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A DiCre recombinase-based system for inducible expression in Leishmania major

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Keywords: DiCre recombinase; Leishmania; inducible expression; DNA damage response; 9-1-1 complex; Rad9-Rad1-Hus1.

Abstract

Here we present the establishment of an inducible system based on the dimerizable Cre recombinase (DiCre) for controlled gene expression in the protozoan parasite Leishmania. Rapamycin-induced DiCre activation promoted efficient flipping and expression of gene products in a time and dose-dependent manner. The DiCre flipping activity induced the expression of target genes from both integrated and episomal contexts broadening the applicability of the system. We validated the system by inducing the expression of both full length and truncated forms of the checkpoint protein Rad9, which revealed that the highly divergent C-terminal domain of Rad9 is necessary for proper subcellular localization. Thus, by establishing the DiCre-based inducible system we have created and validated a robust new tool for assessing gene function in Leishmania.
The genus Leishmania encompasses over 20 species, including those that are the causative agents of devastating human diseases worldwide collectively called leishmaniasis [1]. The Leishmania genome is organized in directional gene clusters that may include hundreds of genes from which transcription occurs in a polycistronic fashion. No canonical RNA Pol II promoters have been identified in this parasite and gene expression regulation seems to have been devolved to post-transcriptional processes [2]. The remarkable genome plasticity of Leishmania, which leads to frequent genome rearrangements, not only impacts gene expression control, but also hinders the genetic manipulation of the parasite [3,4]. Therefore, a dependable and robust genetic toolkit is necessary for effective post-genomic functional studies in this protozoan.

Over the past decades, a collection of genetic manipulation tools for Leishmania has been introduced. For instance, transient and stable transfection, gene replacement and disruption, expression vectors and functional complementation and rescue are well-established and reliable tools [5–7]. More recently, the introduction of protein stabilization strategies [8,9] a tetracycline-inducible system for protein expression [10], the dimerizable Cre recombinase (DiCre)-based system for inducible knockouts [11,12] and the establishment of CRISPR cas9 genome editing [13] has further improved our capacity to address peculiar aspects of this parasite’s biology. Given the stringent regulation of DiCre recombinase activity and the variety of strategies for genetic regulation conferred by the use of loxP recombination sites [14,15], we decided to adapt the DiCre-based inducible system for controlled gene expression in Leishmania major. Our strategy involves the generation of a cell line constitutively expressing DiCre recombinase and carrying an inverted gene of interest flanked by cis orientated loxP sites. These constructs are integrated into the 18S rRNA locus and expressed under the control of the Pol I promoter. The antisense orientation of the gene of interest prevents transcription of coding RNA from the positive strand until activation of DiCre recombinase activity by rapamycin treatment. Once activated, DiCre catalyzes the ‘flip’ of the sequence flanked by cis loxP sites, resulting in transcription of a coding RNA and subsequent protein expression (Figure S1A). To prevent continual gene ‘flipping’ by loxP site recombination we employed left-element mutant (lox66) and right-element mutant (lox77) sites [16]. These mutated lox sequences act as sites of recombination to generate a wild-type loxP site and a double mutant Lox72 site for which DiCre has a dramatically reduced affinity. As such, a single recombination
event is favoured upon DiCre recombinase induction, thereby preventing re-

inversion, leading to continual expression of the gene of interest (Figure S1B).

The advantages of this approach include: (i) the use of fewer transfection

rounds and, consequently, fewer selectable markers when compared to the
tetracycline-inducible system developed for L. mexicana by Kraeva and colleagues
[10]; (ii) the possibility to induce expression of gene products from both chromosomal
and/or episomal contexts; (iii) the system promotes a non-leaky expression that can
be induced in a time and dosage-dependent manner; (iv) the possibility to compare
the expression of endogenous and mutated proteins including the conditional
expression of deleterious gene products.

