Optimized CRISPR-Cas9 Genome Editing for *Leishmania* and Its Use To Target a Multigene Family, Induce Chromosomal Translocation, and Study DNA Break Repair Mechanisms

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**ABSTRACT**  CRISPR-Cas9-mediated genome editing has recently been adapted for *Leishmania* spp. parasites, the causative agents of human leishmaniasis. We have optimized this genome-editing tool by selecting for cells with CRISPR-Cas9 activity through cotargeting the miltefosine transporter gene; mutation of this gene leads to miltefosine resistance. This cotargeting strategy integrated into a triple guide RNA (gRNA) expression vector was used to delete all 11 copies of the A2 multigene family; this was not previously possible with the traditional gene-targeting method. We found that the *Leishmania donovani* rRNA promoter is more efficient than the U6 promoter in driving gRNA expression, and sequential transfections of the oligonucleotide donor significantly eased the isolation of edited mutants. A gRNA and Cas9 co-expression vector was developed that was functional in all tested *Leishmania* species, including *L. donovani*, *L. major*, and *L. mexicana*. By simultaneously targeting sites from two different chromosomes, all four types of targeted chromosomal translocations were generated, regardless of the polycistronic transcription direction from the parent chromosomes. It was possible to use this CRISPR system to create a single conserved amino acid substitution (A189G) mutation for both alleles of *RAD51*, a DNA recombinase involved in homology-directed repair. We found that RAD51 is essential for *L. donovani* survival based on direct observation of the death of mutants with both RAD51 alleles disrupted, further confirming that this CRISPR system can reveal gene essentiality. Evidence is also provided that microhomology-mediated end joining (MMEJ) plays a major role in double-strand DNA break repair in *L. donovani*.

**IMPORTANCE**  *Leishmania* parasites cause human leishmaniasis. To accelerate characterization of *Leishmania* genes for new drug and vaccine development, we optimized and simplified the CRISPR-Cas9 genome-editing tool for *Leishmania*. We show that co-CRISPR targeting of the miltefosine transporter gene and serial transfections of an oligonucleotide donor significantly eased isolation of edited mutants. This cotargeting strategy was efficiently used to delete all 11 members of the A2 virulence gene family. This technical advancement is valuable, since there are many gene clusters and supernumerary chromosomes in the various *Leishmania* species and isolates. We simplified this CRISPR system by developing a gRNA and Cas9 coexpression vector which could be used to delete genes in various *Leishmania* species. This CRISPR system could also be used to generate specific chromosomal translocations, which will help in the study of *Leishmania* gene expression and transcription control. This study also provides new information about double-strand DNA break repair mechanisms in *Leishmania*.

**KEYWORDS**  CRISPR-Cas9, chromosomal translocation, co-CRISPR targeting, homology-directed repair, *Leishmania*, MMEJ, microhomology-mediated end joining, Rad51, double-strand break repair, genome editing, miltefosine transporter, multiple gene family

Received 15 November 2016  Accepted 16 December 2016  Published 18 January 2017

Citation  Zhang W-W, Lypaczewski P, Matlashewski G. 2017. Optimized CRISPR-Cas9 genome editing for *Leishmania* and its use to target a multigene family, induce chromosomal translocation, and study DNA break repair mechanisms. mSphere 2:e00340-16. https://doi.org/10.1128/mSphere.00340-16.

Editor  Ira J. Blader, University at Buffalo

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CRISPR genome editing in Leishmania has been optimized and used to delete A2 multigene family and induce targeted chromosomal translocation
Leishmaniasis is a vector-borne disease caused by the protozoan parasite species of the genus *Leishmania*. Depending on the species, *Leishmania* infection can cause mild self-healing cutaneous leishmaniasis (CL), disfiguring mucocutaneous leishmaniasis (MCL), or fatal visceral leishmaniasis (VL), which is the second-deadliest parasitic disease after malaria (1, 2). Approximately 1 billion people worldwide are at risk of infection, and more than 1.3 million new infections occur each year. Despite decades of research, there is still no vaccine, and treatment of leishmaniasis relies on drugs which are expensive, toxic, and are at risk for resistance development (1, 2).

The *Leishmania* genome contains over 8,000 genes, and most of these genes have unknown functions (3–6). Since its introduction to *Leishmania* research nearly 3 decades ago, the traditional gene targeting method involving homologous recombination using antibiotic selection marker genes has greatly contributed to the understanding of *Leishmania* biology and pathogenesis. This homologous recombination method is, however, time-consuming, limited by available antibiotic selection markers, and not well-suited for introducing point mutations and other genome-editing tasks. Thus, a simpler yet more efficient and versatile genome-editing method is required to accelerate the characterization of *Leishmania* genes for new drug target identification and vaccine development.

We have recently developed vectors expressing the Cas9 nuclease and guide RNA (gRNA) for *Leishmania* spp., and we demonstrated that CRISPR-Cas9 is an effective genome engineering tool for *L. donovani* (7). It was revealed that *L. donovani* mainly uses homology-directed repair (HDR) and microhomology-mediated end joining (MMEJ) to repair Cas9-generated double-strand DNA (dsDNA) breaks and that the nonhomologous end-joining (NHEJ) pathway appears to be absent in *L. donovani*. MMEJ resulted in deletion mutations ranging from 10 to more than 3,000 bp. The activity of different gRNAs can vary significantly. The use of an oligonucleotide donor, antibiotic selection marker donor, and double gRNA expression vector greatly improved the precision and efficiency of CRISPR-Cas9-mediated genome editing (7).

