adenylation site (PAS)	GGAAGTCCAAGTAGCGCG GGTCACACGGCCAAACGA GCCAAGGAGGCCCTACAG GGTGGACGGCTCAACACA GTGTGCTGTGC
OL4297 OL4298 OL4299 OL4300 OL4301 OL4781	R R F R	Primers binding the loxP vector at the 5' splice acceptor site (SAS) and 3' poly-adenylation site (PAS) Upstream and downstream primers for amplification of a floxed	GGAAGTCCAAGTAGCGCG GGTCACACGGCCAAACGA GCCAAGGAGGCCCTACAG GGTGGACGGCTCAACACA GTGTGCTGTGC

Table S1. A list of the oligonucleotides used in this study.

Table S2

pGL No.	Gene ID	Gene Name	Backbone	Description
2313	N/A	diCre	pDONR221	DiCre expression cassette entry vector
2314	N/A	loxP- C-6xHA	pDONR221	LoxP (empty) expression cassette: c-terminal 6xHA tag
2315	N/A	loxP-C-GFP	pDONR221	LoxP (empty) expression cassette: c-terminal GFP tag
2316	N/A	loxP-N-GFP	pDONR221	LoxP (empty) expression cassette: n-terminal GFP tag
2375	LmxM.36.0550	CRK3	pGL631	WT CRK3 ribosomal SSU integration vector
2376	LmxM.36.0550	CRK3 T178E	pGL631	Mutated CRK3 ^{T178E} ribosomal SSU integration vector
2398	N/A	RE9H	pGL631	Red-shifted luciferase bioluminescent protein in G418r pRib
2445	LmxM.36.0550	5' CRK3 flank	pDONR P41-Pr	5' Flank (500bp) ready for Gateway recombination
2446	LmxM.36.0550	3' CRK3 flank	pDONR P2r-P3	3' Flank (500bp) ready for Gateway recombination
2455	N/A	diCre	pDEST R4-R3	DiCre cassette flanked with CRK3 homologous arms
2456	LmxM.36.0550	CRK3	pDEST R4-R3	CRK3-GFP ^{flox} cassette flanked with CRK3 homology
2461	N/A	GFP flox	pGL631	Floxed GFP in pRib: for functional analysis of diCre

Table S2. A list of the plasmids generated in this study.