To create the system, plasmid pGL2339 (Figure S2A) was digested and the
array encoding the blasticidin resistance cassette and the dimerizable Cre
recombinase subunits was integrated into the ribosomal locus to generate the
DiCreSSU cell line (Figures 1A and Table S2). Next, the plasmid pGL2332 (Figure
S2B) was digested and the lox66/lox77-flanked 6xHA-GFP cassette containing the
puromycin resistance marker was integrated into the ribosomal locus of the DiCreSSU

cell line to generate the GFP^flox cell line (Figures 1B and Table S2). Both integration
events were confirmed by PCR analysis, which also ascertained that the 6xHA-GFP
coding sequence was present in the antisense orientation in the GFP^flox cell line
(Figure 1C). To test the system and the DiCre flipping activity, the GFP^flox cell line
was incubated with the DiCre dimerization ligand, rapamycin, and the inversion of the
GFP cassette was confirmed by PCR analysis using the appropriate set of primers
(Figure 1D). Semi-quantitative PCR analysis showed that the flipping reaction is time
dependent and seem to reach its maximal level around 96 hours after induction
(Figure S3). Importantly, flipping of the GFP cassette in the absence of rapamycin
was not detectable in the PCR analyses shown in Figure 1D or Figure S3, indicating
a non-leaky DiCre activity in the GFP^flox cells. Upon rapamycin induction, expression
of 6xHA-GFP was detectable after 12 hours and its levels were dose and time-
dependent (Figure 1E). Consistently, 6xHA-GFP was not detectable in the absence
of rapamycin, further confirming the stringent regulation of the system (Figure 1E).

We further used immunofluorescence analysis (IFA) to examine the expression
profile within the population and observed that 6xHA-GFP was detectable by IFA only
after rapamycin incubation (Figure 1F). Consistent with the western blot analysis, the
IFA also demonstrated the rapamycin dose-dependence of the system (Figure S3)
further confirmed by the quantification of GFP corresponding signal (Figure 1G).

Besides confirming the tight regulation of the system, this set of data also
demonstrates that the system is suitable for subcellular compartmentalization studies in this parasite.

To expand the limits of the system we tested it for the ability to flip sequences from an episome, which can be found in multiple copies in the cell. To that end, the plasmid pGL2332 was transfected into the DiCreSSU cell line to generate the pGFPflox cell line (Figure 1H). PCR analysis confirmed the DiCre background and the presence of the target plasmid (Figure 1I), and the flipping of the 6xHA-GFP cassette upon rapamycin incubation (Figure 1J). Consistently, western blot analysis confirmed the expression of the cassette exclusively in rapamycin treated cells (Figure 1K).

These results further demonstrate the tight control of DiCre activity and greatly extend the applicability of the system as a reliable tool for inducible protein expression not only from the genomic context, but also from episomal targets. It is noteworthy that this system it is not expected to be reversible, which can be a disadvantage if reversion of the expressed phenotype is required. However, the irreversibility of the system can be taken an advantageous feature to be explored in both in vitro and in vivo infections assays where inclusion of selection drugs and rapamycin might not be desired.

To further validate the DiCre flipping tool we used the protein Rad9, which participates in the Leishmania DNA Damage Response as part of the checkpoint clamp 9-1-1 [17,18]. In eukaryotic cells, Rad9 has an unstructured C-terminal domain that corresponds to ~1/3 of the protein and is necessary for its function in signalling genotoxic stress [19]. The unstructured C-domain of Leishmania Rad9 is ~3.5x longer than its human counterpart (Figure 2A) and, so far, no function has been reported for this C-terminal extension. To start exploring the function of this C-terminal domain, Rad9 full length or C-terminal truncated encoding sequences, were cloned into the pGL6000 vector (Figure S2C), to generate the Rad9-6xHAflox and Rad9ΔC'-6xHAflox cassettes, respectively. These constructs were digested and integrated into the ribosomal locus of the DiCreSSU cell line to generate the Rad9flox and Rad9ΔCflox cell lines, respectively (Figure 2B and Table S2). Proper integration was confirmed by PCR analysis (Figure 2C) and, as expected, flipping of the cassettes was detected exclusively upon rapamycin incubation (Figure 2D).