In this study, we further optimized and simplified CRISPR-Cas9-mediated genome editing in *Leishmania* through several approaches. Single vectors capable of expressing both the gRNA and Cas9 nuclease were developed which were functional in all tested *Leishmania* species, including *L. donovani*, *L. major*, and *L. mexicana*. Sequential transfections of a gene-editing oligonucleotide donor significantly eased the isolation of the edited mutants. Cotargeting the miltefosine transporter gene (*MT*) followed by miltefosine selection greatly increased the efficiency of editing a second target gene in parallel. This cotargeting strategy, integrated into a multiple gRNA expression vector, was used to delete all 11 members of the A2 gene family in *L. donovani*, an unattainable task with the traditional gene targeting method (8). We generated specific chromosomal translocations by simultaneously targeting sites from different chromosomes. We demonstrated that RAD51, a DNA recombinase involved in HDR, is essential for *L. donovani*. However, analysis involving the inhibition of RAD51 strongly suggested that MMEJ plays the major role in double-strand DNA break repair in *Leishmania*.

**RESULTS AND DISCUSSION**

**Sequential transfections of an oligonucleotide donor significantly improve CRISPR-Cas9 gene-editing efficiency.** We previously showed that the addition of an oligonucleotide donor improved CRISPR-Cas9 gene disruption efficiency in *L. donovani* (7). In some cases, however, a gene-editing task, such as the introduction of an epitope tag or a point mutation, may involve a site where only low-activity gRNAs can be designed, such as the gRNA-targeting site in the *L. donovani* miltefosine transporter gene *MT* (7). We therefore wanted to determine whether sequential transfections of an oligonucleotide donor containing 25-nucleotide-long flanking homologous sequences to the Cas9 break site would increase the gene-editing efficiency and improve the isolation of the edited mutants (Fig. 1A; see also Data Set S1 in the supplemental material). The *MT* gene was selected for evaluating CRISPR-Cas9 gene-editing efficiency (frequency), since mutations (insertions, deletions, and selected point mutations) in the
MT gene lead to survival (resistance) in the presence of miltefosine (7). Twenty-one days following transfection of vectors expressing gRNAc and Cas9 in L. donovani (21 days is the time required to select for cells with stable expression), sequential transfections with an oligonucleotide donor containing stop codons were carried out every 3 days for a total of four transfections (Fig. 1B). The miltefosine resistance rate was determined 3 days after each oligonucleotide donor transfection. As shown in Fig. 1C, multiple transfections increased the miltefosine resistance rate significantly, compared to gRNAc expression alone. At the end of the fourth donor oligonucleotide transfection, the final resistance rate increased from 0.4% without a donor to nearly 10%, representing a 25-fold increase. This demonstrated that sequential transfections of an oligonucleotide donor can significantly improve CRISPR-Cas9 gene editing (oligonucleotide donor-directed repair) efficiency to make it easier to isolate the edited mutants. Since the growth rate can be altered in edited mutants, it is necessary to clone transfectants into 96-well plates within 2 days after the third or fourth donor transfection. We have successfully used this strategy to generate L. major and L. mexicana centrin gene null mutants (data not shown) and the single-amino-acid substitution mutant for L. donovani RAD51 (see below).

Deletion of the multicopy A2 gene family by cotargeting the miltefosine transporter (MT) gene and selection with miltefosine. One of the key advantages of
CRISPR-Cas9 technology is that the Cas9/gRNA complex will continually scan the genome and generate double-strand DNA breaks until all the targeting sites in the genome have been deleted or mutated. This has been used successfully to inactivate multicopy family genes and endogenous retroviruses in other cell types (9, 10). A2 is a multicopy gene family and an important virulence factor for visceral Leishmania infection (6, 8, 11–13). There are at least 11 copies of the A2 gene of various sizes, which alternate with A2rel genes and are flanked by 5’ A2rel and 3’ A2rel genes. The A2 and MT gRNA-targeting sites and primers used to verify A2 gene deletion are indicated. Note that this putative A2-A2rel gene cluster is based on our previous A2-targeting study (8) plus recent unpublished PacBio genome sequencing data. Because of multicopies and repeated sequences, the A2-A2rel gene cluster loci are not properly assembled in published L. donovani and L. infantum genomes (TriTrypDB). (B) The double- and triple-gRNA expression vectors used to target A2 and LdMT genes. rRNAP, L. donovani rRNA promoter; H, HDV ribozyme; HH, Hammerhead ribozyme. Black boxes represent the 92-bp pyrimidine track. The drawing is not to scale. (C) Western blot analysis of A2 proteins in L. donovani transfected with the double- or triple-gRNA expression vectors (as in panel B), with or without miltefosine selection. Equal loading of cell lysates was verified by reprobing the membrane with anti-HSP83 antibodies. (D) PCR verification of A2 null mutants using A2-specific primers L and R. The genomic DNA quality for each sample was verified by PCR with RAD51-specific primers.