Accordingly, expression of Rad9-6xHA and Rad9ΔC'-6xHA was detectable at comparable levels after induction with rapamycin, as demonstrated by western blot analysis (Figure 2E; upper panel). Anti-Rad9 polyclonal serum was used in western blot analysis to evaluate changes in total Rad9 levels (i.e. Rad9-6xHA plus endogenous Rad9) in the Rad9flox cell line after DiCre activation (Figure 2E; middle
Our analysis indicated that, upon induction, Rad9-6xHA was overexpressed when compared to the endogenous Rad9. Quantification of Rad9 signal revealed that up to ~65% of the total Rad9 expressed in these cells corresponded to the induced version of the protein demonstrating that the system can mediate overexpression. Interestingly, the induction of Rad9-6xHA resulted in significant decrease in the levels of endogenous Rad9. However, the same effect was less pronounced when the truncated Rad9ΔC'-6xHA was expressed (Figure 2E; middle panel and Figure 2F). These data suggest that Leishmania Rad9 levels are under tight regulation, which probably involves the participation of its C-terminal extension. Whether this requires a direct or indirect role of this domain remains to be investigated. We were not able to properly assess Rad9ΔC'-6xHA levels relative to endogenous Rad9 using the polyclonal anti-Rad9 serum. One reason for this is that anti-Rad9 serum detects a faster migrating protein with the same molecular mass as Rad9ΔC'-6xHA, hindering its detection. Similarly to the endogenous Rad9, the level of this protein was also reduced upon induction of Rad9-6xHA (Figure 2E; middle panel). While the presence of this band could represent cross-detection of an unrelated protein, these data suggest that it could be a processed form of Rad9 and further characterization is needed to clarify this.

We also analysed the effect of the deletion of the C-terminal extension on the Rad9 subcellular localization. Using IFA, we observed that in the majority of the cells Rad9-6xHA was almost exclusively found in the nuclear compartment, as previously described for Rad9 [17] (Figures 2G and S4). On the other hand, truncated Rad9ΔC'-6xHA presented a less defined localization being prominently detected in the cytoplasm of the majority of the cells (Figure 2G and S4). Consistently, quantitative analysis of the IFA data showed that Rad9-6xHA signal is concentrated in the region containing the nuclear DNA staining, while Rad9ΔC'-6xHA signal expands beyond the nuclear staining limits (Figure 2H). Based on these data, it is reasonable to conclude that the C-terminal extension is necessary for proper subcellular distribution of Rad9 in Leishmania.

In summary, we have demonstrated that the DiCre-based expression system described here is a valuable addition to the Leishmania genetic manipulation toolkit. Its use to study the parasite Rad9 revealed important features of the protein, produced reagents for future studies and proved its value for functional analyses in this parasite.

Acknowledgments
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**Author contribution**

RERSS performed the experiments, collected data, and critically revised the manuscript. GLAS generated and analysed the pGFP\textsuperscript{flox} cell line. EVS conducted the anti-Rad9 western blot analysis in Figure 2E. SMD and JCM generated pGL2332 and pGL2339 plasmids, provided reagents and critically revised the manuscript. JDD generated the DiCre\textsuperscript{SSU} cell line, planned experiments, performed analysis presented in Figure 2A, helped acquire data of Figure 2G and wrote the manuscript. LROT planned experiments and wrote the manuscript.

**References**


Figure 1. Establishment of an inducible DiCre-based expression system in L. major. (A) Schematic representation (not in scale) of the cassette encoding the truncated forms of DiCre after integration into the 18S ribosomal RNA locus (SSU) of L. major wild type cell (LT252) to generate the DiCre\textsuperscript{SSU} cell line; BLA: blasticidin resistance marker. (B) Schematic representation (not in scale) of the 6xHA-GFP cassette (see Figure S1B for details) after integration into the SSU locus of the DiCre\textsuperscript{SSU} cell line (A) to generate the GFP\textsuperscript{flox} cell line; PAC: puromycin resistance marker; upon rapamycin (RAP) addition, the indicated lox sites mediate the flipping of 6xHA-GFP cassette allowing its expression. In (A) and (B), black arrows indicate approximate annealing position of primers used for PCR analysis. (C) PCR analysis of genomic DNA (gDNA) using the indicated set of primers (annealing positions shown in (A) and (B)); DHFR-TS was the loading control for all PCR analyses presented in this work. (D) PCR analysis of gDNA from GFP\textsuperscript{flox} cells cultivated in the presence (+) or absence (-) of 100nM RAP for 48h; primers are indicated below each panel (see annealing positions in (B)). (E) Western blot analysis of total cell extracts from the GFP\textsuperscript{flox} cells cultivated with RAP for the indicated periods of time; two distinct cell-equivalent amount of extracts was analysed; anti-HA was used to detect 6xHA-GFP; EF1\textalpha was the loading control. (F) Immunofluorescence analysis of GFP\textsuperscript{flox} cells cultivated in the presence (+) or absence (-) of 100nM RAP for 48h; images were acquired with a DMI 6000B inverted microscope (Leica); n and k indicate nuclear and kinetoplast DNA, respectively; scale bar = 5\textmu m. (G) GFP\textsuperscript{flox} cells were subject to immunofluorescence as in (F) after cultivation with RAP for 48h; signal corresponding to 6xHA-GFP from individual cells was quantified using Image J software; bars indicate mean +/- Standard Deviation (S.D.); p values by Kruskal-Wallis test were: (***) = 0.0057; (***) = 0.0008. (H) The plasmid pGL2332 (Figure S1B) was transfected as a circular episome in the DiCre\textsuperscript{SSU} cells to generate the pGFP\textsuperscript{flox} cell line; as in (B), RAP addition is expected to induce expression of 6xHA-GFP; schematic representations are not in scale. (I) PCR analysis of gDNA using the indicated set of primers (annealing positions shown in (A) and (H). (J) PCR analysis of gDNA from pGFP\textsuperscript{flox} cells cultivated in the presence (+) or absence (-) of 100nM RAP for 48h; primers are indicated below each panel (see annealing position in (H)). (K) Western blot analysis of total cell extracts from the pGFP\textsuperscript{flox} cells cultivated in the presence (+) or absence (-) of 100nM RAP for 48h; anti-HA was used to detect 6xHA-GFP; EF1\textalpha was the loading control.
**Figure 2. Expression of Rad9 and Rad9ΔC' using the DiCre-based inducible system.** (A) Predicted amino acid sequence of Rad9 from L. major (LmjF.15.0980) and Homo sapiens (NP_004575.1) were subject to disorder prediction using IUPred (http://iupred.enzim.hu); values above 0.5 (horizontal black dotted line) can be considered as disordered regions; horizontal grey bars above each graph indicate the Rad9 domain (Pfam: PF04139) and the disordered C-terminal domain; C-terminal domain of human Rad9 and the corresponding region in L. major Rad9 is indicated as C; the extended C-terminal of L. major Rad9 is indicated as C'. (B) Full length Rad9 from L. major or a truncated version lacking the C-terminal extension (C'), were cloned as C-terminal fusion with a 6xHA tag to generated the Rad9-6xHA<sup>flox</sup> and Rad9ΔC'-6xHA<sup>flox</sup> constructs (not in scale); Rad9-6xHA and Rad9ΔC'-6xHA constructs are arranged in an inverted orientation, flanked by the indicated lox sequences and were integrated into the ribosomal locus (SSU) of the DiCre<sup>SSU</sup> cells (see Figure 1A) to generate the Rad9<sup>flox</sup> and Rad9ΔC'<sup>flox</sup> cell lines, respectively; black arrows indicate approximate annealing position of oligonucleotides used for PCR analyses; PAC: puromycin resistance marker. (C) PCR analysis of gDNA using the indicated set of primers (see annealing positions in (B)). (D) PCR analysis of gDNA from cells cultivated in the presence (+) or absence (-) of 100nM RAP for 48h; primers are indicated below each panel (see annealing positions in (B)). (E) Western blot analysis of total cell extracts from the indicated cells cultivated in the presence (+) or absence (-) of RAP for 48 hours; anti-HA (upper panel) and anti-Rad9 (bottom panel) was used to detect Rad9-6xHA Rad9ΔC'-6xHA; a protein with a similar migration pattern of Rad9ΔC'-6xHA is detected by anti-Rad9 and is indicated with (*); GAPDH was the loading control. (F) Levels of endogenous Rad9 in western blot analysis shown in (E) were determined and normalized with the GAPDH signal, using Image J software; signal from cells exposed to RAP was plotted as a fraction relative to signal from respective non-induced culture. (G) Immunofluorescence analysis of the indicated cells cultivated in the presence (+) or absence (-) of 100nM RAP for 48h; images are representative of a Z-maximal projection from 14 Z-slices acquired with a multiphoton system coupled with LMS780 AxioObserver microscope (Zeiss); n and k indicate nuclear and kinetoplast DNA, respectively; black bars on the DIC field indicate the section where quantification shown in (H) was performed; n and k indicate nuclear and kinetoplast DNA, respectively; scale bar (white) = 5µm. (H) Signal corresponding to DAPI, Rad9-6xHA and Rad9ΔC'-6xHA in the section indicated by the black bar in (G) was quantified with ImageJ.
A. 5'SSU  FKBP-CRE  ➔  FRB-CRE  ➔  BLA  ➔  3'SSU