CRISPR-Cas9 technology is that the Cas9/gRNA complex will continually scan the genome and generate double-strand DNA breaks until all the targeting sites in the genome have been deleted or mutated. This has been used successfully to inactivate multicopy family genes and endogenous retroviruses in other cell types (9, 10). A2 is a multicopy gene family and an important virulence factor for visceral Leishmania infection (6, 8, 11–13). There are at least 11 copies of the A2 gene of different sizes in L. donovani 15C12D (8, 12) (Fig. 2A). Due to these multiple copies that also alternate with another gene termed A2rel, it was not possible to delete this gene family from L. donovani using the conventional gene-targeting approach (8). We therefore attempted to delete all copies of this multigene family from chromosome 22 of L. donovani by using the CRISPR-Cas9 method. A2 gene-coding sequences are mainly composed of 30-nucleotide repeat sequences encoding 10 amino acid repeats (11, 14) (Fig. 2A). To avoid these repeated sequences from being used as an HDR template, two gRNAs targeting the unique sequences (one targeting near the 5’ end of the A2 coding sequence and the other targeting the 3’ untranslated region) (Fig. 2A) in each of the A2 genes were designed and cloned into the dual gRNA expression vector (Fig. 2B; Data Set S2). In addition to targeting A2 genes, we also investigated whether it was possible to increase the efficiency of A2 gene deletion by coselecting for cells with CRISPR-Cas9 activity. Coselection for CRISPR-Cas9 activity was performed by targeting the L. donovani miltefosine transporter gene (LdMT) and selecting for miltefosine resistance at the
same time as targeting the A2 genes. The rationale was that if one gRNA were used to target the *LdMT* gene and the other gRNA(s) targeted a different gene of interest (in this case, A2), following selection for miltefosine resistance the A2 gene would be targeted with a higher frequency. Thus, the *LdMT* gRNAa coding sequence was added into the gRNA A2a+b construct to generate a triple gRNA expression vector to determine whether A2 genes could be more efficiently deleted in these miltefosine-resistant cells (Fig. 2B).

As shown by Western blotting analysis in Fig. 2C, with prolonged culture and without cloning, the A2 genes could be completely deleted (inactivated) with this CRISPR system (Fig. 2C, lane 5). However, coselection for CRISPR-Cas9 activity with miltefosine significantly reduced the time needed to delete all A2 genes, from 4 months with no selection (Fig. 2C, lane 5) to 6 weeks with miltefosine selection (Fig. 2C, lane 4). In contrast, in the absence of miltefosine there remained detectable but diminished A2 protein expression at 6 weeks (Fig. 2C, lanes 2 and 3). Six weeks is the minimum time to establish miltefosine-resistant cells in culture. The complete deletion of A2 genes in these *L. donovani* cells was further confirmed by PCR analysis with A2-specific primers (Fig. 2D). This demonstrated that this CRISPR system can be used to delete multicopy family genes and that coselection for CRISPR-Cas9 activity can significantly reduce the time needed to obtain these deletion mutants. This is valuable, since there are many tandem gene arrays and supernumerary chromosomes in various *Leishmania* species and isolates (3–6).

This coselection observation is similar to what was reported for *Caenorhabditis elegans* and in human cells, where the co-CRISPR strategy has greatly facilitated detection of genome-editing events; particularly, CRISPR cotargeting the hypoxanthine phosphoribosyltransferase (HPRT) gene in human cells followed by 6-thioguanine selection highly enriched the cotargeting gene edited mutants (15–17). This also agrees with the observation of a bimodal distribution of CRISPR inactivation of porcine endogenous retroviruses (PERVs), where only 10% of clones exhibited complete disruption of all 62 copies of PERV *pol* genes and the remaining clones exhibited no or a low level of editing (10).

**Chromosomal translocation by targeting two sites from different chromosomes simultaneously.** Targeted chromosomal translocations have been successfully generated in various organisms by using CRISPR-Cas9 technology, including various cancer models (18–21). The ability to generate targeted chromosomal translocations in *Leishmania* will help to investigate gene expression and identify mechanisms for the initiation and termination of polycistronic transcription and chromosome stability (22–24). To determine whether specific chromosomal translocations could be generated in *Leishmania*, a dual gRNA expression plasmid was constructed where one gRNA was targeted to the nonessential multidrug resistance gene *Ld241510* in chromosome 24 (25) (Data Set S3) and the other was targeted to the miltefosine transporter gene (*LdMT*) in chromosome 13 (7) (Fig. 3A). The hypothesis was that although most of the dsDNA breaks (DSBs) generated by Cas9 nuclease in chromosomes 13 and 24 would be repaired by intrachromosomal joining, some interchromosomal (translocation) joining could also occur. Following transfection and miltefosine selection, genomic DNA was extracted from the surviving cells, and various PCR primer pairs (one primer specific for chromosome 13 and the other for chromosome 24, each close to one of the two DSB sites) (see Data Set S3 for details), were used to investigate chromosomal translocation. As shown in Fig. 3B, it was possible to detect all four types of chromosomal translocation events resulting from the two DSBs generated simultaneously in chromosomes 13 and 24. This demonstrated that these translocations can occur regardless of the polycistronic transcription direction or the size of the new chromosome generated by translocation. In addition, following cotransfection of two oligonucleotide donors that promoted competitive translocation events (type II or type IV), both translocation PCR products were detected at similar frequencies (Fig. 3C). This further suggests that the frequency of chromosomal translocation events may not be affected by the direction of polycistronic transcription.
FIG 3  Targeted chromosomal translocations generated by targeting sites from two different chromosomes simultaneously. (A) The double-gRNA vector used to coexpress LdBPK_241510.1 targeting gRNA (241510), the LdMT targeting gRNAa (MT), and a schematic of chromosomes 13 and 24, with the Cas9 cleavage sites and the polycistronic transcription directions indicated. (B) Schematic of the 4 types of chromosomal translocations detected following transfection with the gRNA241510+MT coexpression vector. The chromosomal translocation junction sequences joined by MMEJ or a transfected oligonucleotide donor-directed repair are also included. Note that the polycistronic transcription directions and the numbers indicating chromosome size, which could have been altered in the newly generated fused chromosomes after translocation, are directly transferred from the parent chromosomes 13 and 24. (C) PCR detection of type II and type IV chromosomal translocations in cells expressing 241510- and MT-targeting gRNAs following transfection with the mixture of type II and type IV oligonucleotide donors (see the sequences in panel B). Primer 13L2, Ld131590L2; 24L1, Ld241510L1; 24R1, Ld241510R1. (D) Chromosomal translocation detected after L. donovani cells transfected with gRNA A2a+H11001+MT coexpression vector (Fig. 2B). For simplicity, only one A2 gene and one Cas9 cleavage site are represented for the A2-A2rel gene cluster loci in chromosome 22. See the supplemental material for all primer pairs used to detect these chromosomal translocations.
It is interesting that the two gene clusters from chromosomes 13 and 24 were joined together back to back in the type I chromosomal translocation, despite transcription going in opposite directions. In contrast, the other two gene clusters were joined together head to head in the type II chromosomal translocation. Since these two gene clusters in type I chromosomal translocation lose their corresponding transcription initiation sequences, it will be interesting to see how the transcription levels of these gene clusters from the parent chromosomes are affected by these chromosomal translocations. To our knowledge, this could be the first example to show that all four types of chromosomal translocations can be generated (18–21). Interestingly, we also detected a chromosomal translocation event (Fig. 3D) in Leishmania cells transfected with the triple gRNA expression vector described above in Fig. 2B, in which two gRNAs were targeted to the A2 loci in chromosome 22 and one gRNA was targeted to the MT locus in chromosome 13.

As expected, in the absence of oligonucleotide donors, all the chromosomal translocations were joined by MMEJ. It is interesting that two MMEJs were observed in both the detected type III and IV chromosomal translocations. In the type III chromosomal translocation detected, one MMEJ (TCCAC) joined sequences from each side of the break in chromosome 13 before the second MMEJ made the chromosomal translocation joint, which used only a 3-bp microhomology sequence (CCA). In the detected type IV chromosomal translocation, a section of chromosome 24 sequence (more than 200 bp) was reversed, which was likely caused by a flip of the single-strand DNA created by an end resection after the double-strand break and the intramolecular MMEJ (ACGACACCATT). Though the chromosomal translocation events were relatively rare and were enriched by cotargeting the LdMT gene in the current study, it should be feasible to isolate chromosomal translocation mutants in other specific targeting sites by using donors containing drug selection markers.

The RNA polymerase I rRNA promoter is more efficient than the RNA polymerase III U6 promoter in driving gRNA expression in Leishmania. Due to its precise transcription initiation and termination, the RNA polymerase III U6 promoter has been widely used to drive gRNA expression in higher level eukaryotic cells and other protozoan parasites, including L. major (26–30). In contrast, the RNA polymerase I rRNA promoter used for gRNA expression has only been reported in L. donovani (7) and Trypanosoma cruzi (31). We therefore compared genome-editing efficiency when gRNA expression was under control of these two different RNA polymerase promoters. An LdMT (L. donovani miltefosine transporter gene) gRNAa expression vector using the L. donovani U6 promoter was constructed and compared to the rRNA promoter in the pSPneoGRNAaH vector, which expresses the same LdMT gRNAa previously described (7). In addition, we made an expression vector in which the human U6 promoter was used to direct LdMT gRNAa expression (Fig. 4A; Data Set S4). These LdMT gRNAa expression vectors were transfected into Cas9-expressing L. donovani promastigotes, and resistance to miltefosine was determined. As shown in Fig. 4B, the miltefosine resistance rate was much lower in the LdU6 promoter vector-transfected cells than in the rRNA promoter vector-transfected cells at 32 days posttransfection, indicating that the rRNA promoter is more efficient at driving gRNA expression in L. donovani. Interestingly, while the miltefosine resistance rate for cells using the rRNA promoter appeared to be stabilized at 12 to 25% in prolonged culture, the miltefosine resistance rate in cells using the LdU6 promoter was able to reach a similar level after a much longer time period of 73 days posttransfection (Fig. 4B). It is also important to note that because the rRNA promoter is a stronger promoter than the LdU6 promoter (32, 33), it was much easier to obtain G418-resistant transfectants from the rRNA promoter vector than from the LdU6 promoter vector-transfected Leishmania cells (data not shown). Surprisingly, the human U6 promoter was also functional in Leishmania and could mediate gRNA expression, though with a much lower targeting efficiency. Taken together, this comparison demonstrated that the rRNA promoter is much better than the U6 promoter in driving gRNA expression in Leishmania. Thus, it would be interest
to explore the RNA polymerase I promoter for improving gRNA expression in other organisms, including human cells.