B. 5'SSU  lox66  6xHA-GFP  PAC  3'SSU

C.  Kb  WT  GFP^{flox}  

D.  GFP^{flox}  

E.  RAP (nM)  0  100  2  

F.  6xHA-GFP  

G.  6xHA-GFP Signal Intensity

H.  lox66  6xHA-GFP  PAC  

I.  Kb  WT  pGFP^{flox}  

J.  pGFP^{flox}  

K.  pGFP^{flox}  

**  ****  ****  ****
Supplementary Information

A DiCre recombinase-based system for inducible expression in *Leishmania*

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Figure S1

(A) DiCre recognizes cis-inverted loxP sites and catalyses the inversion of any flanked sequence; the same loxP sequences are generated upon inversion, therefore re-inversion can proceed with the same efficiency; loxP sequence (5’ to 3’) is shown in the inset.

(B) Right and left mutated elements lox66 and lox77, respectively, are used to flank the sequence of interest; the sequences (5’ to 3’) for lox66 and lox77 are shown in the insets; red lower case indicate the mutated nucleotides; upon inversion, loxP and lox72 sequences are generated and re-inversion occurs at markedly reduced efficiency; upon inversion, PolI promoter-driven transcription of coding RNA from the positive strand mediates the expression of the gene of interest.
Figure S2. Description of the plasmids and strategies used to generate the cell lines reported in this work.

(A) pGL2339 plasmid bears the cassette encoding the inactive truncated forms of Cre-recombinase (DiCre) FKBP12-Cre59 and FRB-Cre60; to generate the DiCreSSU cell line; the plasmid was digested with PacI and PmeI restriction enzymes and transfected into the L. major WT cell line (LT252); cells bearing the integration into the 18S ribosomal locus were cloned by serial dilution in M199 medium containing 10µg/ml blasticidin.