**Generation of gRNA and Cas9 coexpression CRISPR vectors for use with different Leishmania species.** To further simplify this genome-editing system in *Leishmania*, single coexpression vectors (pLdCN and pLdCH) were constructed where the transcription of gRNA, Cas9, and a drug selection maker (neomycin or hygromycin resistance gene) were placed under control of the same rRNA promoter (Fig. 5A; Data Set S5). The *LdMT* gRNA sequence was used to test whether these vectors could function as predicted. As shown in Fig. 5B, although there were some variations in the targeting efficiency, miltefosine-resistant cells were obtained with both pLdCNgRNAa and pLdCHgRNAa, demonstrating that the gRNA and Cas9 nuclease were properly expressed in both of these coexpression vectors. It is important to note that the *L. donovani* ribosomal promoter also functions well in other *Leishmania* species (32, 33).

So far, we have successfully used the pLdCN vector to delete genes in *L. major*, *L. mexicana*, and *L. donovani* (data not shown). Therefore, a single appropriately designed gRNA construct could be used to target a conserved site (a 20-bp conserved sequence plus NGG, known as the protospacer-adjacent motif) in all three and perhaps more *Leishmania* species.
RAD51, a DNA recombinase involved in HDR, is essential for *L. donovani*. We previously observed that DSBs created by CRISPR-Cas9 in Leishmania were repaired by interallelic HDR, which results in error-free repair, or by MMEJ, which results in deletion mutations (7) (see Fig. 7B, below). RAD51 is a DNA recombinase required for HDR during DSB repair and is not involved in MMEJ (34–37). To determine whether inhibition of the HDR pathway would increase the frequency of MMEJ-mediated DSB repair to improve CRISPR-Cas9-directed gene inactivation efficiency, we attempted to use this CRISPR system to disrupt the *L. donovani RAD51* gene. As shown in Fig. 6A (see also Data Set S6), a plasmid carrying an *L. donovani RAD51*-specific gRNA was transfected into Cas9-expressing *L. donovani* cells, and subsequently a single-strand oligonucleotide donor with stop codons or a bleomycin resistance marker donor PCR product was cotransfected into these cells as described for the experiments shown in Fig. 1.

Although it was possible to disrupt one of the *L. donovani RAD51* alleles by using the donors with or without phleomycin selection, as verified by PCR analysis after cloning, one wild-type RAD51 allele remained in the surviving cells (Fig. 6A and B). Even after two *RAD51* alleles were disrupted when we used a combination of the stop codon oligonucleotide donor and the bleomycin marker gene donor, a third wild-type *RAD51* allele still persisted (Fig. 6A and B). The *RAD51* gRNA-expressing cells did however proliferate slower than control gRNA-expressing cells, likely because the *RAD51* gRNA and Cas9 complex were continually targeting the remaining *RAD51* allele (Fig. 6C). Indeed, after cloning the *RAD51* gRNA-expressing cells (*Ld* *RAD51*+/−) in 96-well plates, at least 10 of these single-cell clones died out after continuous culture for 2 weeks. Interestingly, after cloning, many of these putative *RAD51* null mutants were able to continue multiplying slowly, as a clump to as many as 100 parasites before crashing, indicating it would take some time to dilute and degrade the remaining wild-type *RAD51* mRNA and proteins in these null mutant cells (Fig. 6D). Taken together, this demonstrated that like in mammalian cells, the *RAD51* gene is essential for *L. donovani*.

In contrast to *L. donovani*, a *RAD51* gene null mutant has been generated in *L. infantum* (37). Since a low frequency of homologous recombination events could still be detected in *L. infantum RAD51* null mutants, *L. infantum* may have developed ways (other recombinases) to compensate for *RAD51* deficiency (37). Interestingly, while it was possible to generate *RAD51* null mutants in *Trypanosoma brucei* and *L. infantum,*
FIG 6 RAD51 is essential for L. donovani. (A) Strategies used to generate RAD51 disruption mutants and a mutant with a single conserved amino acid substitution. To generate various RAD51 mutants, L. donovani cells were transfected with Cas9- and RAD51-targeting gRNA expression vectors followed by transfection of oligonucleotide donors (with stop codons and an EcoRI site or with a conservative amino acid substitution) and/or the bleomycin selection marker donor. Genomic DNA from these L. donovani cells (clones) were subjected to PCR, restriction enzyme digestion, and sequencing analysis. (B) PCR and restriction enzyme analysis of RAD51 single- and double-allele disruption mutants. (Left) PCR amplification of the RAD51 sequence with primers L and R, followed by EcoRI digestion. Lane 1, wild-type L. donovani; lane 2, RAD51+/− mutant with a single RAD51 allele disrupted by the oligonucleotide donor containing stop codons and an EcoRI site. Note that although the EcoRI-digested bands (479 and 435 bp) were detected, the 894-bp wild-type RAD51 allele band remained in this single RAD51 disruption mutant. (Middle) PCR analysis of bleomycin resistance clones (RAD51+/− mutants) with primers L and R. Both the 1,428-bp bleomycin marker insertion band and the 894-bp wild-type RAD51 allele band were detected in all these bleomycin resistance clones. Note that sequencing indicates that the additional bands detected are rearrangements of the 1,428-bp ble marker bands. (Right) PCR and EcoRI digestion analysis of Rad51+/−/+ mutants with one allele disrupted with stop codons and an EcoRI site containing oligonucleotide donor and the other allele with a bleomycin selection marker donor. PCR bands (not shown) similar to those in the middle panel, including the 1,428-bp ble marker bands and the approximately 900-bp bands, were obtained from these mutants. The approximate 900-bp bands were then extracted from the gel and subjected to complete EcoRI digestion. Note that the 894-bp wild-type (WT) RAD51 allele band remained in these mutants. These are representative data of more than 100 clones analyzed. (C) Growth curves of L. donovani cells targeted by RAD51 gRNA: RAD51+/− (Oligo donor), RAD51+/− mutant with the oligonucleotide donor (stop codons) insertion, RAD51+/− Ble donor, RAD51+/− mutant with bleomycin selection marker donor insertion, and RAD51+/−, wild-type L. donovani cells expressing a control gRNA targeting the LdMT gene. The data are representative of three independent experiments. (D) Microscope images showing that disruption of all RAD51 alleles is lethal for L. donovani. The RAD51+/− (Continued on next page)
RAD51 could also be essential for *L. major* and *T. cruzi*, as no RAD51 null mutants have been reported in these latter parasites despite attempts (38–41).

Although it was not possible to disrupt or introduce stop codons into both wild-type RAD51 alleles in surviving *L. donovani* cells, we were able to edit both RAD51 alleles with an oligonucleotide donor designed to generate a single conserved amino acid substitution (A189G) (Fig. 6A, E, and F). This may be the first example of engineering a chromosomal single amino acid change in *Leishmania*, further revealing the importance and versatility of CRISPR-Cas9 gene editing for *Leishmania*.

**Microhomology-mediated end joining plays a dominant role in double-strand DNA break repair in Leishmania.** Since we were not able to generate a RAD51 null mutant, as it is essential for *L. donovani*, we attempted to inhibit RAD51 activity by using the RAD51 inhibitors B02 and RI-1 (42–44). We reasoned that impairing HDR with these inhibitors could induce higher levels of MMEJ and increase CRISPR-Cas9 gene inactivation efficiency. However, there was no increase in *LdMT* gene inactivation through MMEJ in *LdMT* gRNA-expressing cells after we used various concentrations of these RAD51 inhibitors (Fig. 7A), suggesting that HDR may play a less dominant role than previously anticipated for DSB repair in *Leishmania* (7). The B02 and RI-1 inhibitors did, however, confirm that RAD51 is essential, since *L. donovani* cells could not survive.

**FIG 6 Legend (Continued)**

mutant cells, which continue expressing RAD51-targeting gRNA were cloned in 96-well plates, and cell growth was monitored by microscopy. The image for RAD51−/− cells was taken 1 week after cloning; the images for RAD51−/− cells were taken 3 weeks after cloning. (E) Partial sequence of the oligonucleotide donor with mutations resulting in a single conservative amino acid substitution of the RAD51 protein (A189G) and inactivation of the RAD51 gRNA-targeting site. (F) Direct sequencing of the PCR product amplified from an *L. donovani* clone, showing both alleles of RAD51 have been mutated to the sequence of the oligonucleotide donor (see panels A and E).
in medium containing more than 15 μM B02 or 70 μM RI-1, though the general toxicity of these RAD51 inhibitors might also have contributed in part to the death of these Leishmania cells.

The role that HDR and MMEJ play in DSB repair was further investigated by comparing the repair rates of MT genes in wild-type L. donovani (LdMT+/+) cells and L. donovani cells with one MT gene replaced with a bleomycin gene (LdMT+/-) by using traditional gene replacement. LdMT+/+ and LdMT+/- mutants were then transfected with vectors expressing Cas9 and three different LdMT gene-targeting gRNAs designated a, b, and c (7). If HDR indeed played a major role in DSB repair, the miltefosine resistance rate (i.e., mutation rate) would be expected to be much higher in LdMT+/- cells than in wild-type LdMT+/+ cells. This is because there is no wild-type MT allele available as an HDR template, and cleavage of the single wild-type MT allele would result in miltefosine resistance either through HDR or through MMEJ (Fig. 7C). In the wild-type LdMT+/+ cells, the majority of the DSBs generated in one of the MT alleles would be repaired by HDR (error free) with the remaining intact wild-type allele as the template, and it would therefore take longer for resistance to develop (Fig. 7B). Interestingly, however, there was no increase in miltefosine resistance rates in LdMT+/- cells compared to the wild-type LdMT+/+ cells (Fig. 7D). This strongly suggests that MMEJ is more efficient than HDR in DSB repair in Leishmania, which agrees with the above RAD51 inhibition data.

Given that significant increases in gene-editing efficiency have been observed when using oligonucleotide and double-stranded DNA donors with only 25 nt (bp) micro-homology flanking sequences in this and our previous studies (7), these donors could actually be the ideal templates for MMEJ rather than the HDR templates (45, 46). This interesting alternative is important, as we and authors of study reports on other organisms have previously attributed HDR as the sole mechanism of oligonucleotide donor-directed DSB repair (7, 47, 48). This would not, however, explain why the double-stranded donors with long homology arms (100 to 1,000 bp) were often less efficient in DSB repair than the short single-strand oligonucleotide donors (47, 48). These results also argue that Cas9 cleavage is quite efficient; it can quickly cleave the second allele following the first allele cleavage or it can generate DSBs on both alleles simultaneously.