(B) pGL2332 plasmid bears the coding sequence for N-terminal 6xHA-tagged GFP; this fusion is inverted in relation to the 5’ and 3’ SSU sequences, and flanked by lox66 and lox77 sites; to generate the GFP\textsuperscript{flox} cell line, pGL2332 was digested with PacI and PmeI restriction enzymes and transfected into the DiCre\textsuperscript{SSU} cell line; cells bearing the integration into the 18S ribosomal locus were cloned by serial dilution in M199 medium containing 10µg/ml puromycin and 10µg/ml blasticidin.

(C) To generate pGL6000 plasmid, pGL2332 was digested with KpnI and Nhel restriction enzymes to remove the 6xHA-GFP coding sequence and ligated to a PCR product bearing the 6xHA tag and a multiple cloning site containing SdaI, NdeI and BglII restriction sites; the resulting vector allows for cloning of genes of interest as a C-terminal 6xHA fusion flanked by the lox66/lox77 sites; to generate the Rad9\textsuperscript{flox} and Rad9\textsuperscript{ΔC\textsuperscript{flox}} cell lines, pGL6002 and pGL6004 plasmids, respectively, were digested with PacI and PmeI restriction enzymes and transfected into the DiCre\textsuperscript{SSU} cell line; selection and cloning of transfectant parasites was done as in (B).
Figure S3

(A) Schematic representation (not in scale) of the 6xHA-GFP cassette after integration into the SSU locus of the DiCre<sup>SSU</sup> cell line to generate the GFP<sup>flox</sup> cell line; PAC: puromycin resistance marker; upon rapamycin (RAP) addition, the indicated lox sites mediate the flipping of 6xHA-GFP cassette allowing its expression; a, c, d and e indicate approximate annealing position of oligonucleotides used for the PCR analyses shown in (B).

(B) Genomic DNA (gDNA) was extracted from GFP<sup>flox</sup> cell line after cultivation with 100nM RAP for the indicated period of time; ~10ng of gDNA was subjected to semi-quantitative PCR analysis using the indicated set of primers; ~15% of the PCR reaction was resolved in agarose gels; the number of cycles performed in each PCR reaction is indicated at right; annealing position for primers c+d and a+e, which detect only the molecules that has not been flipped, is shown in (A); DHFR-TS was used as the loading control.

(C) Signal corresponding to PCR products using primers c+d and a+e shown in (A) was measured using Image J software and then normalized with DHFR-TS signal; mean values obtained with each set of primers were calculated for each time point and are expressed relative to the 0 hours time point; vertical lines indicate standard error of the mean.

Figure S3. Time course analysis of DiCre-mediated flipping efficiency.

(A) Schematic representation (not in scale) of the 6xHA-GFP cassette after integration into the SSU locus of the DiCre<sup>SSU</sup> cell line to generate the GFP<sup>flox</sup> cell line; PAC: puromycin resistance marker; upon rapamycin (RAP) addition, the indicated lox sites mediate the flipping of 6xHA-GFP cassette allowing its expression; a, c, d and e indicate approximate annealing position of oligonucleotides used for the PCR analyses shown in (B).

(B) Genomic DNA (gDNA) was extracted from GFP<sup>flox</sup> cell line after cultivation with 100nM RAP for the indicated period of time; ~10ng of gDNA was subjected to semi-quantitative PCR analysis using the indicated set of primers; ~15% of the PCR reaction was resolved in agarose gels; the number of cycles performed in each PCR reaction is indicated at right; annealing position for primers c+d and a+e, which detect only the molecules that has not been flipped, is shown in (A); DHFR-TS was used as the loading control.