In summary, we have described the optimization and simplification of CRISPR-Cas9 genome editing in Leishmania. Sequential transfections of oligonucleotide donors and cotargeting MT significantly increase gene-editing efficiency. The single gRNA and Cas9 coexpression vectors that were successfully used in all three tested species further simplify this CRISPR-Cas9 system for Leishmania. With this optimization, it was possible to delete the multicopy A2 gene family and to generate targeted chromosomal translocations, which will further advance studies on pathogenesis and polycistronic transcription control in Leishmania. By directly observing the death of CRISPR-targeted RAD51 null mutant clones, we demonstrated that the RAD51 DNA recombinase is essential for L. donovani, confirming that this CRISPR system can be effectively used to determine the essentiality of a Leishmania gene. We have presented evidence to argue that MMEJ plays a more important role than HDR in DSB repair in L. donovani. The CRISPR-Cas9 technology described within has greatly improved the ability to manipulate the Leishmania genome and provides new knowledge about DNA repair in L. donovani. We foresee that CRISPR-Cas9 will soon be widely used in Leishmania research and will eventually help control and eliminate leishmaniasis.

MATERIALS AND METHODS

Leishmania strains and culture medium. L. donovani 1 S/C2D, L. major Friedlin v9, and L. mexicana (MNYC/BZ/62/M379) used in this study were routinely cultured at 27°C in M199 medium (pH 7.4) supplemented with 10% heat-inactivated fetal bovine serum, 40 mM HEPES (pH 7.4), 0.1 mM adenine, 5 mg liter⁻¹ hemin, 1 mg liter⁻¹ biotin, 1 mg liter⁻¹ biopterine, 50 μU ml⁻¹ penicillin, and 50 μg ml⁻¹ streptomycin. Cultures were passaged to fresh medium at a 20-fold dilution once a week. Leishmania cells after transfection with the CRISPR vector and donor DNA were sometimes cultured at 33°C or 37°C for 2 to 3 days to improve gene-editing efficiency (7).
Plasmid construction. All primer sequences used in this study are listed in the supplemental material (Data Sets S1 to S6).

The LdU6gRNAahas plasmid was generated as follows. (i) A 184-bp PCR fragment containing the 98-bp L. donovani rRNA polymerase III U6 promoter was amplified from L. donovani genomic DNA with primers LdU6pf and LdU6pr, which also contains the LdMT gRNA guide coding sequence, and digested with HindIII and BbsI. (ii) The fragment from step 1 was cloned into the HindIII and BbsI sites of the pSPneo vector to create HumanU6gRNAplasmid. (iii) The LdMT gRNA guide coding sequence was then inserted into LdonovanirNA plasmid after BbsI digestion.

The plLCN plasmid was generated as follows. (i) A 495-bp PCR fragment containing the human U6 promoter and gRNA coding sequence was amplified from plasmid pX330 (26) with primers pX330gRNAF1 and pX330gRNAar. (ii) The fragment from step 1 was digested and cloned into the HindIII and BamHI sites of pSPneo vector to create HumanU6gRNA plasmid. (iii) The LdMT gRNA guide coding sequence was then inserted into the LdonovanirNA plasmid after BbsI digestion.

The humanU6gRNAhas plasmid was generated as follows. (i) A 740-bp PCR fragment containing the human U6 promoter and gRNA coding sequence was amplified from plasmid pX330 (26) with primers pX330gRNAF1 and pX330gRNAar. (ii) The fragment from step 1 was digested and cloned into the HindIII and BamHI sites of pSPneo vector to create HumanU6gRNA plasmid. (iii) The LdMT gRNA guide coding sequence was then inserted into the LdonovanirNA plasmid after BbsI digestion.

The plCD plasmid was generated as follows. (i) A 495-bp PCR fragment containing the 180-bp L. donovani rRNA promoter, the gRNA and HDV ribozyme coding sequences, and the 92-bp pyrimidine track was obtained from the pSPneo vector (7) with primers LdrRNApNde1 and pSPneoRHind3. (ii) The PCR fragment from step 1 was cloned into Ndel and HindIII sites of the pSP72 vector. (iii) The 6,595-bp HindIII and BglII fragment containing the Cas9 and hygromycin resistance genes from plPhyg-Cas9 (7) was subsequently cloned into the corresponding sites of the pSP72 vector in step ii to generate the plCD plasmid.

The plCDN plasmid was generated as follows. (i) A 495-bp PCR fragment containing the 180-bp L. donovani rRNA promoter, the gRNA and HDV ribozyme coding sequences, and the 92-bp pyrimidine track was obtained from the pSPneo vector (7) with primers LdrRNApNde1 and pSPneoRHind3. (ii) The fragment from step 1 was cloned into the HindIII and BamHI sites of the pSP72 vector. (iii) The 4.4-kb HindIII and BglII fragment, which contains the LdrRNAP and gRNA 241510 coding sequence from the pSPneo vector (49) to generate pLPneoCas9. (iv) The 6.4-kb HindIII and BglII fragment containing the Cas9 and neomycin resistance genes from the pSPneoCas9 plasmid was subsequently cloned into the corresponding sites of the pSP72 vector in step ii to generate the plCDN plasmid.

The gRNA 241510+MT(gRNAa) coexpression vector was generated by inserting the 360-bp HindIII and BamHI fragment, which contains the LdRNAP and gRNA 241510 coding sequence from the pSPneo vector (7) after removing the 180-bp HindIII and BamHI fragment.

The gRNA A2a+b coexpression vector was generated as follows. (i) A 276-bp PCR fragment containing gRNA A2a, HDV, and hammerhead ribozyme coding sequences was amplified with primers Ld220670a and Ld220670b from the gRNA 241510 MT coexpression vector. (ii) The PCR product from step 1 was digested with BbsI and inserted into the BbsI-digested pSPneoRiNAH vector.

The gRNA A2a+b+MT(gRNAa) triple expression vector was generated by inserting the 579-bp HindIII and BamHI fragment, which contains LdRNAP, gRNA A2a+b, and ribozyme coding sequences from the gRNA A2a+b coexpression vector into the HindIII and BglII sites of the pSPneoRiNAH vector after removing the 180-bp HindIII and BamHI fragment.

gRNA and primer design and synthesis. Since current gRNA design tools developed from data for higher-order eukaryotic cells are not necessarily suitable for Leishmania (7), we selected gRNA guide sequences based on the relatively high activity scores in all of the following three design programs and not off-target sites. This method used a target sequence (table) to determine mitofusine resistance rates (7). The microhomology sequences are required for MMEJ but should be avoided if a donor will be used. Sequence scan for CRISPR (SSC) (based on human and mouse data; http://grna.cregd.jega.edu/) provides useful information on off-target sites and microhomology sequences flanking the DSB (9). The microhomology sequences are required for MMEJ but should be avoided if a donor will be used. Sequence scan for CRISPR (SSC) provides useful information on off-target sites and microhomology sequences flanking the DSB (9). The microhomology sequences are required for MMEJ but should be avoided if a donor will be used.

The following experimental procedures were performed as previously described: single gRNA guide sequence cloning into various gRNA expression vectors (7); Leishmania transfection and a limiting dilution assay to determine mitofusine resistance rates (7); Leishmania genomic DNA extraction, PCR, and sequencing analysis (7); A2 Western blot analysis (14).

RAD51 inhibition assay. The stock solutions (10 mM) of Rad51 inhibitors B02 and RI-1 (catalog numbers SML0364 and 1274; Sigma) were prepared in dimethyl sulfoxide and stored at 4°C (42–44). Immediately before use, the stock solutions were diluted with Leishmania culture medium to a 1 mM working concentration for RI-1 and 100 μM for B02. These inhibitors were then directly added into Cas9 and LdMT gRNA-expressing Leishmania culture medium (1 × 10^6 promastigotes per ml) to a final concentration of 0 to 20 μM for B02 and 0 to 100 μM for RI-1. The proper concentrations of these inhibitors in culture were maintained by adding fresh inhibitors once every 3 days for a total 4 times in a 2-week period before measuring the mitofusine resistance rate.

Generation of LdMT^+/− single-knockout cells via the traditional gene replacement method. The specific LdMT^−/− bleomycin-targeting fragment was generated by overlapping PCR. (i) The 666-bp LdMT^−/− 5′ flank fragment targeted by overlapping PCR. (ii) The 666-bp LdMT^−/− 5′ flank fragment was flanked by primers Ld131590F and Ld131590R, the 334-bp bleomycin expression cassette with primers 1315908F and 1315908R, and the 666-bp LdMT^−/− 3′ flank fragment targeted by overlapping PCR. (iii) The 334-bp bleomycin expression cassette with primers 1315908F and 1315908R and the 666-bp LdMT^−/− 3′ flank fragment were PCR amplified separately. (iv) The 334-bp bleomycin expression cassette with primers 1315908F and 1315908R to generate
the specific 1,857-bp LdMT bleomycin-targeting fragment. L. donovani promastigotes transfected with this LdMT bleomycin-targeting fragment were selected with 100 μg/ml bleomycin. The LdMT+/− single-knockout clones were verified by PCR with primer pair Ld1315905F1 and 1315908B1.

Accession number(s). The pLdCN, pLdCH, and pSPneogRNA241510 plasmids have been deposited in Addgene under numbers 84290, 84291, and 84292, respectively. The partial sequences for these plasmids are provided in the supplemental material (Data Sets S3 and S5).

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mSphere.00340-16.

DATA SET S1, DOCX file, 0.04 MB.
DATA SET S2, DOCX file, 0.02 MB.
DATA SET S3, DOCX file, 0.01 MB.
DATA SET S4, DOCX file, 0.01 MB.
DATA SET S5, DOCX file, 0.03 MB.
DATA SET S6, DOCX file, 0.02 MB.

ACKNOWLEDGMENTS

We thank Feng Zhang for plasmid pX330, Marc Ouellette and Barbara Papadopoulos for plasmids pSPneo and pSPhyg, and Dan Zylberstein for anti-HSP83 serum.

This work was supported by Canadian Institute of Health Research grant MOP125996 to G.M.

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