(C) Signal corresponding to PCR products using primers c+d and a+e shown in (A) was measured using Image J software and then normalized with DHFR-TS signal; mean values obtained with each set of primers were calculated for each time point and are expressed relative to the 0 hours time point; vertical lines indicate standard error of the mean.
**Figure S4.** GFP\textsuperscript{flox} cell line was cultivated in the presence of the indicated rapamycin concentrations for 48 hours; cells were fixed and subjected to immunofluorescence analysis using mouse anti-HA as primary antibody and anti-mouse conjugated with Alexa-Fluor 488 as secondary antibody (false coloured green); images were acquired with a DMI 6000B inverted microscope (Leica); scale bar = 25\textmu m
Figure S5. Rad9^flox^ and Rad9ΔC^flox^ cell lines were cultivated in the presence (+) or absence (-) of rapamycin for 48 hours; cells were fixed and subjected to immunofluorescence analysis using mouse anti-HA as primary antibody and anti-mouse conjugated with Alexa-Fluor 488 as secondary antibody; images were acquired as a series of 14 Z-slices with a multiphoton system coupled with LMS780 AxioObserver microscope (Zeiss); images are representative of a Z-maximal projection from 14 Z-slices; scale bar = 25 µm.
**Supplementary Table 1: Primers used in this work**

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<td>Duncan et al., 2016</td>
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<td>b</td>
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<tr>
<td>d</td>
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<td>e</td>
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<td>f</td>
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**Supplementary Table 2: Cell lines used in this work**

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<tr>
<td>Rad9ΔClox</td>
<td>SSU DiCre BLA SSU lox66((Rad9ΔC::6xHA)_AS)lox77 PAC</td>
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*AS = Anti Sense*
**Experimental procedures**

**Parasites culture**
*L. major* LT252 (MHOM/IR/1983/IR) and all other cell lines were cultured as promastigotes at 26°C in M199 medium plus 10% heat-inactivated fetal bovine serum. Generation of cell lines was done using the transfection protocol previously described (Kapler et al., 1990).

**PCRs and cloning**
Genomic DNA was extracted with DNeasy Blood & Tissue Kit (QIAGEN) according to manufacturer instructions. PCRs were performed using Taq DNA Polymerase (Thermo) for products up to 2kb or Phusion® High-Fidelity DNA Polymerase (NEB) for products with more than 2kb.

For generation of pGL6002 plasmid, the sequence coding for the 718 amino acids of full length Rad9 was amplified from gDNA using Rad9NdelFw and Rad9ABCsdalRv as primers (Table S1). For generation of pGL6004 plasmid, the sequence coding for the first 370 amino acids of Rad9 was amplified from gDNA using Rad9NdelFw and Rad9ABCsdalRv as primers (Table S1).

**Antibodies and western blotting analyses**
Rabbit anti-LmRad9 antibody was previously described (Damasceno et al., 2013) and was used at 1:3 000 dilution. Rabbit anti-GAPDH antibody was kindly provided by Dr. Paul Michels and was used at the 1:15 000 dilution. The commercial antibodies used were as follows: mouse anti-HA (Sigma-Aldrich) at 1:1000 dilution, mouse anti-EF1α (Merck Millipore) at 1:50 000 dilution. HRP-conjugated anti-mouse and anti-Rabbit antibodies (GE Life Sciences) were used at 1:10 000 and 1: 30 000 dilution, respectively.

Unless otherwise indicated, total cell extracts equivalent to ~10^6 cells were resolved by SDS-PAGE, transferred to PVDF membrane (GE Life Sciences) and analyzed with the indicated antibodies. Bands were detected with ECL Prime Western Blotting Detection Reagent (GE Life Sciences) and visualized with Hyperfilm ECL (GE Life Sciences).

**Immunofluorescence**
Cells were fixed with 4% paraformaldehyde for 10 min at room temperature. Fixed cells were adhered to poly-L-lysine coated glass slides and permeabilized with 0.3% Triton X-100 for 30 min. Cells were pre-blocked with 2% BSA in 1x PBS for 1 hour. Anti-HA antibody was used at 1:1000 dilution to detect 6xHA-GFP, Rad9-6xHA and Rad9ΔC-6xHA. Primary antibody was visualized with anti-mouse secondary antibody conjugated with Alexa Fluor 488 or Alexa Fluor 594 (Invitrogen). Slides were mounted with ProLong® Gold Antifade Reagent (Thermo) and 2µg/ mL Hoechst 33342 (Thermo).
Plasmids
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